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In vitro propagation of *Cyathea atrovirens* (Cyatheaceae): spore storage and sterilization conditions

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Abstract: *Cyathea atrovirens* occurs in a wide range of habitats in Brazil, Paraguay, Uruguay and Argentina. In the Brazilian State of Rio Grande do Sul, this commonly found species is a target of intense exploitation, because of its ornamental characteristics. The *in vitro* culture is an important tool for propagation which may contribute toward the reduction of extractivism. However, exogenous contamination of spores is an obstacle for the success of aseptic long-term cultures. This study evaluated the influence of different sterilization methods combined with storage conditions on the contamination of the *in vitro* cultures and the gametophytic development of *C. atrovirens*, in order to establish an efficient propagation protocol. Spores were obtained from plants collected in Novo Hamburgo, State of Rio Grande do Sul, Brazil. In the first experiment, spores stored at 7°C were surface sterilized with 0.5, 0.8 and 2% of sodium hypochlorite (NaClO) for 15 minutes and sown in Meyer’s culture medium. The cultures were maintained in a growth room at 26±1°C for a 12-h photoperiod and photon flux density of 100µmol/m²/s provided by cool white fluorescent light. Contamination was assessed at 60 days, and gametophytic development was scored at 30, 60, 120 and 130 days of *in vitro* culture, analyzing 300 individuals for each treatment. There was no significant difference in culture contamination among the different sodium hypochlorite concentrations tested, and all treatments allowed for the development of cordiform gametophytes at 130 days of culture. In the second experiment, spores stored at 7 and -20°C were divided into two groups. Half of the spores were surface sterilized with 2% of NaClO for 15 minutes and the other half was not sterilized. All spores were sown in Meyer’s medium supplemented with one of the following antibiotics: nystatin, Micostatin® and actidione. The culture conditions and the procedures used for evaluating contamination and gametophytic development were the same described for the first experiment. No contamination was observed in spores stored at -20°C and treated with NaClO and actidione. In all treatments, cordiform gametophytes presenting antheridia were observed at 120 days. The percentages of these gametophytes increased from 120 to 130 days and no significant differences were observed among treatments. Archegonia were observed on cordiform gametophytes at 130 days. The findings provide data relevant to *in vitro* propagation procedures of this species, which may increase the availability of plants for ornamental purposes, therefore contributing to the reduction of the exploitation of endangered tree ferns species. Rev. Biol. Trop. 62 (1): 299-308. Epub 2014 March 01.

Key words: actidione, antibiotic, gametophyte, germination, *in vitro* culture, surface sterilization, tree fern.

Tree ferns are an important component of tropical rainforests and among them Cyatheaceae is a noteworthy family, being represented by approximately 170 species in the Neotropics (Tryon & Tryon, 1982) and 23 species in South and Southeastern regions of Brazil (Windisch & Santiago, 2013). *Cyathea atrovirens* (Langsd. & Fisch.) Domin (Cyatheaceae) occurs in a wide range of habitats in Brazil, Paraguay, Argentina (Ponce, 1996) and Uruguay (Marquez & Brussa, 2011). In the Brazilian State of Rio Grande do Sul, the species is commonly found in open or moderately shaded humid or swampy places and in areas largely affected by
human action, such as roadsides (Lorscheitter, Ashraf, Windisch & Mosbrugger, 1999). *Cyathea atrovirens* forms 6m high caudices, with a sheath of adventitious roots at the persistent petiole bases. Petioles and leaves are, respectively, up to 1.10 and 3m long, and the laminas are bipinnate-pinnatifid to pinatilobate (Sehnem, 1978; Fernandes, 1997). The spores are tetratrahedal-globose and present a triangular equatorial limb with prominent rounded angles and straight to slightly depressed sides (Lorscheitter et al., 1999).

Habitat destruction and fragmentation are the main factors related to the reduction of tree fern populations. In addition, tree ferns are exploited because of their gardening potential (Windisch, 2002). Leaves (Tryon & Tryon, 1982) and entire plants (Schmitt & Windisch, 2012) of *C. atrovirens* are used for ornamental purposes. Caudices of older plants, when presenting a sheath of adventitious roots at the base, are used to manufacture fiber handicrafts (Fernandes, 2000).

