

Development of *In vitro* Tetraploid plants of *Hevea brasiliensis*

Abstract

Increase in global consumption of natural rubber necessitates crop improvement of *Hevea* aimed at increased productivity. As conventional breeding of *Hevea* is very elaborate and time consuming. Hence in the present study development of tetraploids through chromosome doubling of diploid callus obtained from cultured immature inflorescence of *Hevea* using colchicines were attempted. Chromosome doubling of the diploid callus occurred when treated with 1.25 μ M colchicine for 3 days. In higher concentrations as well as at longer exposure periods, the callus texture and viability were affected. 48 % embryo induction and a maturation frequency of 45 % were obtained. Embryo germination and plant regeneration with a germination frequency (30 %) and a regeneration frequency (20 %) were obtained. Cytological and flow cytometric analyses confirmed the tetraploid nature of the colchicines treated callus. *In vitro* tetraploid plant developed through these *in vitro* techniques can be further used in *Hevea brasiliensis* breeding.

Keywords: *Hevea brasiliensis*, Colchicine, *In vitro*, Tetraploids

Introduction

Tetraploids are polyploids with four sets of chromosomes per cell. In nature, a large number of angiosperm species are available with one or more episodes of polyploids which often results in good quality, high yielding plants with increased resistance to environmental stress, pests and diseases. Differential responses are observed in morphological and physiological characters of species due to tetraploidy. The cells of a tetraploid are much larger than that of diploid, as tetraploids have twice the number of sets of chromosomes per cell. Greater the number of chromosomes per cell, greater is the proportion of cell contents relative to cell wall material. It usually exhibits increased biomass mainly due to their high photosynthetic potential compared to diploids. In *Hevea brasiliensis* the economic life starts after 6-7 years of planting. Any attempt to reduce the immature (juvenile) phase would be quite rewarding. Even a reduction in the immature phase by 6 month or 1 year would enable the farmer to harvest the crops earlier and help to fetch the farmer with a reasonable income. *In-vitro* approaches to increase the vigour and biomass will naturally lead to shortening of the immature phase thereby enabling early tapping. The yield per tree per tap increases with increase in girth of trees due to increase in length of tapping cut. Polyploids either arise spontaneously or produced artificially. Artificial induction plays an important role in polyploid breeding. Manmade synthetic polyploids from wild plants have contributed to improvement of cotton, wheat and peanut [1]. With the advent of *in vitro* techniques for chromosome doubling using antimitotic agents, polyploids have been produced in a large number of species. *In vitro* induction of tetraploids through colchicine treatment has been achieved in many plants such as *Pyrus communis* [2], *Morus alba* [3], *Cinchona ledgeriana* [4] etc. Colchicine ($C_{22}H_{25}NO_5$), which is an alkaloid contained in seeds and bulbs of *Colchicum autumnale* L, has affinity for tubulin, a microtubule-sub unit protein, and inhibits spindle function thereby preventing both cell and nuclear division. *In vitro* induction

45 of polyploids by treating the diploids with colchicine has been successful in many plants like
46 oil palm, sesame and ginger [4].

47 Tetraploid/polyploid plants of *Hevea*, once generated through this technique, will be
48 having greater vigor and increased biomass which can lead to a reduction in the immaturity
49 period. Also, development of such plants with increased biomass may result in high girthing
50 trees which can be employed for cultivation as latex-timber clones. Hence the objective of the
51 study was to develop polyploid callus through colchicine application and regeneration of
52 polyploid plant by confirmation done using flow cytometer and cytological analysis.

53 **Materials and Methods**

54 **Plant material**

55 In this study, embryogenic callus derived from immature inflorescence, which is
56 diploid in nature, was used as the target material for colchicine treatment.

57 Embryogenic callus from immature inflorescence was raised using the earlier developed
58 protocol [5]. The immature inflorescence were washed thoroughly in running tap water for 10
59 min and surface sterilized with 0.1% (w/v) mercuric chloride solution containing two drops of
60 Tween-20 for three minutes followed by rinsing 3 times with sterile distilled water. These
61 explants were cut into small pieces and cultured for callus induction on MS basal medium
62 supplemented with growth regulators 2, 4-D (4.5 μM), NAA (2.7 μM) and Kinetin (2.3 μM).
63 The calli induced were cultured over modified MS medium supplemented with Kinetin (4.6
64 μM), BA (0.44 μM) and GA₃ (1.4 μM) for embryogenic callus induction. After 3-4
65 subcultures, cultures were transferred to same medium with high phytigel and charcoal.
66 Embryogenic callus emerged in this medium was used for chromosome doubling through
67 colchicine treatment (Fig. 1).



