Induction and Cytological Studies of Coffea arabica cv. Catimor CIFC 7963 Callus

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ABSTRACT

This study investigated the effect of activated charcoal (AC) and 4amino-3,5,6-trichloro-2-pyridinecarboxylic acid (picloram) on callus induction and occurrence of mitotic aberrations in coffee cells. Explants from the first pair of expanded leaves (1st leaves) of Coffea arabica "Catimor CIFC 7963" were cultured on Murashige and Skoog (MS) medium supplemented with $1 \mu M$ 2,4-dichlorophenoxyacetic acid $(2,4-D) + 5 \mu M$ 6-benzlyaminopurine (BA). The treatments included the following rates of AC application: 0.008%, 0.08%, 0.1%, and 1% (w/v). We also evaluated the effects of growth in MS medium supplemented with 1 μ M 2,4-D + 5 μ M BA + 10 mM picloram vs. 10 mM picloram as the sole plant growth regulator. The results showed that cells of the 1^{st} leaves on MS medium + 1 μ M 2,4-D + 5 μ M BA + 0.008% AC were induced to callus (46.88% callus induction). In addition, callus induction (90.63% and 59.38%) was observed in the leaves cultured on MS medium + 1 μ M 2,4-D + 5 $\mu M BA + 10 mM$ picloram and on MS medium+10 mM picloram, respectively. However, the 4-month-old friable calli developed on MS medium containing picloram showed mitotic aberrations. Some of these included binucleated cells, a micronucleus in binucleated cells, and vagrant chromosome cells. These results highlight that both 0.008% AC and 10 mM picloram were able to induce the development of 4-month-old friable callus in a high proportion of C. arabica "Catimor CIFC 7963" cells, but while the former produced no mitotic aberrations, the latter induced mitotic aberrations in cells.

Keywords: *Coffea arabica*, Mitotic aberration, Friable callus, Picloram, Activated charcoal

INTRODUCTION

Coffea arabica "Catimor CIFC 7963" is a hybrid coffee plant, obtained by crossing C. arabica "Caturra rojo" with a hybrid of Timor (a natural cross of C. arabica and C. canephora) (Silva et al., 2005). It is a self-fertilizing species (tetraploid plant, 2n = 4x = 44) (Etienne and Bertrand, 2003). Propagation of coffee plants via vegetative organs provides true-to-type regenerated plants. In addition, micropropagation using somatic embryogenesis can facilitate largescale vegetative propagation of hybrid varieties in clonal form. However, during this process, the polyploid cells may undergo abnormal division and generate aneuploid cells in polyploid plants. Chromosome segregation errors can result from changes in the relative dosage of products of genes located on the mis-segregated chromosomes (Potapova and Gorbsky, 2017). Thus, successful mass production of embryogenic cells and somatic embryos (SEs) is contingent on the genetic stability of regenerated plants. Embryogenic coffee cells are frequently generated using older embryogenic cultures, and hence, the phenotypic variants due to aneuploidy can be passed on through mitotic aberrations in cells. It has been demonstrated that the chromosomal variability is directly linked to somaclonal variation (SV) in embryogenic coffee cells (Etienne et al., 2016). SV is described as the phenotypic variation observed during the clonal propagation of plants. It is a major cause of concern when strict clonal propagation of elite plants is required to ensure true-to-type in vitro propagation.