The *in vitro* culture is an important tool for plant propagation which may contribute toward the reduction of exploitation of natural populations (Caldecott, Jenkins, Johnson & Groombridge, 1996; Giudice, Giacosa, Luna, Yañez & de la Sota, 2011). Spore germination is considered the most efficient method of *in vitro* culture of ferns and lycophytes (Pence, 2008) and studies with different species can be cited (e.g. Kiss & Kiss, 1998; Bertrand, Albuerner, Fernández, González & Sánchez-Tamés, 1999; Fernández, Bertrand & Sánchez-Tamés, 1999; Kuriyama, Kobayashi & Maeda, 2004; Souza, Medeiros & Mendes, 2007).

In general, studies analyzing the viability of the spores and the initial ontogenetic development of gametophytes do not refer to contamination (Bertrand, Albuerner, Fernández, González & Sánchez-Tamés, 1999; Fernández et al., 1999; Chen, Cheng, Liu & Jiao, 2008), despite exogenous contamination of spores is an obstacle for the success of aseptic long-term cultures (Dyer, 1979; Simabukuro, Dyer & Felippe, 1998). The substances most commonly used for plant tissue surface sterilization are ethylic alcohol and compounds with chlorine, like calcium hypochlorite and sodium hypochlorite (Wu, Ping-Tin, Li-Ping & Long-Qing, 2009). Recently, the sodium dichloroisocyanurate was described as an efficient sterilizing agent for spores (Barnicoat, Cripps, Kendon & Sarasen, 2011). Addition of antimicrobial agents to the culture medium has also been tested. Among antibiotics that can be used to avoid the *in vitro* contamination of plants (Kyte & Kleyn, 1996), Micostatin® (Ranal, 1991), nystatin (Dyer, 1979; Simabukuro et al., 1998; Quintanilla, Amigo, Pangua & Pajaron, 2002) and streptomycin (Cox, Bhatia & Ashwath, 2003) were tested for ferns, although without total elimination of the microorganisms from the cultures. The treatment of plant tissues with sodium hypochlorite, followed by the addition of the fungicide Benlate® (Benomil - Dupont) to the culture medium has also been used for fern species (Brum & Randi, 2006; Begnini & Randi, 2009; Viviani & Randi, 2008; Santos, Lehmann, Santos & Randi, 2010). However, this fungicide had its manufacture discontinued since 2001 (Dupont, 2012). Unfortunately, most surface sterilization treatments may also reduce fern spore germination (Hamilton & Chaffin, 1998; Simabukuro et al., 1998).

Storage conditions under different temperatures have been tested focusing on the survival and germination of spores (Simabukuro et al., 1998; Rogge, Viana & Randi, 2000; Quintanilla et al., 2002; Begnini & Randi, 2009; Santos et al., 2010). The viability of spores from different species has been investigated after cold storage (for a revision, see Ranker & Hauffler, 2008), and low temperature was considered as minimizing bacterial and fungal contamination without decreasing spore viability (Quintanilla et al., 2002).

The aim of this study was to test the influence of different sterilization methods combined with storage conditions on the contamination of the *in vitro* cultures and the gametophytic development of *C. atrovirens*, in order to contribute to the establishment of an efficient propagation protocol contributing to management programs of this species.
MATERIAL AND METHODS

Fertile leaves of *C. atrovirens* were collected in the Parque Municipal Henrique Luis Roessler (29°40'54" S - 51°06'56" W, at 16.4m in elevation), which is a conservation area of the Rio dos Sinos basin in the municipality of Novo Hamburgo, State of Rio Grande do Sul, Brazil. The leaves were wrapped in smooth paper and kept at room temperature to induce dehiscence of sporangia. Spores were filtered through lens cleaning tissue and stored. Two different experiments were carried out as follows.

