Induction and characterization of oil palm 
(*Elaeis guineensis* Jacq.) pro-embryogenic masses

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**ABSTRACT**

Oil palm is one of the most economically valuable oil seed plants, but the expansion of plantations has been limited by availability of seedlings, as the conventional propagation is through seeds, which have low germination rates. One possible solution for the large-scale production is the use of somatic embryogenesis. The aim of this study was to evaluate the effects of auxins 2,4-D and picloram on the induction of pro-embryogenic masses in *E. guineensis* hybrid leaf explants and characterize, regarding embryogenic characteristics, with cytochemical and ultrastructural analyses. Specifically, in vitro plantlets leaves fragments were inoculated in Y3 culture medium supplemented by 2.4-D or picloram at different concentrations (0.0, 1.0, 3.0, 6.0 and 9.0 mg l⁻¹). After 90 days the presence/absence of cell masses were evaluated. Both growth regulators efficiently induced cellular masses regardless of the concentrations applied. As the cell masses were not homogeneously formed, they were classified according to color and shape into four types: TYPE 1 - elongated and translucent, TYPE 2 – uneven and translucent, TYPE 3 - globular and beige, TYPE 4 – globular and white. Based on the anatomical and ultrastructural features, TYPE 2, 3 and 4 cell masses were considered to have the highest embryogenic potential and therefore may be most suited to large-scale vegetative propagation of oil palm.

**Key words**: embryogenic potential, growth regulators, tissue culture, transmission electron microscopy.

**INTRODUCTION**

The oil palm (*Elaeis guineensis* Jacq. var Tenera) is one of the most economically valuable oil seed plants due to its high oil yield per bunch - reaching up to 6,000 kg ha⁻¹ for certain genotypes. Palm oil is of high quality and is widely used in the food, medicine and cosmetic industries. Moreover, as with other vegetable oils, palm oil can be used to create biodiesel through mixing with petrodiesel or through transesterification (Benjumea et al. 2008, Ghassan et al. 2003).

In order to expand oil palm cultivation and increase biofuel supply, the Brazilian government has recently launched the National Biofuels Programme (Agroanalysis 2007). However, expansion of oil palm plantations has been limited by availability of seedlings, as the conventional propagation is through seeds, which have low germination rates and require a substantial period (1-3 years) to produce seedlings (Martine et al. 2009, Luis et al. 2010).
In vitro propagation is an efficient alternative for the large-scale propagation of many species (Kanchanapoom and Domyos 1999, Steinnacher et al. 2007), including the oil palm (Duval et al. 1988). Somatic embryogenesis is an in vitro propagation method that allows the production of plants from diploid cells without gamete fusion and is one of the most promising methods for cultivation of palm oil (Thomas and Rao 1985). This method ensures the maintenance of important agronomic characteristics and the healthy quality of seedlings. Somatic embryogenesis can occur in vitro in two ways: (i) indirectly, with an intermediate phase characterized by callus formation, and; (ii) directly, without previous callus formation (Guerra et al. 1998). During the indirect somatic embryogenesis different types of callus are frequently produced, some of which have potential to generate plants. Identification of callus cells with embryogenic potential can be achieved through cytological and ultrastructural analysis (Fillipi et al. 2001, Steiner et al. 2005, Moura et al. 2008), thereby greatly enhancing the efficiency of the in vitro propagation process.

The aim of this work was to evaluate the effect of the growth regulators 2,4-D and picloran on the induction of pro-embryogenic masses from oil palm (Elaeis guineensis Jacq. var Tenera) leaf explants. Embriogenic potential of explants was evaluated through cytochemical and ultrastructural analysis.

**MATERIALS AND METHODS**

**PLANT MATERIAL**

This research was performed at the Central Laboratory of Molecular Biology and at Laboratory of Electron Microscopy and Ultrastruttural Analysis (LME), both belonging to the Federal University of Lavras, Minas Gerais State, Brazil. Immature fruits of E. guineensis hybrid Tenera were provided by the company Denpasa, located in Para State, northern Brazil. The fruits (collected approximately 90 to 100 days after pollination) were washed in sodium hypochlorite (1.25%) and broken to remove the epicarp, mesocarp and endocarp thereby exposing the kernels. These were washed in water and placed in a laminar flow chamber for disinfection. The kernels were immersed in 70% ethanol for 30 seconds, then placed in sodium hypochlorite (1.25%) containing Tween, and finally they were washed three times in sterile distilled water under continuous stirring. After disinfection, the embryos were isolated from the kernels and inoculated in a Petri dish containing a modified Y3 culture medium (Eeuwens 1978) supplemented with 45 g L⁻¹ of sucrose, 0.6% (w/v) of agar and pH adjusted to 5.7 ± 0.1. The inoculated embryos were maintained on a photoperiod of 16 hours at 26 ± 2 °C. Every 30 days the embryo were subcultured to flasks containing fresh culture medium.