68
69 **Fig. 1 Embryogenic calli derived from immature Inflorescence**

70

71 **Colchicine treatment**

72 Colchicine treatment of the embryogenic callus was carried out using two different methods.

73 *Direct exposure to colchicine*

74 The embryogenic callus was suspended in different concentrations (0.25 - 2.5 μM) of
75 filter sterilized colchicine solution and incubated for different time intervals (2 - 24 h) with

76 continuous shaking at 1000 rpm, after which they were transferred to callus proliferation
77 medium.

78 *Different levels of colchicine (0.75 - 7.5 μ M) were incorporated in the callus proliferation*
79 *medium and the embryogenic callus was cultured in these media for different time intervals*
80 *(2-10 days) followed by transfer to proliferation medium without colchicine.*

81 Filter sterilized solution of colchicine was added to the **callus induction medium** just
82 before solidification, mixed well and poured into petri dishes and allowed to solidify. A stock
83 solution of 1000 ppm colchicine (Plant Cell Culture Grade-Sigma) was prepared in distilled
84 water and kept in amber coloured bottle, since the solution is light sensitive.

85 **Callus proliferation**

86 Modified MS medium supplemented with 2, 4-D (4.5 μ M), Kin (0.9 μ M), BA (0.8 μ M)
87 and GA₃ (0.14 μ M) was used as the callus proliferation medium. After colchicine treatment, the
88 calli were transferred to the proliferation medium. Sub culturing to the same medium was carried
89 out at one month interval for callus proliferation. Cytological analysis was carried out using this
90 callus to confirm the ploidy.

91 **Embryo induction**

92 The proliferated calli were transferred to different embryo induction media consisting
93 of three basal media namely Nitsch, MS and WPM fortified with different levels of BA (0.9 -
94 4.6 μ M) and GA₃ (0.57 - 2.9 μ M). Observations on embryo induction were recorded after 2-3
95 months in this medium and percentage of embryo induction was calculated.

96 **Embryo maturation**

97 The developing embryos were transferred to embryo maturation media consisting of
98 modified MS and WPM supplemented with different levels of kinetin (2.3 – 9.3 μ M) and
99 ABA (1.1 – 3.8 μ M). Basal medium without any growth regulators was also tried. Effect of
100 phytigel on embryo maturation was evaluated by solidifying with different levels of phytigel
101 (0.2 - 1.0 %).

102 Promotive effect of three amino acids viz. L- glutamine, L- asparagine and L- alanine
103 on embryo maturation was evaluated by adding different concentrations (5, 10, 15, 20 mM)
104 and combinations of these amino acids into the maturation medium. Four different
105 combinations were tried as follows-

- 106 (1) L glutamine (5mM) + L asparagine (5mM) + L alanine (5mM)
- 107 (2) L glutamine (10mM) + L asparagine (15mM) + L alanine (15mM)
- 108 (3) L glutamine (15mM) + L asparagine (10mM) + L alanine (10mM)
- 109 (4) L glutamine (20mM) + L asparagine (20mM) + L alanine (20mM)

110 **Embryo germination and plant regeneration**

111 Two different basal media, modified MS and WPM were used for germination and
112 plant regeneration experiments. The growth regulators experimented for germination was BA
113 (2.2 - 8.8 μ M) and IBA (2.5 - 9.9 μ M). For plant regeneration, the germinated embryos were
114 transferred to media fortified with various levels of IAA (1.7 -5.7 μ M), BA (6.6- 13.3 μ M)
115 and GA₃ (1.4 μ M) along with organic supplements like coconut milk (10 % v/v) and banana
116 powder (500 mg/l).

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118

119 **Statistical analysis**

120 All experiments were conducted in completely randomized design (CRD) and
121 analyzed using SPSS 16.0 software. The data was subjected to square root and arc sine
122 transformation and analyzed using ANOVA with a significance of $p \leq 0.05$.