**Experiment I:** In the first experiment, spore samples (~30mg) stored at 7°C for 30 days were distributed in microtubes of 1.5mL and surface sterilized in 1mL of sodium hypochlorite (NaClO) solution at concentrations of 0.5, 0.8 and 2.0% (v/v) for 15min. Spores immersed in 1mL of sterile distilled water instead of NaClO were used as control. After removing the supernatant, spores were rinsed in 1mL of sterile distilled water, and centrifuged at 2 000rpm for 3min. The washing and centrifugation steps were repeated three times. The spores were sown in petri dishes (9cm diameter, 10mg/dish) containing 30mL of Meyer’s medium (Meyer, Anderson & Swanson, 1955), supplemented with 0.25% (w/v) Phytagel®, with pH adjusted to 6.0 before autoclaving. For each NaClO concentration tested and for the control, three repetitions were made. All procedures were carried out in a laminar hood. The cultures were maintained in a growth room at 26±1°C for a 12-h photoperiod and photon flux density of 100µmol/m²/s provided by cool white fluorescent light. Once a month, 2mL of sterile distilled water (pH 6.0) were added to the culture medium in each dish, to stimulate the gametophytic development (Mendoza-Ruiz & Pérez-García, 2009).

To assess the contamination at 60 days of culture, a quantitative analysis was performed, using two printed paper grids, each containing 21 fields (each field of 1.5x1.5cm), a black one and a white one, for better visualization of fungi and bacteria. The grids were placed separately under each dish, covering the entire area, and the number of fields presenting macroscopically visible contamination was counted.

Gametophytic development was scored at 30, 60, 120 and 130 days of *in vitro* culture. Three slides (one slide per dish) from each treatment were analyzed, and 100 individuals (spores or young gametophytes) were counted on each slide (300 individuals per treatment). Individuals were classified according to their developmental stage (Rechenmacher, Schmitt & Droste, 2010) in the following classes: non-germinated spores (NG), gametophyte with chlorocyte and rhizoid (G1); filamentous gametophyte (G2); laminar gametophyte (G3) and cordiform gametophyte (G4). The criterion for germination was the emergence of the chlorocite or the rhizoid (Ranal, 1999).

**Experiment II:** In a second experiment, spore samples (~30 mg) stored at 7 and -20°C for 30 days were distributed in microtubes of 1.5mL. For each storage temperature, spores of three tubes were surface sterilized with 2% (v/v) NaClO for 15min, and spores of three tubes were immersed in sterile distilled water for the same time (control), according to the procedure described for the first experiment. The spores were sown in Petri dishes (9cm in diameter, 10 mg/dish), with 30mL of Meyer’s medium (pH 6.0 before autoclaving), supplemented with one of the following antibiotics: (a) 1mL of nystatin (Sigma-Aldrich) 10 000U/mL, (b) 1mL of Micostatin® (Bristol-Myers Squibb), a commercial antibiotic containing 100 000U/mL of nystatin, and (c) 0.5g/L of actidione (Fluka, Sigma-Aldrich). The antibiotics were added to the medium after autoclaving and partial cooling. For each treatment, that combined a different antibiotic and storage temperature with or without previous surface sterilization, three dishes were used, totaling 36 dishes. The culture conditions and the procedures used for evaluating contamination and gametophytic development were the same described for the first experiment.
The statistical analyses of the data were performed using the SPSS 17.0 and the BioEstat 5.0 softwares. The Shapiro-Wilk and Levene tests, respectively for goodness of fit for normality and for the homogeneity of variance (0.05) were applied before the analysis. Data related to contaminated fields in the first experiment, as well as data of gametophytic development in both experiments were submitted to the nonparametric Kruskal-Wallis test followed by the Student-Newman-Keuls test, at 0.05 significance. Data of gametophytic development in the second experiment were submitted to analysis of variance (ANOVA) and the pairwise comparison between means was performed by the parametric Tukey test at 0.05 significance (Zar, 1999).

RESULTS

Experiment I: In the first experiment, contamination was observed in all cultures. A minor number of contaminated fields were visualized in cultures when spores were treated with 2.0% of NaClO (9.6) compared to 0% (21.0), 0.5% (14.0) and 0.8% of NaClO (18.6), although these differences were no statistically significant (Kruskal-Wallis, H=5.0600, p=0.1675). Germination and gametophytic development were observed in cultures from the three NaClO concentrations tested (Table 1). At 30 days of culture, most gametophytes were on laminar stage and cordiform gametophytes only have been observed in treatments with 2.0% of NaClO (Table 1). At 60 days,