Arabica coffee hybrid plants should be studied for multiplication using somatic embryogenesis processes, as it can help achieve: (1) a reduction in production costs, and (2) a guarantee that the propagated plants are true-to-type (Etienne and Bertrand, 2003). In particular, the use of embryogenic callus has been frequently associated with an increase in genetic instability and SV in the regenerated plants (Jähne et al., 1991; Rival et al., 1997; Lu et al., 2006). The SV in embryogenic-cell-derived plants is probably related to the use of 2, 4-D for division of cells in an embryogenic undifferentiated state (Lambé et al., 1997; Von Aderkas and Bonga, 2000). In addition, picloram, which works as an auxin, can stimulate DNA and RNA synthesis along with the transcription of variable proteins in plants, resulting in uncontrolled cell division to induce plant callus formation (Ozel et al., 2018). However, the uncontrolled cell division is associated with incomplete replication of the genome, and can lead to chromosomal aberrations (Arora et al., 2013). The frequency of SV has been shown to increase exponentially with the age of the embryogenic cell culture (Etienne and Bertrand, 2003). The types of phenotypic variants observed in coffee plants have been characterized based on the color of juvenile leaves, giant, dwarf, and thick leaves (bullata), variegata, angustifolia, and multi-stems. These variants were rare (1.3%) for plants derived from embryogenic callus in 3-month-old embryogenic cell suspensions. However, the frequency of variants produced from 6-, 9-, and 12-month-old cell suspensions was 6%, 10%, and 25%, respectively (Etienne and Bertrand, 2003). Thus, the multiplication time for embryogenic cells culture was shortened to 6 months and the level of auxin was reduced (0-1.4 μ M 2, 4-D), as the frequency of variation is affected by these parameters.

Chromosomal variability can also be a source of SV. A cytological technique has been used to quantify the SV and genetic stability of plants derived from a tissue culture system (Raha and Roy, 2003). In addition, variation in plant tissue culture, particularly in callus cultures, can also be caused by mitotic aberrations (Larkin and Scowcroft, 1981). In this present study, the mitotic aberrations in leaf embryogenic callus cultures were investigated. Activated charcoal (AC) promotes embryogenesis, and improves the growth and morphogenic responses of explant. AC adsorbs many impurities, and modulates the culture medium pH by altering the mineral availability in the culture medium, in order to induce embryogenesis (Chee and Tricoli 1988; Van Winkle and Pullman 2003; Van Winkle et al., 2003; Thomas, 2008; Sáenz et al. 2010; Cheng et al., 2013). Thus, a selected leaf explant from C. arabica "Catimor CIFC 7963" was used to study the effects of treatment with various concentrations of AC and plant growth regulators for various durations of callus culture. In addition, we carried out a morphological characterization of the callus initiated from leaf sections, and studied the callus induction as well. The purpose of this study was to evaluate the optimal culture age of C. arabica "Catimor CIFC 7963", and the cytotoxicity of picloram on mitotic divisions of embryogenic callus, for true-totype propagation.

MATERIALS AND METHODS

Plant preparation and tissue culture

A sample of 32 explants of leaves from greenhouse-grown 2-year-old coffee trees (C. arabica "Catimor CIFC 7963") were used for callus induction. All explants from the basal part of the leaf were washed under running tap water for 5 min to remove dust particles. The explants were then disinfected for 10 min in a 15% (w/v) solution of commercial bleach (NaOCl) containing a few drops of Tween 80 (Merk Schuchardt), followed by three successive washes with sterile double distilled water to remove all traces of NaOCl. After surface sterilization, the margin and the midrib of each leaf were removed and the remaining leaf tissue was cut into pieces of size $1 \text{ cm} \times 1 \text{ cm}$. The sterilized explants were placed on MS medium (Murashige and Skoog, 1962), supplemented with picloram and acid-washed AC, for plant tissue culture (Phytotechnology Laboratories[®]), combined with or without 2,4-D and BA. Eight treatments were used to evaluate the effects of the plant growth regulators (2,4-D and BA (µM), picloram (mM), and AC (w/v)%) as follows: treatment A: 0 2,4-D, 0 BA, 0 picloram, 0 AC, treatment B: 1 2,4-D, 5 BA, treatment C: 1 2,4-D, 5 BA, 0.008 AC, treatment D: 1 2,4-D, 5 BA, 0.08 AC, treatment E: 1 2,4-D, 5 BA, 0.8 AC, treatment F:

1 2,4-D, 5 BA, 1 AC, treatment G: 1 2,4-D, 5 BA, 10 picloram, and treatment H: 0 2,4-D, 0 BA, 10 picloram. All media were adjusted to pH 5.7–5.8. The cultures were incubated in the dark at 25 ± 2 °C and subcultured at 4-week intervals on the same media for 4 months. Subsequently, the numbers of explants with calli were counted, and the percentage of explants with calli was calculated using the formula:

Percentage of explants with calli = $\frac{\text{Number of explants with calli}}{\text{Total number of explants}} \times 100$

After 4 months, the calli were examined and the fresh weight per explant was determined before investing the cytological behaviors.