123 **Confirmation of ploidy**

124 *Cytological analysis*

125 Proliferated calli from various colchicine treatments was subjected to cytological
126 analysis [6]. After plant regeneration, developing root tips of the regenerated plants were also
127 subjected to cytology in order to reassure ploidy of the regenerants. Callus with actively
128 dividing cells were pre-treated with 0.2 mM 8-hydroxyquinoline for 5 h at 4°C. After this
129 pretreatment the solution was drained off, the callus was washed with distilled water and
130 transferred to cold freshly prepared fixative, Carnoys fluid II (3:1 ethanol acetic acid), for 48
131 h at room temperature. Afterwards the fixative was drained off and the callus was washed
132 thoroughly to remove traces of fixative, if any. Then the samples were stained with 1 %
133 Snows carmine for 4 hrs. The samples were smeared in 45% acetic acid with a glass rod and
134 mounted on slides as per standard protocol. The slides were observed under a light trinocular
135 microscope (Leica).

136 *Flow cytometric analysis*

137 The callus which showed good embryogenic response was subjected to flow
138 cytometry analysis. For sample preparation, the callus was crushed in galbriath's* buffer and
139 kept for 5 min incubation. The suspension containing the nuclei was mixed by pipetting up
140 and down several times and then filtered through a 50 µm nylon mesh. The filtrate containing
141 the nuclear suspension was stained with 50 µg/ml propidium iodide and incubated at room
142 temperature for 5 min. 50 µg/ml RNase was then added and mixed and this mixture was used
143 for ploidy analysis [7]. The position of peak G1 nuclei of the control (Diploid callus derived
144 from immature anther) was established at channel 400 on a 1024-channel scale, after which
145 the instrument setting was kept constant and the test samples were run under the same
146 parameters.

147 *(MgCl₂ 45mM, MOPS 20mM, Sodium citrate 30mM, Triton X 100- 0.1% (vol/vol),
148 pH-7 and stored at -20°C as 10 ml aliquots)

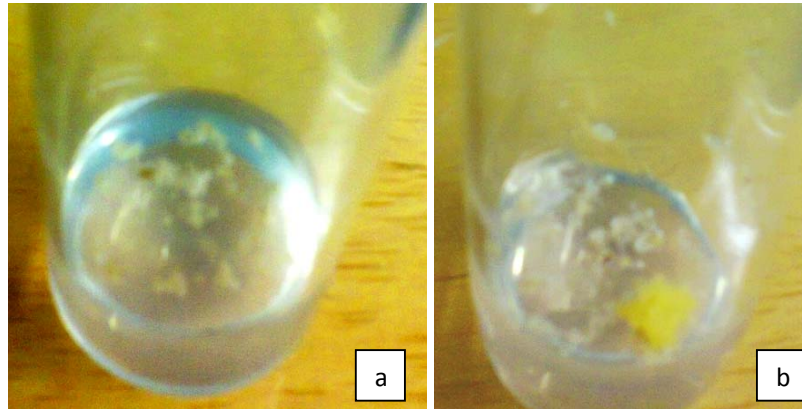
149 **Results**

150 Among the two different methods of colchicine application, addition of colchicine in the
151 culture medium was found to be ideal. In the first method of direct exposure to colchicine,
152 both colour and texture of the callus changed after colchicine treatment. Texture of the callus
153 changed from friable to spongy and simultaneously yellowish colour of the callus first turned
154 white and gradually became brown in colour. Rate of such changes was proportional to the
155 colchicine concentration to which the callus was exposed. At higher concentrations of
156 colchicine (1.0 – 2.5 µM) the calli turned brown and spongy within 12 h whereas in the calli
157 exposed to lower concentrations of colchicine (0.25-1.0 µM), these changes took place only
158 slowly. Also it was observed that duration of colchicine treatment did not have any
159 significant effect. The spongy, brown callus obtained after direct exposure to colchicine, did
160 not undergo any further growth or development, instead it gradually dried up. All the
161 treatments responded almost in the same way.

162 In the second method embryogenic calli were cultured, for different duration over
163 media supplemented with colchicine. The colchicine treated calli kept in the proliferation

164 medium first turned white in colour and watery, irrespective of the concentration of
165 colchicine and duration of exposure (Fig. 2a). However, emergence of new yellow
166 embryogenic callus occurred, within 4-6 weeks, from the calli exposed to lower
167 concentrations of colchicine (Fig. 2b). In this experiment this concentration of colchicine is
168 crucial for the emergence of new callus.