<table>
<thead>
<tr>
<th>Days of culture</th>
<th>Stage1</th>
<th>NaClO (%)</th>
<th>0.0</th>
<th>0.5</th>
<th>0.8</th>
<th>2.0</th>
<th>H²</th>
<th>p³</th>
</tr>
</thead>
<tbody>
<tr>
<td>30</td>
<td>NG</td>
<td>6.3±1.5a</td>
<td>0a</td>
<td>4.0±5.1a</td>
<td>3.6±4.7a</td>
<td>5.494</td>
<td>0.139</td>
<td></td>
</tr>
<tr>
<td></td>
<td>G1</td>
<td>11.6±3.2a</td>
<td>1.6±0.5b</td>
<td>1.0±0.1b</td>
<td>3.3±1.5ab</td>
<td>9.000</td>
<td>0.029</td>
<td></td>
</tr>
<tr>
<td></td>
<td>G2</td>
<td>31.3±7.6a</td>
<td>9.6±3.0a</td>
<td>14.7±5.7a</td>
<td>12.6±2.0a</td>
<td>7.333</td>
<td>0.062</td>
<td></td>
</tr>
<tr>
<td></td>
<td>G3</td>
<td>50.8±5.8a</td>
<td>88.8±3.5b</td>
<td>80.3±10.0ab</td>
<td>79.3±1.1ab</td>
<td>8.157</td>
<td>0.043</td>
<td></td>
</tr>
<tr>
<td></td>
<td>G4</td>
<td>0a</td>
<td>0a</td>
<td>1.0±1.7a</td>
<td>0.692</td>
<td>0.875</td>
<td></td>
<td></td>
</tr>
<tr>
<td>60</td>
<td>NG</td>
<td>2.5±0.7a</td>
<td>0a</td>
<td>0.6±1.1a</td>
<td>0a</td>
<td>4.540</td>
<td>0.209</td>
<td></td>
</tr>
<tr>
<td></td>
<td>G1</td>
<td>6.0±1.4a</td>
<td>0a</td>
<td>0a</td>
<td>0a</td>
<td>4.500</td>
<td>0.212</td>
<td></td>
</tr>
<tr>
<td></td>
<td>G2</td>
<td>26.0±19.7a</td>
<td>5.4±4.6a</td>
<td>13.5±8.6a</td>
<td>1.4±2.3a</td>
<td>6.208</td>
<td>0.102</td>
<td></td>
</tr>
<tr>
<td></td>
<td>G3</td>
<td>65.5±17.6a</td>
<td>92.3±3.2a</td>
<td>82.6±7.3a</td>
<td>94.3±5.6a</td>
<td>6.040</td>
<td>0.102</td>
<td></td>
</tr>
<tr>
<td></td>
<td>G4</td>
<td>0a</td>
<td>2.3±1.5a</td>
<td>3.3±2.8a</td>
<td>4.3±3.5a</td>
<td>3.994</td>
<td>0.262</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Days of culture</th>
<th>Stage1</th>
<th>NaClO (%)</th>
<th>0.0</th>
<th>0.5</th>
<th>0.8</th>
<th>2.0</th>
<th>H²</th>
<th>p³</th>
</tr>
</thead>
<tbody>
<tr>
<td>120</td>
<td>G2</td>
<td>0.6±1.1a</td>
<td>1.1±1.7a</td>
<td>0a</td>
<td>0.622</td>
<td>0.733</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>G3</td>
<td>94.8±1.5a</td>
<td>88.3±1.5a</td>
<td>81.3±12.0a</td>
<td>4.356</td>
<td>0.113</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>G4</td>
<td>4.6±2.0a</td>
<td>10.6±3.2a</td>
<td>18.7±12.0a</td>
<td>3.922</td>
<td>0.141</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Days of culture</th>
<th>Stage1</th>
<th>NaClO (%)</th>
<th>0.0</th>
<th>0.5</th>
<th>0.8</th>
<th>2.0</th>
<th>H²</th>
<th>p³</th>
</tr>
</thead>
<tbody>
<tr>
<td>130</td>
<td>G2</td>
<td>0a</td>
<td>1.0±1.7a</td>
<td>0a</td>
<td>0.600</td>
<td>0.741</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>G3</td>
<td>93.4±2.0a</td>
<td>87.0±2.6a</td>
<td>75.7±10.0a</td>
<td>5.956</td>
<td>0.051</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>G4</td>
<td>6.6±2.0a</td>
<td>12.0±4.3a</td>
<td>24.3±10.0a</td>
<td>5.067</td>
<td>0.079</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

1 NG = no germinated spores; G1 = gametophyte with chlorocyte and rhizoid; G2 = filamentous gametophyte; G3 = laminar gametophyte; G4 = cordiform gametophyte.
2 H = Kruskal-Wallis test.
3 p = significance level.
4 NE = not evaluated at 60 days due to contamination.

