Chemical sterilization of culture medium for in vitro multiplication of Cochlospermum regium

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ABSTRACT: Cochlospermum regium roots are used in popular medicine and its extract has diverse phytochemical molecules some with antimicrobial activity, consequently exposing this species to genetic erosion risks. Thus, the objective of this study was to develop an in vitro multiplication protocol using chemical sterilization of culture medium. Therefore, explants obtained from apical buds of C. regium seedlings were inoculated into with 0.05mg L−1 NAA and 1mg L−1 BAP sterilized by chemical agent sodium hypochlorite (NaOCl) at 0.001%, 0.003% and 0.005% of active chlorine (Cl). Autoclaved culture medium was used as control. Result showed that the contamination by bacterial at 91 days of cultivation was significantly (P<0.05) controlled by autoclaving, 0.001% and 0.005% Cl. Moreover, the callus induction in the culture medium with 0.001% and 0.005% Cl was, respectively, 30% and 20% major than autoclaving sterilization. There was not significant (P>0.05) in the percentage of shoot induction among the sterilization preparations methods, and 65% of the explants survived in the presence of culture medium with 0.005% Cl. Histological analyses indicated that the Cl did not have any deleterious effects on morphogenic events. These results indicated that the chemical sterilization using 0.001% - 0.005% Cl controlled the fungal and bacterial multiplication in the culture medium and no affected the C. regium explants development, becoming it an alternative to autoclaving method.

Key words: autoclaving, cerrado biome, active chlorine, in vitro cultivation.

INTRODUCTION

The Cerrado is the second biggest Brazilian biome with 4,400 endemic plant species. This biome has been lost large area and so it is considered an important biodiversity hotspots designated for priority conservation (MYERS et al., 2000). Several plants from Cerrado are known as source of medical compounds and food interests (CARAMORI et al., 2004). Cochlospermum regium (Schrank) Pilger, popularly known as ‘algodãozinho do cerrado’ or ‘algodãozinho do cerrado’, is a medicinal species belonging to Bixaceae family and is commonly found in Cerrado (SÓLON et al., 2009; BATISTA et al.,...
2014). Roots and barks from *C. regium* plants are used in popular medicine to treat female reproductive system infections, intestinal inflammation, rheumatoid arthritis, ulcers, leucorrhoea, cancer and gastritis (TRESZVENZOL et al., 2006; SOUZA & FELFILLI, 2006; MOREIRA & GUARIM-NETO, 2009; SOLON et al., 2009; MELO et al., 2011; BATISTA et al., 2014). Phytochemical analyses of *C. regium* extract indicated the presence of essential oils, flavonoids, saponins, triacylbenzenes, tannins, phenol derivatives, steroidal triterpenes and gallic acid derivatives, providing support to justify its use as popular phytotherapeutic (BRUM et al., 1997; FELFILLI, 2006; MOREIRA & GUARIM-NETO, 2009; TRESVENZOL et al., 2006; SOUZA & FELFILLI, 2006). Moreover, essential oils and hydroalcoholic extracts of *C. regium* roots showed antimicrobial activity against *Staphylococcus aureus*, *Pseudomonas aeruginosa* and *Salmonella typhimurium* (BRUM et al., 1997; SOLON et al., 2009; MELO et al., 2011; INACIO et al., 2014).

The use in an extractive way of *C. regium* roots for phytotherapeutic preparations, the destruction of its habitat and the lack agronomic studies that enable its large-scale cultivation exposes this specie to genetic erosion risks and has placed it on the priority list of native medicinal species for *ex situ* conservation (CAMILLO et al., 2009; INACIO et al., 2010). Thus, tissue culture, through the technique of micropropagation, becomes an alternative for the propagation and conservation. Even though advantages of micropropagation, some problems persist and do not meet commercial interests, especially those associated with the high costs of implementing and maintaining the technique, high energy consumption and time required preparing material of culture medium (MACEK et al., 1995; BRONDANI et al., 2013).