Chromosome squash

After 2 and 4 months, the calli were subjected to cytological analysis. They were kept in Carnoy's solution (absolute alcohol:glacial acetic acid, 3:1) for 24 h and then stored in 70% ethyl alcohol at 4 °C until analysis. The cell walls of the calli were hydrolyzed in 1 N HCl for 15 min and then stained in 1% aceto-orcein for 10 min before the tissue was squashed under a coverslip. The mitotic stages in the calli cells were observed under a light microscope (Olympus Optical Co., Ltd) using a 100x objective lens in immersion oil. Selected representative stages of cells were photographed. The normal and abnormal cells were then analyzed. The mitotic index (MI) was calculated using the formula:

Mitotic index (MI) = $\frac{\text{Number of dividing cells}}{\text{Total number of cells scored}} \times 100$

Statistical analysis

Data were analyzed using one-way ANOVA, and significance differences among treatment means were determined using Duncan's multiple range test at P < 0.05. The statistical package for SPSS version 14.0 software was used (Freund and Wilson, 2003).

RESULTS

Effects of AC on cell proliferation and callus induction in *C. arabica* "Catimor CIFC 7963" leaf explants

In treatments B and C, calli were obtained from leaf explants after 1 month, while in treatments A (control), D, and E, groups of small cells were obtained. While cell division was induced in all leaf sections of these three treatments, the fresh weight could not be examined due to insufficient proliferation. Our results showed that none of the groups of small cells (treatments A, D, and E) showed

continuous development to callus after 2 months. Furthermore, in treatment F, some explants showed no response and exhibited necrosis after 2 months. This result revealed that increasing the concentration of AC beyond 0.08% (w/v), to 0.01% or 1%, resulted in decreased cell proliferation into callus formation among the surviving explants. The highest percentage of callus induction (100%) on medium was obtained for explants (0.20 ± 0.07 g/explant) cultured on MS medium supplemented with 1 μ M 2,4-D + 5 μ M BA (treatment B). Moreover, the addition of 0.008% AC resulted in no significant difference in the fresh weight of 4-month-old calli (0.18 ± 0.07 g/explant), but did induce friable calli in all the explants. We found that both compact and friable calli were obtained after 2–4 months on MS medium supplemented with 2,4-D + BA (treatment B). Interestingly, the MS medium supplemented with 2,4-D + BA containing 0.008% AC induced only friable calli from explants (Table 1). This result indicated that AC had a significant effect on the development of friable calli.

Table 1.	Effects of AC o	n cell proliferation	and callus indu	uction in C.	arabica
	"Catimor CIFC	7963" leaf explants	5.		

MS medium	Supple	ement	AC (%w/v)	Percentage of callus induction	Callus (4 months old)		
	2,4-D	BA		(%)	Fresh weight	Callus type	
	(µM)	(µM)			(g/explant)		
A	-	-	-	56.25 ^b	_*	group of small cells	
В	1	5	-	100 ^a	$0.20\pm0.07^{\rm a}$	compact callus/	
						friable callus	
С	1	5	0.008	46.88 ^b	$0.18\pm0.07^{\rm a}$	friable callus	
D	1	5	0.08	15.63°	_*	group of small cells	
Е	1	5	0.1	6.25 ^{cd}	_*	group of small cells	
F	1	5	1	0^d	_*	-	

Note: * No callus, data collected after 4 months for fresh weight callus/explant. ^{a,b,c,cd,d} represent differences among treatment means within the same column (P < 0.05).