• Means followed by different letters within each row differ significantly according to the Student-Newman-Keuls test (p ≤ 0.05).

TABLE 1
Influence of sodium hypochlorite (NaClO) on the percentage of gametophytes of Cyathea atrovirens in vitro cultures
gametophytes in the first stage (with chloro-
cytes and rhizoids) were present only in the
treatment without NaClO and most gameto-
phytes were in laminar stage. In the following
observations, there was an increase in the
percentage of cordiform gametophytes in all
treatments with NaClO. The highest percentage
of this type of gametophytes was observed after
treatment with 2% of NaClO, although without
significant difference to the other treatments
with this sterilizing agent (Table 1). Antheridia
were observed in the control and in treatments
with 0.5 and 2.0% of NaClO at 60 days, and
also in treatment with 0.8% at 120 days. Arche-
gonia were seen only in the treatment with 2%
NaClO, at 120 days of cultivation.

Although there was no statistically diffe-
rence in culture contamination among the
different sodium hypochlorite concentrations
tested in the first experiment, the treatment
with 2% NaClO allowed for numerically lower
contamination and higher percentage of cor-
diform gametophytes at 130 days of culture.
Therefore, this treatment was used in the
second experiment. The lowest mean numbers
of contaminated fields were observed when
spores were surface sterilized with NaClO
combined with the addition of antibiotics to the
culture medium (Table 2). For the spores stored
at 7°C and previously surface sterilized, the use
of actidione did not differ significantly from
the use of nystatin, which led to significantly
less contamination than using Micostatin®.
When there was no prior sterilization of spores
stored at 7°C, the use of actidione provided
significantly less contaminated fields in rela-
tion to the use of nystatin and Micostatin®. For
spores stored at -20°C and previously surface
sterilized, no significant difference among the
antibiotics tested was verified. Contamination
was low in treatments with nystatin and Micos-
tatin®, and the use of actidione eliminated con-
tamination totally. For spores stored at -20°C,
without prior sterilization with NaClO, the use
of actidione showed the lowest mean contami-
nation values, with significant difference com-
pared to the other antibiotics tested (Table 2).

**Experiment II:** In the second experiment,
the development of gametophytes was studied
when subjected to antibiotic treatments and pre-
vious sterilization of spores with NaClO. At 30
days, the percentages of laminar gametophytes
were significantly higher in treatments with
Micostatin® than in treatments with actidione,
and cordiform gametophytes could already
be observed in treatments with this antibiotic
(Table 3). At 60 days, most gametophytes were
in laminar stage, and cordiform gametophytes
were registered in treatments with nystatin and
Micostatin®. There was an increase in the num-
ber of laminar gametophytes from 60 to 120
days of culture in the treatments with nystatin
and actidione. In the presence of Micostatin®,

**TABLE 2**
Influence of storage temperatures and sterilization treatments on contamination1 of *Cyathea atrovirens* in vitro cultures

<table>
<thead>
<tr>
<th>Antibiotic</th>
<th>Storage temperature / Surface sterilization</th>
<th>7°C</th>
<th>-20°C</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>NaClO</td>
<td>No NaClO</td>
<td>NaClO</td>
</tr>
<tr>
<td>Nystatin</td>
<td>0.3±0.5a</td>
<td>21.0±0b</td>
<td>5.0±4.0a</td>
</tr>
<tr>
<td>Micostatin®</td>
<td>8.7±4.0b</td>
<td>21.0±0b</td>
<td>4.6±1.1a</td>
</tr>
<tr>
<td>Actidione</td>
<td>0.3±0.5a</td>
<td>2.3±0.5a</td>
<td>0±0a</td>
</tr>
<tr>
<td>F2</td>
<td>12.2</td>
<td>3136.0</td>
<td>4.0</td>
</tr>
<tr>
<td>p3</td>
<td>0.008</td>
<td>&lt;0.001</td>
<td>0.077</td>
</tr>
</tbody>
</table>