**SOMATIC EMBRYOGENESIS INDUCTION**

Leaves of in vitro plants were used for induction of pro-embryogenic masses. The leaves explants (approximately 0.5 cm) were inoculated with the adaxial side in contact with the Y3 culture medium (Eeuwens 1978) supplemented with picloran (4-amin acid-3,5,6 - triclor-2-pyridinecarboxylic acid) or 2,4-D (2,4-dichlorophenoxyacetic) at concentrations of 0, 1, 3, 6 e 9 mg.L⁻¹. Culture media were suplemented with sucrose (3%), solidified with agar (0.6%) and the pH were adjusted to 5.7±0.1. After inoculation, the explants were maintained in a growth chamber in the dark at a temperature of 27 ± 2°C. After 90 days the percentage, morphology and color of masses were evaluated.

The presence and absence of cell masses in the leaf explants were assessed through analysis of variance and the averages were compared by Generalized Linear Models. All statistical analysis was performed using SAS® v. 9.3.

**HISTOCHEMICAL AND MORPHOLOGIC ANALYSIS OF CELL MASSES**

The cell masses were fixed in FAA (formaldehyde, acetic acid and alcohol) for 72 hours and transferred...
CHARACTERIZATION OF PRO-EMBRYOGENIC MASSES

After fixing, masses were placed in a 50% alcohol + resin solution overnight and were then transferred to pure resin for 48 h. Finally, they were embedded in Leica resin according to the manufacturer's protocol. Embedded samples were sectioned with a thickness of 5 mm using a rotary microtome and stained with toluidine blue 0.05% solution or lugol solution. The stained sections were then mounted on slides and observed with a photonic Zeiss Scope.A1 microscope with attached camera.

TRANSMISSION ELECTRON MICROSCOPY OF THE CELL MASSES

For analysis in the transmission electron microscope (Zeiss EM 109), samples of cell masses were immersed in fixative (modified Karnovsky, 2.5% glutaraldehyde, 2.0% paraformaldehyde, 0.05 M cacodylate buffer, pH 7.2) for 24 hours and prepared according to the protocol described by Bossola and Russel (1999).

RESULTS AND DISCUSSION

Both growth regulators (2,4-D and picloran, added to the culture medium Y3) efficiently induced cellular masses in leaves of the oil palm *E. guineensis* var. tenera. These masses cellular formed regardless of the growth regulator type and of the concentration used (Figure 1). Moreover, callus formation was not detected in the absence of growth regulators (Figure 2).

Figure 1 - Callus percentage on oil palm tenera hybrid (*Elaeis guineensis*) leaves induced with 2,4D and picloran.

The morphology and staining of the cell masses were not homogeneous, and it was possible to classify them into four types (Figure 2): TYPE 1, elongated and translucent; TYPE 2, uneven and translucent; TYPE 3 globular and beige, and; TYPE 4 globular and white.

These results are similar to those of Bravin et al. (2006), who describe the proliferation of four different callus types on leaf explants of *Hypnea musciformis*. The formation of different types of callus has also been observed in oats *Avena sativa* (Lamb et al. 2002). In this case the phenotypically distinct callus were classified as embryogenic (yellow and friable), organogenic (whitish) or unable to generate plants (watery and translucent).

A greater proportion of TYPE 1 cell masses were generated in the culture medium with the lowest dose of 2,4-D (1 mg L⁻¹) (Figure 1a). In contrast, culture media containing the picloran had a low proportion of this type of cell masses (Figure 3a). Type 2 cell masses were only generated on the treatment contained growth regulator 2,4-D at a concentration of 3 mg.L⁻¹ (Figure 3b), while TYPE 3 cell masses were observed more frequently on the 1,6 and 9 mg L⁻¹ picloran treatments (Figure 3c). Type 4 cell masses were obtained on the 3 mg.L⁻¹ treatment of 2,4-D and on the 1 mg.L⁻¹ and 9 mg.L⁻¹ picloran treatments (Figure 3d).