Effects of picloram with or without 2, 4-D and BA on callus induction in *C. arabica* "Catimor CIFC 7963" leaf explants

Calli from leaf explants were obtained in treatments B, G, and H. Moreover, the MS medium supplemented with 10 mM picloram with or without $1 \mu M 2$, $4-D + 5 \mu M BA$ (treatments G and H, respectively) had a significant effect on the development of friable callus. The highest fresh weight of 4-monthold callus (0.32 ± 0.06 g/explant) was obtained on MS medium supplemented with 1 μ M 2,4-D + 5 μ M BA containing 10 mM picloram (treatment G). Moreover, the addition of 10 mM picloram in MS medium supplemented with $1 \mu M 2$, $4-D + 5 \mu M BA$ resulted in no significant effect on callus induction (90.63%), but picloram did show a significant effect on the fresh weight of callus. The fresh weight of callus obtained with the combination of 2, 4-D + BA was not significantly different from that obtained with picloram as the sole plant growth regulator, indicating that the effect of picloram as auxin was similar to that of 2, 4-D in callus induction. Our results showed that the picloram also affected the cell division, without any cytokinin application, in treatment H. However, cells of the leaf explants developed into friable calli, resulting in reduced callus induction (59.38%) and reduced fresh weight of callus (0.13 ± 0.05 g/explant) (Table 2). The calli from treatments B, C, G, and H were fixed in Carnoy's solution to investigate their cytological behavior.

MS	Supplement		Picloram Percentage		Callus (4 months old)		
medium	2,4-D (µM)	ΒΑ (μΜ)	(mM) (%)	of callus induction (%)	Fresh weight (g/explant)	Callus type	
A	_	_	_	56.25 ^b	_*	group of small	
						cells	
В	1	5	-	100 ^a	$0.20\pm0.07^{\rm b}$	compact callus/	
						friable callus	
G	1	5	10	90.63 ^a	$0.32\pm0.06^{\rm a}$	friable callus	
Н	-	-	10	59.38 ^b	$0.13\pm0.05^{\text{b}}$	friable callus	

Table 2. Effects of picloram with or without 2, 4-D and BA on callus inductionin *C. arabica* "Catimor CIFC 7963" leaf explants.

Note: * No callus, data collected after 4 months for fresh weight callus/explant. ^{a,b} represent differences among treatment means within the same column (P < 0.05).

Callus characterization of *C. arabica* "Catimor CIFC 7963" (2 and 4 months old)

All calli were cultured on media for 2–4 months. At the end of the second month, the green calli presented a white yellowish color in treatments B and C (Figure 1A and 1B). Thereafter, up to 4 months, the calli presented a black brownish color (Figure 1E and 1F). Asynchronous development of SEs was observed under the stereo microscope. Different stages of SEs, including globular embryos and elongated embryos, were clearly visible within 2 months in treatments B and C (Figure 1A and 1B). Treatment B and C not only produced SEs, but also produced roots via indirect organogenesis within 2 months (Figure 1A-B and 1E-F). Treatments G and H showed no established SEs. Our results revealed that AC and picloram produced more developed and larger sized friable calli, respectively. Moreover, the white friable calli of treatments G and H showed no color change even after 4 months (Figure 1C-D and 1G-H).



Figure 1. Callus of *C. arabica* "Catimor CIFC 7963" on media at 2 and 4 months. (A, E) MS medium + 1 μ M 2, 4-D + 5 μ M BA, (g) globular embryo (e) elongate embryo (r) root. (B, F) MS medium + 1 μ M 2, 4-D + 5 μ M BA + 0.008% AC (w/v). (C, G) MS medium + 1 μ M 2, 4-D + 5 μ M BA + 10 mM picloram. (D, H) MS medium + 10 mM picloram.

Cytological study of calli cells (2 and 4 months old)

Mitotic division was observed every 2 months for the calli cells cultured on the four media (treatments B, C, G, and H). Cytological analysis showed 1,829 actively dividing cells (2 months old) at various stages of mitosis (interphase, prophase, metaphase, anaphase, and telophase), as shown in Figure 2A-E. The calli contained several types of cells, including those both round and elongated ones. Moreover, cytokinesis was observed, where the cell divided to form two daughter cells. The dividing structure, known as the cell plate, formed in the center of the cytoplasm, and a new cell wall developed between the two daughter cells (Figure 2F). Our results showed that 10 mM picloram in combination with