Autoclaving is an energetically expensive operation but is the most commonly sterilization method used in the laboratory. Chemical sterilization of culture medium with sodium hypochlorite (NaOCl) is an alternative method for asepsis of *in vitro* culture of several species, such as *Sequoia sempervirens* (RIBEIRO et al., 2011), *Chrysantemum* (DEEIN et al., 2013), *Eucalyptus benthamii* (BRONDANI et al., 2013), *Hyptis leucochepala* and *H. platani folia* (NEPOMUCENO et al., 2014). Nevertheless, the necessary conditions for the active chlorine not to cause damage to the growth and development of *C. regium* tissues plant are not established. Therefore, the objective of this study was to develop an *in vitro* multiplication protocol of *C. regium* using chemical sterilization of culture medium.

**MATERIALS AND METHODS**

Seeds of *C. regium* were randomly collected between September and October of 2014 from mature fruits of 28 plants located in Cerrado area in the National Park of Chapada dos Guimaraes, ‘Chapada dos Guimaraes’, Mato Grosso, Brazil.

Seeds were washed with distilled and autoclaved water, and subsequently were immersed in sodium hypochlorite solution (i.e., NaOCl, 2.0-2.5% of active chlorine) for 20 minutes. After, triple wash with distilled and autoclaved water was performed, and the seeds were *in vitro* inoculated in MS culture medium (MURASHIGE & SKOOG, 1962) for germination and growth. After 40 days, explants were obtained from apical buds of *C. regium* seedlings and inoculated into test tubes (2×10cm) containing 5mL of MS culture medium supplemented with 0.05mg L$^{-1}$ of α-naphthaleneacetic acid (NAA) and 1mg L$^{-1}$ of 6-benzylaminopurine (BAP).

Culture medium was prepared with distilled water, adding 6g L$^{-1}$ of agar and 30g L$^{-1}$ of sucrose and the pH value was adjusted to 5.8 with HCl (0.1 M) or NaOH (0.1 M). The culture medium was sterilized by addition of the chemical agent sodium hypochlorite (NaOCl) in the concentrations of active chlorine (Cl) of 0.001%, 0.003% and 0.005%. Autoclaved culture medium was used as control, and was performed at 121°C (~ 1kgf cm$^{-2}$) for 20 minutes.

The first subculture occurred at 28 days, totaling 4 subcultures (28, 49, 70 and 91 days). At each subculture, the explants were transferred to the same MS culture medium as previously described. The procedure of manipulation of the explants was carried out in laminar flow chamber under aseptic conditions. The explants were grown in a growth room with controlled temperature at 25°C±2°C, 16 hours of photoperiod (32μmol m$^{-2}$ s$^{-1}$). During the cultivation time (91 days) each explant was evaluated about the percentage of fungal and bacterial contamination, callus induction, number of shoots, oxidation and survival.

To histological analysis, callus samples of *C. regium* were fixed in formaldehyde and modified glutaraldehyde (1% glutaraldehyde, 4% paraformaldehyde in sodium phosphate buffer – 0.1 M NaH$_2$PO$_4$.H$_2$O at pH 7.2) (KARNOVSKY, 1965) and subjected to six series of vacuum (~600mmHg) for 30 minutes each. After 30 days at 4°C the samples were dehydrated by ethyl-alcohol series with increasing concentrations for 15 minutes in each solution from the gradient (10, 20, 30, 40, 50, 60, 70, 80, 90 and 100%, v v$^{-1}$). Then, the samples were immersed in growth conditions.
hydroxyethyl methacrylate resin (Historesin, Leica®, Hildeberg, Germany) and the blocks containing the samples were prepared according to manufacturer’s instructions, being stored for 28 days at 24°C. Blocks containing the samples were sectioned longitudinally or transversely to a thickness of 5μm using an automatic rotary microtome Microm HM 355S (Thermo Scientific). Sections were stained with toluidine blue (0.05%, v v⁻¹) in sodium phosphate buffer and citric acid (SAKAI, 1973) for 30 minutes and mounted on slides with Entellan synthetic resin (Merck®). Histological slides were analyzed and photographed with a light microscope (Opton) and images were captured in micrometric scale.