Callus external characteristics may be species-specific, hindering standardization of external morphology criteria to identify embryogenic potential. Such potential can, however, be potentially identified on the basis of cytological features. Histochemical analysis of TYPE 1 cell masses indicated the presence of dispersed elongated cells (Figure 4), without the deposition of starch grains (Figure 4b). TYPE 2 cell masses consisted of small isodiametric cells (Figure 4c) with clear accumulation of starch grains which were very apparent with lugol (Figure 4d). TYPE 3 cell masses were also characterized by small isodiametric cells and accumulation of starch grains (Figure 4e). These masses also contained globular structures with differentiation sites representing meristematic centers.
with initial formation of procambium. Some cells were observed shedding around these globular structures (Figure 4f). Type 4 cell masses also contained globular structures formed by small isodiametric cells, contain starch grains and cells shedding around the globular structures (Figure 4g) and other cells (Figure 4h).

Different cell mass formations have been observed in the callus of Araucaria angustifolia (Steiner et al. 2005). In this case, some cell masses contained small isodiametric cells which formed embryogenic clusters from which subsequently originated proembryos. Other masses had elongate and vacuolated cells similar to the TYPE 1 masses observed in the present study. These masses did not produce embryos.

Cell masses belonging to TYPES 2, 3 and 4 were characterized by the deposition of starch grains (Figure 4d, 4f and 4h). Starch storage in the embryo, embryogenic cells or in adjacent cells usually indicates the acquisition of embryogenic competence (Moura 2008) as starch grains being produced to support and initiate the development of somatic embryos. Large amounts of starch have been observed in embryogenic callus cells of Gentiana punctata, as source of energy for intense cell division and for development of the embryo (Mikula et al. 2004). On the peach palm (Bactris gasipaes) the accumulation of starch has been observed to precede the development of somatic embryos (Steinmacher et al. 2011).

Figure 2 - Stereomicrographs of callus produced on oil palm tenera hybrid (Elaeis guineensis) leaves. A) Type 1 - elongated and translucent; B) Type 2 - uneven and translucent; C) Type 3 - globular and beige; D) Type 4 - globular and white. Bar = 0.25 mm.
Similar observations were made in embryogenic cells of *Feijoa sellowiana*, which are characterized by a well-developed nucleus with prominent nucleoli and starch grains (Canhoto et. al 1996). Once again, the starch grains are thought to provide energy for the development of the somatic embryos, suggesting active regulation of starch grain accumulation in embryogenic cells (Martin et al. 2000).

The ultrastructural analysis indicated that TYPE 1 cell masses had large vacuoles (Figure 5a), diffuse cytoplasm (Figure 5b) and intercellular spaces (Figure 5c) within elongated cells (Figure 1a). Such elongated cells are not embryogenic and the vacuolation is considered as an early marker of cell death (Filonova et al. 2000, Lam et al. 2000). Based on these characteristics, Type 1 cell masses probably have no embryogenic potential.

TYPE 2 cells were characterized by a nucleus with prominent nucleoli, dense cytoplasm, high nucleus/cytoplasm ratio, the presence of amiloplasts close to the nucleus (Figure 6), thick walls with little intercellular space, mitochondria and numerous plasmodesmata (Figure 6). TYPE 3 cells had a nucleus with nucleoli apparent, many amiloplasts, mitochondria, endoplasmic reticulum, phenols, lipids, and thin cell walls (Figure 8). It was also possible to observed thick cell walls with few intercellular spaces in TYPE 4 cells, which also contained many mitochondria and amiloplasts (Figure 9).

Callus cells of the coconut palm (*Cocos nucifera*) have a similar ultrastructure to TYPE 2 cell masses, being characterized by large nuclei with two nucleoli, cell walls with uniform thickness, amiloplasts and plasmodesmata (Verdeil et al. 2001). Plasmodesmata are channels that connect neighboring cells allowing the exchange of structural and functional molecules at a faster rate than transport through membranes (Concenço et al. 2007).