- Means followed by different letters within each column differ significantly according to the Tukey test (p≤0.05).
1. From a total of 21 fields.
2. F=ANOVA test.
3. p=significance level.
the number of laminar gametophytes decreased, because some of them have already developed into cordiform gametophytes. However, no statistical differences were found on laminar gametophyte percentages among treatments. In all treatments, cordiform gametophytes presenting antheridia were observed at 120 days. At 130 days, no filamentous gametophytes were found, and in all treatments most gametophytes were of the laminar type. The percentages of cordiform gametophytes increased from 120 to 130 days and no significant differences were observed among treatments (Table 3). Archegonia were observed on cordiform gametophytes at 130 days.

### DISCUSSION

Although without significant statistical difference among treatments, the contamination of cultures coming from spores sterilized with 2% NaClO was about 50% lower than the contamination of cultures from spores not treated with this substance. Due to exogenous contamination, surface sterilization of spores is the first step in an aseptic culture of ferns (Dyer, 1979). Cultures initiated with unsterilized biological material tend to contaminate after 10 days, as noted by Camloh (1999) for *Platycerium bifurcatum* (Cav.) C. Chr. and Rechenmacher et al. (2010) for *C. atrovirens*.

### TABLE 3
Influence of storage temperatures and sterilization treatments on the percentage of gametophytes of *Cyathea atrovirens* in vitro cultures

<table>
<thead>
<tr>
<th>Days</th>
<th>Stage</th>
<th>NaClO/Nystatin</th>
<th>NaClO/Micostatin®</th>
<th>NaClO/Actidione</th>
<th>H²</th>
<th>p³</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>7°C</td>
<td>-20°C</td>
<td>7°C</td>
<td>-20°C</td>
<td>7°C</td>
</tr>
<tr>
<td></td>
<td></td>
<td>mean±standard deviation (%)</td>
<td>mean±standard deviation (%)</td>
<td>mean±standard deviation (%)</td>
<td>mean±standard deviation (%)</td>
<td>mean±standard deviation (%)</td>
</tr>
<tr>
<td>30</td>
<td>NG</td>
<td>2.0±1.0b</td>
<td>5.0±1.0ab</td>
<td>8.0±0.0a</td>
<td>4.0±1.4ab</td>
<td>7.6±4.0</td>
</tr>
<tr>
<td></td>
<td>G1</td>
<td>10.2±4.0bc</td>
<td>12.7±1.5abc</td>
<td>10.4±3.2bc</td>
<td>6.0±0.0c</td>
<td>31.0±11.0a</td>
</tr>
<tr>
<td></td>
<td>G2</td>
<td>32.6±20.5ab</td>
<td>20.7±9.8ab</td>
<td>5.6±2.5b</td>
<td>3.0±1.4b</td>
<td>42.4±12.5a</td>
</tr>
<tr>
<td></td>
<td>G3</td>
<td>55.2±18.9ab</td>
<td>61.6±9.2ab</td>
<td>75.0±4.5a</td>
<td>86.0±1.4a</td>
<td>14.0±12.7b</td>
</tr>
<tr>
<td></td>
<td>G4</td>
<td>0a</td>
<td>0a</td>
<td>1.0±1.0a</td>
<td>1.0±1.4a</td>
<td>0a</td>
</tr>
</tbody>
</table>

| 60   | NG    | 1.6±0.5ab | 3.7±0.5a | 0b | 0b | 4.0±1.0a | 1.6±1.5ab | 13.755 | 0.017 |
|      | G1    | 0b | 3.0±1.0ab | 0b | 0b | 4.3±0.5a | 0b | 11.703 | 0.039 |
|      | G2    | 7.4±2.5ab | 15.7±13.8a | 0b | 0b | 8.4±2.5a | 7.0±3.6ab | 12.007 | 0.035 |
|      | G3    | 88.0±6.2ab | 77.6±13.3b | 95.7±3.2a | 98.0±1.0a | 83.3±1.5b | 91.4±4.0ab | 12.795 | 0.025 |
|      | G4    | 0.6±0.57ab | 0b | 4.3±3.2a | 2.0±1.0a | 0b | 0b | 12.516 | 0.028 |