The experiment was conducted in a completely randomized design in a factorial scheme (4×5), with split-plots subdivided in 4 culture medium preparations (Factor 1 - autoclaved, 0.001%, 0.003% and 0.005% Cl) and 5 subcultivations (Factor 2 - 0, 28, 49, 70 and 91 days). For this, 4 replicates composed of 5 experimental units were used for each replicate. Each experimental unit was composed of one explant. The data collected from experiment were submitted to Shapiro-Wilk’s test (P<0.05) for normal distribution and to Hartley’s test (P<0.05) to verify the variance homogeneity among treatments. When necessary, the data were transformed through Box-Cox’s test (P<0.05). After, the data were submitted to ANOVA (P<0.05 and P<0.01). Duncan’s test was performed when significance was detected by ANOVA. Regression equations were estimated for the sub-cultivations periods, selecting the best adjusted model. Histology was characterized by descriptive analysis.

RESULTS AND DISCUSSION

According to the analysis of variance there was a significant effect for all variables in relation to the effects of culture medium preparation (Factor 1) and sub-cultivations (Factor 2). However, there was no interaction between factors (Factor 1 × Factor 2).

The figure 1A shows the C. regium explants development at each subculture (28, 49, 70 and 91 days) and the formation of callus in the base of the explant, this one being oxidized with the passage of subcultures. Fungal (Figure 1B) and bacterial (Figure 1C) contamination increased during the 91 days but the fungal presence was 15% major than bacteria from the 49 days and approximately 50% of the explants presented oxidation signals (Figure 1D). The percentage of callus (Figure 1E) and shoot induction (Figure 1F) has reached approximately 65% and 47%, respectively, after 49 days and at the 91 days the survival of the explants decreased 40% (Figure 1G). These data shows the in vitro multiplication viability of C. regium as evidenced before (INÁCIO et al., 2011) and allowed to verify that C. regium requires several in vitro subcultures to present morphogenic response in relation to the multiplication of shoots, achieving maximum production at 60 days of cultivation.

The use of active chlorine for the chemical sterilization of culture medium aiming to the in vitro propagation of plants has already been proven for several species (RIBEIRO et al., 2011; BRONDANI et al., 2013; NEPOMUCENO et al., 2014); however it was not recorded in literature for C. regium. Explants were transferred at each subculture, to the same culture medium composition but with different sterilization preparations such as autoclaving, 0.001%, 0.003%, and 0.005% Cl. After 91 days, the percentage of contamination by fungal was no significant different among the sterilization preparations of culture medium (Figure 2A). However, the contamination by bacterial was significantly controlled by autoclaving, 0.001% and 0.005% Cl, while the use of 0.003% Cl was significantly the sterilization method less efficient for bacterial proliferation control (Figure 2B). In the presence of culture medium without active chlorine (i.e., autoclaving method) only 25% of the explants presented oxidation signals, while with active chlorine the oxidation was at least 30% higher (Figure 2C). Callus induction in the culture medium with 0.001% and 0.005% Cl was, respectively, 30% and 20% major than autoclaving preparation (Figure 2D) while the percentage of shoot induction was no significant different among the sterilization preparations methods (Figure 2E). Finally, 65% of the explants survived in the presence of culture medium with 0.005% Cl after 91 days (Figure 2F). These results indicated that 0.001% and 0.005% Cl satisfactorily controlled the fungal and bacterial multiplication in the culture medium and that the presences of 0.005% Cl no affected the C. regium tissue development during 91 days.