![Figure 3 - Callus type percentage of oil palm tenera hybrid (*Elaeis guineensis*) induced on different 2,4D and picloran. A) Type 1, B) Type 2, C) Type 3, and D) Type 4.](image-url)
Figure 4 - Histochemical analysis of oil palm tenera hybrid (Elaeis guineensis) callus leaves induced with 2,4D and picloran
A) Callus Type 1 stained with toluidine blue. Elongated and dispersed cells (arrows); B) Callus Type 1 stained with Lugol;
C) Callus Type 2. Cells strongly stained with toluidine blue (arrows); D) Callus Type 2 stained with Lugol. Starch grains (arrows);
E) Callus Type 3 stained with toluidine blue. Globular structure with procambium (Pc); F) Callus Type 3 with starch grains stained with lugol (arrows). G) Callus Type 4 stained with toluidine blue. Isodiametric small cell H) Callus Type 4. Starch grains stained with lugol. Procambium.
Figure 5 - Transmission electron micrographs of the callus Type 1 A) Large vacuole, narrow cytoplasm and cell wall rupture (arrows). B) Cells. C) Large intercellular spaces (arrows), cell wall (CW) and large vacuole (Va).
Embryogenic cells undergo changes when acquire the competence to form pro-embryos as: cell wall thickening, surrounding tissues senescence, increase of amiloplasts around the nucleus, no spherical nucleus and consequently numerous mitotic divisions. After of the pro-embryos formation the cells return to meristematic cells features with spherical nucleus, less amiloplasts around the nucleus, thin cell wall and formations of plasmodesmata (Verdeil et al. 2001).

Date-palm (*Phoenix dactylifera*) somatic embryos are characterized by small cells with meristematic characteristics, dense nuclei and cytoplasm (Aslam et al. 2011), and are therefore histologically similar to the globular structures observed in TYPE 3 cell masses. These embryogenic characteristics (large nuclei with prominent nucleoli, mitochondria, and starch grains) have also observed in the callus of Inga (Stein et al. 2010). In *Feijoa sellowiana*, cells with embryogenic characteristics had well-developed nuclei with prominent nucleoli, many mitochondria and starch granules (Canhoto et al. 1996). The presence of numerous mitochondria is related to high cell metabolism because this organelle is responsible for cellular respiration.
Figure 8 - Transmission electron micrographs of the callus Type 3. A) Large nucleus, nucleolus and thick cell wall. B) Endoplasmic reticulum (arrows). C) Numerous amyloplasts (arrows). D) Mitochondria (arrows) E) and F) Phenols (arrows).
Figura 9 - Transmission electron micrographs of the callus Type 4. A) Thin cell wall. B) Numerous mitochondria (arrows) and C) amyloplasts (arrows). Cell Wall (CW).
CONCLUSIONS

The culture medium supplemented with the auxins 2,4-D and picloram successfully promoted the formation of four types of cell masses, regardless of the concentrations of growth regulator.

TYPE 1 cell masses had no embryogenic characteristics, being composed of elongated vacuolated cells and with degraded cell walls - indicating apoptosis.

The growth regulator picloram (at a concentration of 1 mg.L$^{-1}$) is most effective at inducing Type 3 and 4 pro-embryogenic masses.

Based on ultrastructural characteristics and on production of pro-embryogenic masses, TYPE 3 cell masses are most suitable for regeneration of oil palm through somatic embryogenesis.

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REFERENCES


RESUMO

O dendezeiro é uma das oleogínas mais importante economicamente, mas a expansão das plantações tem sido limitada pela disponibilidade de mudas, como a propagação convencional é através de sementes, que têm baixas taxas de germinação. Uma possível solução para produção em larga escala é por meio da embriogênese somática. O objetivo deste trabalho foi avaliar os efeitos das auxinas 2,4-D e picloram na indução de massas pró-embriogênicas em explantes foliares de Elaeis guineensis híbrido Tenera e caracterizá-las, considerando as características embriogênicas, com análises citoquímicas e ultraestruturais. Para indução das massas pró-embriogênicas, fragmentos foliares de plântulas in vitro foram inoculados em meio de cultivo Y3 suplementado com 2,4-D ou picloram em diferentes concentrações (0,0; 1,0; 3,0; 6,0 e 9,0 mg.L$^{-1}$). Após 90 dias foram avaliadas presença e ausência das massas celulares. Ambos reguladores de crescimento foram eficientes na indução destas massas, independentemente, das concentrações utilizadas. Como as massas celulares não eram homogêneas, estas foram classificadas quanto à cor e ao formato em quatro tipos: TIPO 1 – translúcido e alongado, TIPO 2 – translúcido e aquoso, TIPO 3 – bege e globular e TIPO 4 - branco e globular. De acordo com as características anatômicas e ultraestruturais os as massas celulares dos TIPOS 2, 3 e 4, foram consideradas com maior potencial embriogênico e portanto podem ser as mais promissoras para a propagação vegetativa em larga escala para o dendezeiro.

Palavras-chave: potencial embriogênico, reguladores de crescimento, cultura de tecidos, microscopia eletrônica de transmissão.