| 1120 | G1    | 0a | 0a | 0a | 0a | 0.3±0.5a | 0a | 0.789 | 0.978 |
|      | G2    | 0a | 0a | 0a | 0a | 2.7±2.5a | 0a | 3.158 | 0.676 |
|      | G3    | 93.7±3.5a | 94.0±3.6a | 83.0±13.2a | 74.6±25.7a | 95.7±2.8a | 91.6±2.5a | 7.154 | 0.209 |
|      | G4    | 6.3±3.6ab | 6.0±3.6ab | 17.0±12.7a | 25.4±25.7a | 1.3±1.5b | 8.4±2.5ab | 11.168 | 0.048 |

| 130  | G3    | 90.0±2.6a | 91.0±2.6a | 80.6±12.6a | 72.0±25.2a | 93.7±2.0a | 89.0±3.6a | 9.301 | 0.098 |
|      | G4    | 10.0±2.6a | 9.0±2.6a | 19.4±12.6a | 28.0±25.2a | 6.3±2.0a | 11.0±3.6 ab | 9.301 | 0.098 |

- Means followed by different letters within each line differ significantly according to the Student-Newman-Keuls test (p≤0.05).
- NG=no germinated spores, G1=gametophyte with chlorocyte and rhizoid, G2=filamentous gametophyte, G3=laminar gametophyte, G4=cordiform gametophyte.
- H=Kruskal-Wallis test.
- p=significance level.
On the other hand, although surface sterilizing treatments are commonly referred to as drastically reducing germination (Simabukuro et al., 1998), this did not occur in the present work. The reduction of microorganisms of in vitro cultures observed in the present study for spores sterilized with NaClO, also was recorded for other tree ferns. Low fungal contamination and the elimination of bacterial contamination on Cyathea delgadii Sternb. (Cyatheaceae) were observed by Simabukuro et al. (1998) when using NaClO concentrations of 1.0, 3.0 and 5.0%. Souza et al. (2007) reported contamination by fungi and bacteria in about 50% of the culture dishes when spores of Dicksonia sellowiana Hook. (Dicksoniaceae) where sterilized in 2% of NaClO.

The surface sterilization of spores with NaClO associated with the addition of the fungicide Benlate® (benomyl) to the nutrient medium has been used as an alternative to eliminate contamination of fern cultures in Brazil (Renner & Randi, 2004; Brum & Randi, 2006; Begnini & Randi, 2009; Viviani & Randi, 2008; Santos et al., 2010). However, this fungicide is not a viable alternative, since it is no longer available in the market of this country. Nystatin is the active ingredient of the commercial antibiotic Micostatin®. In the present study, nystatin allowed less contamination than Micostatin®, when combined with the use of NaClO in spores stored at 7°C, although a 10-fold higher concentration of nystatin is present in 1mL of Micostatin® compared with the use of pure nystatin. Ranal (1991, 1999) did not find contamination by fungi in cultures of different fern species from spores stored at 4°C and developed in medium with Micostatin® (E.R. Squibb 10 000U/mL). On the other hand, according to Kozai (1991) and Souza et al. (2007), the presence of sucrose among the inactive ingredients of Micostatin® may favor the development of fungi and bacteria and thereby increase the rate of contamination.

Nystatin was previously used for spore surface sterilization. Filippini, Duz & Randi (1999) cited the use of NaClO and nystatin on in vitro culture of D. sellowiana. Quintanilla et al. (2002) considered the use of 100U/mL of nystatin efficient to reduce fungus and bacteria in wet storage of spores of five threatened fern species: Culcita macrocarpa C. Presl (Culci-taceae), Dryopteris aemula (Aiton) O. Kuntze (Dryopteridaceae), D. corleyi Fraser-Jenkins (Dryopteridaceae), D. guanchica Gibby and Jermy (Dryopteridaceae) and Woodwardia radicans (L.) Sm. (Blechnaceae). However, in these studies, no numerical data of contamination were presented. Simabukuro et al. (1998) observed that the best treatment for elimination of bacteria and fungi, in C. delgadii spore cultures, was the surface sterilization with 0.5% of calcium hypochlorite for two minutes and addition of 100U/mL of nystatin to the medium. However, contamination by fungi and bacteria persisted, with 18.3 and 48.3% of contaminated points, respectively, and only 52.7% of the spores germinated.