 $1 \mu M 2,4-D + 5 \mu m BA$ (treatment G) induced the highest fresh weight of callus/explant (0.32 ± 0.06 g/explant) (Table 2). This phenomenon caused mitotic aberrations in the 4-month-old calli. The results showed that almost 2,337 calli cells (4 months old) had normally dividing nuclei, with treatments B and C showing no abnormal calli cells. In contrast, binucleated cells were observed in MS medium containing 1 μ M 2, 4-D + 5 μ m BA + 10 mM picloram (treatment G). In binucleated cells, the nuclei were typically of the same size, owing to the inhibition of cell plate formation (Figure 2G). In addition, micronuclei in binucleated cells and vagrant chromosome cells, as a consequence of spindle poisoning, were observed in MS medium containing only 10 mM picloram (treatment H) (Figure 2 G-I). The percentage of these abnormal cells was low (1%, data not shown). The natural consequence of such abnormality was genetic mosaicism of the callus. The cells observed develop into polyploid and aneuploid plants. This result suggested that 10 mM picloram had significant effects on mitotic aberration in the cells of 4-month-old calli, and a higher number of polyploid and aneuploid cells were seen with increasing age of the culture.



Figure 2. Normal and abnormal calli cells. (A) Interphase (100x), (B) prophase (100x), (C) metaphase (100x), (D) anaphase (100x), (E) telophase (100x), (F) cytokinesis (100x), (G) binucleated cell (100x), (H) vagrant chromosome cell (100x), and (I) micronucleus in binucleated cell (100x).

MI study of calli cells (2 and 4 months old)

As for the distribution of cells across the mitotic phases, the highest number of cells was seen in prophase. The number of calli cells in metaphase, anaphase, and telophase decreased on all media (Table 3). Cytological analysis revealed normal mitotic activity in 2-month-old calli cells (Figure 2A-E), whereas the 4-month-old calli cells in treatments G and H showed mitotic aberrations (Figure 2G-H), with a rather low MI in treatment C (8.46) but a higher MI value in treatment B (4.62) (Table 3). The MI was lower in 4-month-old calli cells compared with the 2-month-old calli cells in treatments G (4.90 and 4.96, respectively) and H (4.74 and 4.82, respectively). This result revealed that aging of the cell culture on medium containing 10 mM picloram was mutagenic in coffee, as there was a reduction in the MI of the 4-month-old calli cells. In contrast, treatments B and C resulted in an MI increase in the 4-month-old calli cells compared with the 2-month-old calli cells (4.62 and 2.66, respectively) and (8.46 and 4.41, respectively) (Table 3), and they showed normal mitotic division (Figure 2A-E). Moreover, we found that the presence of auxin at a low level (1 μ M 2, 4-D) in treatments B and C resulted in organized, normal growth phases in the callus, with no mitotic aberrations.

Table 3. MI and number of calli cells (2 and 4 months old) undergoing mitoticdivision in *C. arabica* "Catimor CIFC 7963".

	of xed	No. of cells in mitotic phases	IM	No. of cells by mitotic phases			
dium	No. c analyz cells			Prophase	Metaphase	Anaphase	Telophase
S me	2/4	2/4 months	2/4	2/42/4monthsmonths oldmonths	2/4	2/4	2/4
N	months old		months		months old	months old months old	
		old	old		old		
В	489/606	13/28	2.66/4.62	5/25	3/2	2/1	3/0
С	340/544	15/46	4.41/8.46	7/45	4/1	3/0	1/0
G	524/469	26/23	4.96/4.90	18/18	3/2	2/1	3/2
Н	539/718	26/34	4.82/4.74	20/30	2/1	2/2	2/1

DISCUSSION

Effects of AC on cell proliferation and callus induction in *C. arabica* "Catimor CIFC 7963" leaf explants