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Fig. 2 Colchicine treatment of diploid callus

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a Callus after colchicine treatment

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b Emergence of friable yellow callus after

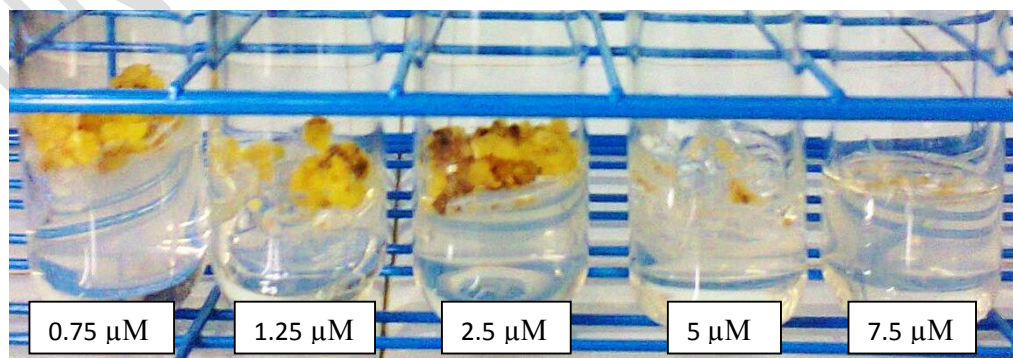
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colchicine treatment

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176 As evident in Fig.3, lower levels of colchicine (0.75-2.5 μM) did not affect the viability
177 of the treated callus and led to the emergence of new callus whereas those cultures exposed to
178 higher levels (5 and 7.5 μM) of colchicine did not give rise to any new callus, instead they
179 just got dried up on prolonged culture. Also it was observed that emergence and proliferation
180 rate of the callus varied with the period of exposure. Highest callus proliferation frequency of
181 73% was observed in the cultures exposed to 1.25 μM colchicine for 3 days. With the
182 increase in concentration of colchicine, along with increase in days of exposure, callus
183 proliferation rate was reduced. On the high side of colchicine (5 and 7.5 μM) and longer
184 duration of exposure (8 and 10 days), no callus emergence could be observed (Table 1).

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Fig. 3 Callus proliferation rate of calli exposed to different concentrations of colchicine

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190 **Table 1 Effect of colchicine concentration and exposure time on callus proliferation rate**

Colchicine concentration (μM) →	0.75	1.25	2.5	5.0	7.5
Days of exposure ↓					
2	46.5(42.9)	53.0(46.7)	33.8(31.0)	6.5 (14.7)	0.00 (0.33)*
3	57.5(49.3)	73.2(59.0)	47.5 (43.5)	3.5 (10.7)	0.00 (0.33)
6	25.5(30.3)	33.5(35.3)	6.5(14.7)	3.5(10.7)	0.00 (0.33)
8	7.0(15.3)	14.0(21.9)	0.00(0.33)	0.00 (0.33)	0.00 (0.33)
10	4.0(11.5)	4.0(11.5)	0.00 (0.33)	0.00 (0.33)	0.00 (0.33)

191

*Callus proliferation rate (%)

*CD= 1.61

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* CD- Coefficient of determination

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The data were subjected to arc sine transformation and transformed means are given in parenthesis

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In the embryo induction process MS medium was found to be most effective for induction of embryogenic callus and subsequent embryo formation (Fig.4a&b). Highest embryo induction frequency of 48% could be obtained in MS medium in the presence of 2.3 μM GA₃ and 1.8 μM BA. At lower concentrations of growth regulators, no embryo induction could be obtained. At higher concentrations also embryo induction was found to be less (Table 2).

200

Table 2 Effect of different concentrations of GA₃ and BA in embryo induction

GA ₃ (μM) →	0.57	1.2	1.73	2.3	2.9
BA (μM) ↓					
0.9	0.00(0.33)	0.00(0.33)	27.0 (31.3)	31.5(34.1)	23.0 (28.6)*
1.8	0.00(0.33)	0.00 (0.33)	35.5(36.57)	48(43.5)	37.5(37.5)
2.7	0.50(3.03)	5.5(13.5)	22.5(28.3)	27.5(31.6)	12.5(20.6)
3.6	1.0(5.7)	4.5(12.2)	16.0(23.5)	19.0(25.8)	5.5(12.0)
4.6	1.0(5.7)	3.5(10.7)	5.0(12.9)	7.5(15.8)	2.0(8.1)