Actidione is an antibiotic indicated as inhibitor of bacteria and fungi (Panthier, Diem & Dommergues, 1979). In the present study, the use of actidione without previous surface sterilization of spores stored at 7 and -20°C led to significantly less contamination than the other antibiotics tested. The associated use of this antibiotic with previous surface sterilization of spores stored at -20°C allowed the total decontamination of the cultures. Rojas & Ron-dón (1995) showed in vitro growth inhibition of Fusarium decemcellulare Brick using actidione associated with Benlate, and suggested the use of these substances in Mangifera indica L. (Anacardiaceae) field cultures to control this harmful fungus.

Although sterilizing agents are used for in vitro culture of Cyatheaceae, there were no found reports in literature concerning the qualitative and quantitative influence of them on the development of gametophytes. In the present study, the data at 30 days of culture suggest that the gametophytic development has been delayed in treatments with actidione compared to the treatments with Micostatin®, in which higher percentages of laminar gametophytes and even cordiform gametophytes were observed. This delay was not maintained during
the experiment, and there were no significant differences between treatments at the end of the experiment. Furthermore, the presence of reproductive structures at 120 and 130 days corroborate with the findings of Rechenmacher et al. (2010) for C. atrovirens cultures without using sterilizing agents, suggesting that the antimicrobial action of these agents does not seem to influence negatively the development of gametophytes. The increasing number of reproductive structures observed during the evaluations suggests the continuity of the developmental process after the study period of 130 days.

The surface sterilization of spores stored at -20°C associated with the use of actidione in the culture medium may be an efficient method for an aseptic in vitro culture of C. atrovirens, since it showed to eliminate the contamination and to allow the development of gametophytes with reproductive structures. Moreover, the findings provide data relevant to in vitro propagation procedures of this species, which may increase the availability of plants for ornamental purposes, therefore contributing to the reduction of the exploitation of endangered tree ferns species.

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RESUMEN

Propagación in vitro de Cyathea atrovirens (Cyatheaceae): almacenamiento de esporas y condiciones de esterilización. Cyathea atrovirens (Langsd. & Fisch.) Domin (Cyatheaceae) se presenta en una amplia gama de hábitats en Brasil, Paraguay, Uruguay y Argentina. Debido a sus características ornamentales, la especie es objeto de intensa explotación. El cultivo in vitro es una herramienta importante para la propagación lo que puede contribuir a la reducción del impacto de las actividades extractivas. Sin embargo, la contaminación exógena de esporas es un obstáculo para el éxito de cultivos asépticos a largo plazo. Este estudio evaluó la influencia de diferentes métodos de esterilización en combinación con las condiciones de almacenamiento sobre la contaminación de los cultivos in vitro y el desarrollo gametofítico de C. atrovirens. En el primer experimento, las esporas almacenadas a 7°C se esterilizaron superficialmente con 0.5, 0.8 y 2% de hipoclorito de sodio (NaClO) durante 15 minutos y se sembraron en medio de cultivo de Meyer. Aunque no hubo diferencia en la contaminación de los cultivos entre las concentraciones de hipoclorito de sodio de las diferentes pruebas, en el tratamiento con 2% NaClO se observó un mayor porcentaje de gametofitos cordiformes a los 130 días. En el segundo experimento, las esporas almacenadas a 7 y -20°C fueron divididas en dos grupos. La mitad de las esporas se esterilizaron con 2% de NaClO durante 15 minutos y la otra mitad no fue esterilizada. Todas las esporas se sembraron en medio de Meyer suplementado con uno de los siguientes antibióticos: nistatina, Micostatin® o actidiona. No se observó contaminación de las esporas almacenadas a -20°C y tratadas con NaClO y actidiona. En todos los tratamientos, se observaron gametofitos cordiformes con anterídios y arquegonios. Los resultados proporcionan datos relevantes para la propagación in vitro de C. atrovirens, que pueden aumentar la disponibilidad de las plantas para fines ornamentales, contribuyendo así a la reducción de la exploración de las especies de helechos arborescentes en peligro de extinción.

Palabras clave: actidiona, antibiótico, cultivo in vitro, esterilización superficial, gametófito, germinación, helecho arborescente.

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