The callus induction in the treatment with 1 μ M 2,4-D + 5 μ M BA + 0.008% AC (w/v) did not differ significantly from the one with 1 μ M 2,4-D + 5 μ M BA treatment (without the AC). Interestingly, AC at 0.008% had a high rate of induction of only friable calli from explants, in contrast with the medium

containing 2,4-D + BA, which produced both compact and friable calli (Table 1). These results support the finding from a previous study that AC at a concentration of 0.05% (w/v) can act as a promoter of embryogenesis in different genotypes of pepper (Cheng et al., 2013). Our results showed that AC at a concentration of 0.008% induced cell proliferation into friable callus, which is a characterization of embryogenic callus (Table 1). Thereafter, up to 2 months, some friable calli developed into somatic embryos or roots (Figure 1B). Kaouther et al., (2011) reported that AC at 0.3% was effective in controlling oxidation and improving the percentage of callus induction in palms. Therefore, owing to the strong adsorptive properties exhibited by AC in tissue culture media, it could remove the growth-inhibitory substances produced by plant tissue (Cheng et al., 2013). Moreover, AC adsorbed not only the plant growth-inhibitory substances in the medium, but also the plant growth regulators and nutrients (Liu, 1993). Thus, the increasing of AC effects on the survival explant. This could also explain why the treatment of 2,4-D + BA with a concentration of AC beyond 0.08% (w/v), to 0.01% and 1%, showed a decreasing trend of cell proliferation into callus formation in surviving explants (Table 1).

Effects of picloram with or without 2,4-D and BA on callus induction in *C. arabica* "Catimor CIFC 7963" leaf explants

The highest fresh weight of 4-month-old callus $(0.32 \pm 0.06 \text{ g/explant})$ was obtained for MS medium supplemented with 1 μ M 2,4-D + 5 μ M BA, containing 10 mM picloram (treatment G). In addition, treatment H showed that 10 mM picloram was the sole plant growth regulator, which affected cell division without the presence of cytokinin hormones (Table 2). This result was supported by a previous report that 4 µM picloram as the sole plant growth regulator induces callus formation in C. arabica "Rubi", "Catuaí vermelho 81", and "IAPAR 59" (Cid et al., 2004). According to Ozel et al., (2018), a high concentration of picloram affects the root cells of Allium cepa, which exhibited a decreased MI and mitotic abnormalities such as anaphase bridges, C-mitosis, multipolarity, fragments, and lagging chromosomes and polyploidy. We found that addition of 10 mM picloram resulted in high rate of callus formation in C. arbica "Catimor CIFC 7963" and relatively unaffected calli cells, with the frequency of mitotic abnormality limited to around 1%. Picloram produced a larger size of friable calli compared with the MS medium supplemented with 1 μ M 2, 4-D + 5 μ M BA (Figure 1A and 1C). This result revealed that picloram served as a superior auxin than 2, 4-D hormone. This could be due the more effective uptake and rapid mobilization of picloram at the target sites (Karun et al., 2004). The superior effect of picloram for embryogenic callus induction has been reported in many plants such as palm (Elaeis guineensis Jacq.), cherry (Prunus incisa Thunb.), kodo millet (Paspalum scrobiculatum L.), and C. arabica (Cid et al., 2004; Kaur and Kothari, 2004; Kaouther et al., 2011; Jayanthi et al., 2015).

Cytological and MI analysis of calli cells (2 and 4 months old)

The highest proportion of cells was seen in prophase, whereas the number of calli cells in metaphase, anaphase, and telophase was relatively low across all media (Table 3). Furthermore, we found that the rate of division of 4-month-old calli cells decreased at prophase in treatments G and H compared with treatments B and C. Picloram treatments decreased the rate of prophase activity, thereby diminishing the MI. The reduction in MI could have been due to blocking of DNA synthesis at the G1 or S phase (Schneiderman et al., 1971; Sudhakar et al., 2001) or blocking of the G2 phase (Van't, 1968). Our results showed that the reduction in the MI was clearly related to long-term culture, as well as the cytotoxic effect of picloram and 2,4-D, resulting in three types of structural aberrations: binucleated cells, micronuclei in binucleated cells, and vagrant chromosome cells (Figuure 2G-I).