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CD – 5.0

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* Percentage of embryo induction

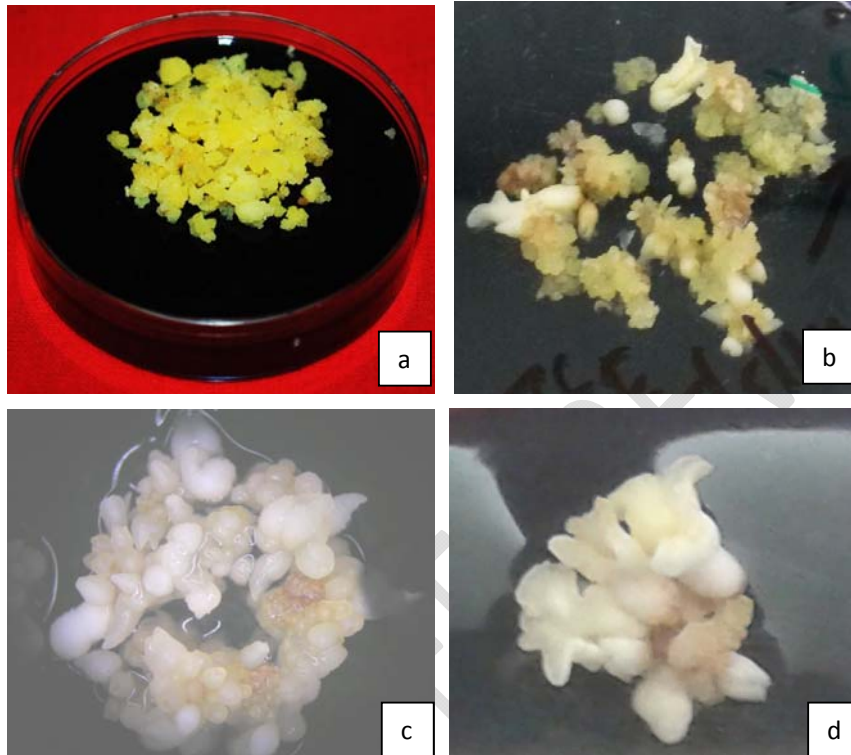
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The data were subjected to arcsine transformation and transformed means are given in parenthesis

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205 Embryos at different developmental stages like globular, heart shaped and
206 cotyledonary stages were obtained (Fig. 4c& d). It was also observed that the percentage of
207 embryo induction as well as the quality of the embryos depends on the colchicine
208 concentration in the treatment phase. Normal and healthy embryos were obtained from the
209 calli exposed to 1.25 μ M colchicine. Calli exposed to higher colchicine levels gave rise to
210 abnormal embryos, that too at a low frequency.

211



212

213

Fig. 4 Embryo induction from colchicine treated calli

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a Embryogenic callus

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b Induced embryos

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c& d Embryos at different stages of development

217

and cotyledonary

218

In case of embryo maturation a maturation frequency of 30 % could be obtained in
219 modified MS medium fortified with 4.7 μ M Kin and 1.9 μ M ABA (Table 3). In the hormone
220 free medium, maturation percentage was quite low. No increase in the embryo maturation
221 frequency was obtained by the addition of amino acids. However, quality of the embryos
222 could be improved in medium supplemented with the amino acid combination containing L
223 glutamine (15 mM) + L asparagine (10 mM) + L alanine (10 mM). Well developed, normal
224 and healthy matured embryos could be obtained from this combination.

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Table 3 Combined effects of Kin and ABA in embryo maturation