The presence of 0.009% and 0.005% Cl in the culture medium for in vitro multiplication of Eucalyptus pellita (TEIXEIRA et al., 2008) and E. benthamii (BRONDANI et al., 2013), respectively, did not affect the shoot viability to the conventional autoclaved preparation. Results also showed that que the presence of 0.005% Cl did not affect shoot induction rate in C. regium explants (Figure 2E). These results show a difference among species regarding the most appropriate active chlorine concentration to
be added to the culture medium, which may be linked to sensitivity to the toxic effects.

As for the oxidation of the explants, results differ from those reported by BRONDANI et al. (2013), which observed oxidation only at concentrations equal to or greater than 0.005% active chlorine when working with *E. benthamii*, whereas for *C. regium* the oxidation of the explants was detected in all treatments. This fact can be justified by the exudation of endogenous substances by the *C. regium* explants after excision, besides the active chlorine to increase the oxidation of tissues. This behavior was observed for each subculture, especially when it was necessary to perform some cut in the explant. However, there are no studies that identify the nature of exuded substances from *C. regium* explants after excision. An alternative to reduce oxidation would be to avoid unnecessary cuts in the explants, as well as the reduction of subculture time, which according to JONES and SAXENA (2013) may contribute to the reduction of tissue oxidation.

Figure 1 - *In vitro* multiplication of *C. regium*. (A) Stages of *in vitro* development of buds and shoots at 28, 49, 70 and 91 days of cultivation. Average percentage of (B) fungal and (C) bacterial contamination, (D) oxidation, (E) callus (F) and shoot induction, and (G) survival in 91 days of cultivation. (B-G) Significant by F test (*P*<0.05).
To verify whether the active chlorine would be affecting the occurrence of morphogenic events, histological analysis from explants of *C. regium* in *vitro* grown at 91 days were performed. Histological analysis showed the presence of small and isodiametric cells in the callus formed indicating areas with intense meristematic activity (Figure 3A). There is presence of unipolar meristematic centers in phase of differentiation (Figure 3B and 3C) and potentially embryonic cells, with asymmetric divisions, similar to that reported in pro-embryonic structures (Figure 3D). The presence of meristematic zones presupposes the formation of adventitious buds and the presence of cells with pro-embryonic characteristics shows the possibility of the occurrence of somatic embryogenesis, which originates from somatic cells, without the occurrence of gametes fusion (CARNEIRO et al., 2014), being an excellent alternative for the asexual propagation and even for the development of synthetic seeds. Thus, these histological results indicate that the active chlorine did not have any deleterious effects on morphogenic events.

According results, the active chlorine supplemented to the culture medium up at concentration of 0.001% - 0.005% was feasible for the *in vitro* multiplication of *C. regium*, when compared with the autoclaving method. Moreover, the chemical sterilization of culture medium demonstrated adequate control of fungal and bacterial contamination, positive response to calogenesis, shoot induction and up to 75% explants survival. Thus, the possibility of chemical sterilization of culture medium with active chlorine was feasible for the *in vitro* multiplication of *C. regium*, becoming an alternative to autoclaving method.

Figure 2 - Autoclaving and chemical sterilization of the culture medium for *in vitro* multiplication of *C. regium* apical buds. Average percentage of (A) fungal, (B) bacterial contamination, (C) oxidation, (D) callus, (E) shoot induction and (F) survival in *C. regium* explants at 91 days of *in vitro* culture. Auto. = Autoclaving method; Chemical sterilization with active chlorine at 0.001%, 0.003% and 0.005%. Means followed by the same letters show no difference by Duncan’s test (*P*<0.05). Data are presented as: mean (± standard error).
CONCLUSION

The chemical sterilization of culture medium by use of active chlorine at 0.001% - 0.005% was efficient for in vitro multiplication of *C. regium*, obtaining similar results to autoclaving method.

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CONFLICTS OF INTEREST

The authors declare no conflict of interest. The founding sponsors had no role in the design of the study; in the collection, analyses, or interpretation of data; in the writing of the manuscript, and in the decision to publish the results.

REFERENCES


*Ciência Rural*, v.48, n.9, 2018.
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