The mitotic aberration frequency was 1% (data not shown). According to Landey et al. (2013), the variant frequency increased in embryogenic suspensions of C. arabica after 6 months of culture in a medium containing 4.52 mM 2, 4-D, while after 12 months, the frequency reached 25%. Thereafter, they observed SV in 11- and 27-month-old embryogenic callus cultures, which affected the aneuploidy regenerated plants. This result suggested that cell culture ageing is highly mutagenic in coffee and chromosomal rearrangements are directly linked to SV (Landey et al., 2015). Our results showed that 10 mM picloram induced a Figure cell division and rapid rate of (Table C-D). 1 1 a consequence, there was a disorganized rapid growth phase in the callus, which contributed to the SV (Karp, 1994; Rani and Raina, 2000). This result was supported by the finding of a previous report that 1 mg/L picloram shows a significantly higher induction rate than 0.1 mg/L IAA and 0.1 mg/L IBA, but the SEs obtained show teratological abnormalities such as fused or polycotyledonary embryos (Kaouther et al., 2011). We demonstrated that the effect of culture age was associated with prolonged exposure to 10 mM picloram. Many altered mechanisms, observed during the mitosis of calli cells of C. arabica, can produce chromosome alterations such as polyploidy, aneuploidy, sticky chromosomes, double prophases, and lagging chromosomes (Menéndez-Yuffá et al., 2000). Our results showed that cells of C. arabica "Catimor CIFC 7963" with a micronucleus or binucleated cells and vagrant chromosome cells would be differentiated into polyploid and aneuploid plants, respectively.

SV detection has used to verify ploidy instability in embryogenic cells of *C. arabica* in vitro using flow cytometry (FCM) (Clarindo et al., 2012). This report revealed that FCM was able to detect the non-true-to-type calli cells (4 months old), and highlighted the octaploid or aneuploid cells. Therefore, these authors recommended a culture age limit of 4 months for true-to-type mass propagation of *C. arabica* cell suspensions (Clarindo et al., 2012). Similarly, our results indicated that the 4-month-old calli cells of *C. arabica* "Catimor CIFC 7963" exposed to 10 mM picloram showed mitotic aberrations. However, to the

best of our knowledge, the 4-month-old friable calli cultured on MS medium supplemented with 1 μ M 2,4-D + 5 μ M BA + 0.008% AC did not have cytotoxic effect on cells. In contrast, treatment with 1 μ M 2, 4-D + 5 μ M BA + 10 mM picloram affected 4-month-old calli cells, and resulted in a mitotic aberration frequency of 1% compared with the 3-month-old embryogenic cell suspension of *C. arabica*, which showed a frequency of 1.3% (Etienne and Bertrand, 2003). Moreover, our culture medium resulted in a lower frequency of SV (1%) than the embryogenic calli of *C. arabica* "Catimor CIFC 7963" (25%) cultured on ½MS supplemented with 1 mg/L 2,4-D + 8 mg/L BA, followed by transfer to the same medium containing 0.8 mg/L NAA for 3 months (Menéndez-Yuffá et al., 2000).

CONCLUSION

The results of this study showed that leaf explants of C. arabica "Catimor CIFC 7963" could be induced to form compact and friable calli on MS medium supplemented with 1 μ M 2,4-D + 5 μ M BA. Furthermore, friable calli were produced on MS media supplemented with 1 μ M 2,4-D + 5 μ M BA + 0.008% AC (w/v), 1 μ M 2,4-D + 5 μ M BA + 10 mM picloram, or 10 mM picloram. Our results revealed that AC at a concentration of 0.008% induced cell proliferation into friable callus at a slower rate than 10 mM picloram. However, cytotoxic effects of 10 mM picloram in this experiment caused mitotic aberrations in the long-term culture (4 months old). In contrast, 0.008% AC did not show any cytotoxic potential in cultured calli during the 4 months. Based on these results, we recommended an upper limit of 4 months for true-to-type mass propagation of C. arabica "Catimor CIFC 7963" friable calli with 10 mM picloram. The friable calli should be induced into SEs within 2-3 months, and the obtained seedlings would show a decreased SV. Given that SV was observed in the cytological study, it is advisable to conduct further analyses of the SEs using other methods, such as DNA molecular markers or chromosome count methods. These will confirm the genetic stability of the coffee plant SEs.

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