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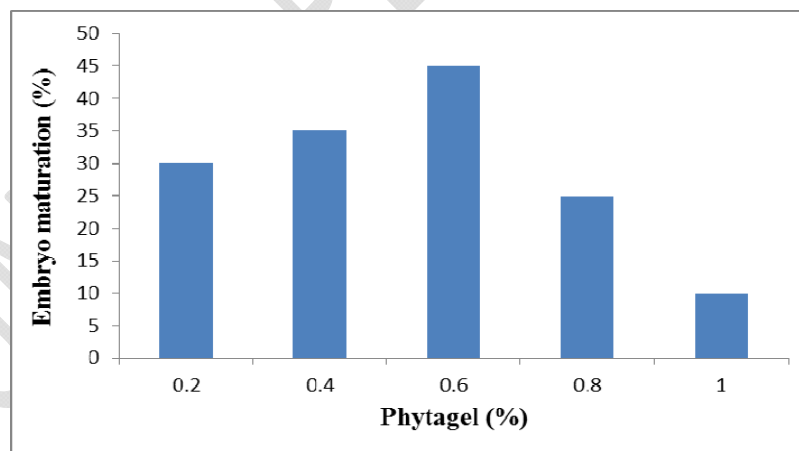
Kin (μM) \rightarrow	2.3	4.7	6.9	9.3
ABA (μM) \downarrow				
1.1	5.0(2.2)	21.5(4.6)	16.5(4.6)	12.0(3.4)
1.9	23.5(4.8)	30.2(5.5)	26.5(5.1)	22.5(4.7)
2.7	19.0(4.3)	26.5(5.1)	22.0(4.6)	18.0(4.2)
3.8	16.0(3.9)	14.0(3.7)	10.0(3.1)	6.5(2.5)

CD= 0.25

***Plant regeneration percentage**

The data were subjected to square root transformation and transformed means are given in parenthesis

Maturation frequency could be further enhanced by increasing the phytagel concentration. It was noticed that in a medium containing all the standardized parameters including basal medium, growth regulators and amino acids, an increase in the maturation frequency from 30 to 45 % could be achieved when the phytagel concentration was increased from 0.2 to 0.6 % (Fig. 5). At higher concentrations of phytagel the maturation frequency decreased, reaching 10% in the presence of 1.0% phytagel. Bipolar differentiation could be observed in some of the matured embryos (Fig. 6a).



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Fig. 5 Effect of phytagel concentration on embryo maturation

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Better embryo germination and plant regeneration were obtained in the presence of WPM, rather than MS medium. Mature embryos in the cotyledonary/late torpedo stage germinated (Fig. 6b) in WPM supplemented with 7.3 μM IBA and 6.6 μM BA. A germination frequency of 30 % was obtained with this growth regulator combination. Number of germinating embryos was quite low in media containing lower levels of growth regulators

259 (Table 4). Some of the embryos showed abnormalities in development. Some had multiple
 260 cotyledons, some became dormant and some showed only root development.

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Table 4 Effect of BA and IBA on embryo germination

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BA(μ M)→	2.2	4.4	6.6	8.8
IBA(μ M) ↓				
2.5	0.0(1.0)	1.5(1.6)	3.0(2.0)	2.0(1.73)*
4.9	4.5(2.3)	10.5(3.4)	14.5(3.9)	9.5(3.2)
7.3	11.5(3.5)	19.0(4.4)	30.0(5.4)	14.5(3.9)
9.9	5.0(2.4)	6.5(2.7)	9.5(3.2)	6.5(2.7)

CD=0.24

273

*** Percentage of embryo germination**

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The data were subjected to square root transformation and transformed means are given in parenthesis

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Among the various growth regulator combinations experimented for plant regeneration, 20% plant regeneration could be achieved in the combination of IAA (2.8 μ M), BA (8.8 μ M) and GA₃ (1.4 μ M) (Fig.6c). In other combinations, the regeneration frequencies were low (Table 5).

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Table 5 Effect of IAA and BA in presence of GA₃ 1.4(μ M) on plant regeneration

IAA (μ M)→	1.7	2.8	4.0	5.7
BA(μ M) ↓				
6.6	6.5(2.7)	10.5(3.4)	8.0(3.0)	7.0(2.8)
8.8	7.5(2.9)	20(4.5)	14.5(3.9)	4.5(2.3)
11.1	6.5(2.7)	15.5(4.0)	10.5(3.4)	3.5(2.1)
13.3	4.0(2.2)	2.0(1.7)	1.0(1.4)	0.0(1.0)

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CD=0.27

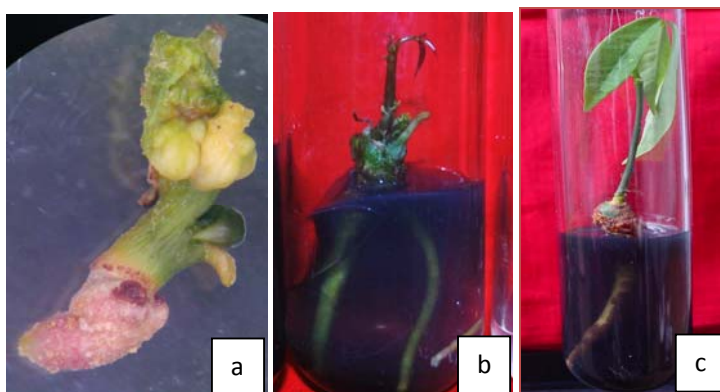
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***Plant regeneration percentage**

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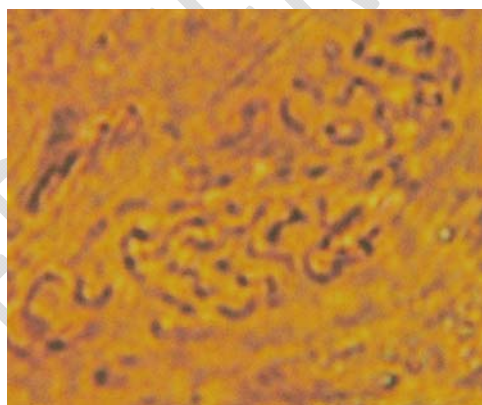
The data were subjected to square root transformation and transformed means are given in parenthesis

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287
 288 **Fig. 6 Plant regeneration from matured embryo obtained**
 289 **from colchicine treated calli**
 290 **a) Embryo maturation**
 291 **b) Embryo germination**
 292 **c) Regenerating plant**
 293

294 Cytological studies have revealed a chromosome count of $4n=72$ in the colchicine
 295 treated callus, observed at a magnification of X 400 (Fig. 7). Also from the root tip of one
 296 regenerated plant similar chromosome count was obtained. This confirms the tetraploid
 297 nature of the regenerated plant from the colchicine treated diploid callus.



298
 299 **Fig. 7 Chromosome count of colchicine treated callus $4n=72$ (X 400)**
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301 Using flow cytometry the ploidy of the colchicine treated callus was determined and
 302 from the histogram it can be observed that the fluorescence intensity of nuclei from the test
 303 sample got the highest peak at channel 780 which is double the value of control sample (Fig.
 304 8). This confirms that after treating the diploid embryogenic callus with colchicine, the nuclei
 305 content increased twice as that of the diploid callus, thereby resulting in the development of
 306 tetraploids.

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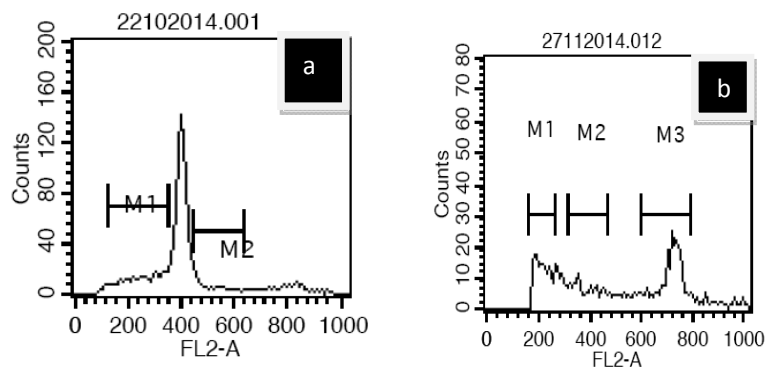


Fig. 8 Histogram showing peaks for
a Diploid (Immature anther derived callus)
b Tetraploid (Colchicine treated embryogenic callus)

Discussion

Development of synthetic polyploids of *Hevea brasiliensis* by applying colchicine on the axillary buds started by 1984 [8]. Success obtained through this method was very low due to the chances of occurrence of chimeras and the difficulty in stabilizing the polyploids. In *Hevea*, when seeds are used for colchicine treatment, it leads to loss of clonal integrity. Usage of explants like shoot apex and nodal segments for treatment, are not preferred because of the chances of obtaining chimeras. In the present study we have used callus as the explant for colchicine treatment. As the somatic embryo arises from a single cell, this method ensures complete tetraploidy of the regenerated plants and the chance for obtaining chimeras is remote. Also the clonal integrity can be maintained except for somaclonal variations. There are also reports in mulberry on the induction of tetraploidy through colchicine treatment of germinating seeds, seedlings and vegetative buds. Percentage of success was maximum (47 %) with callus explant compared to other explants [9].

Direct exposure may lead to abnormalities in cell division which can cause chromosome imbalance leading to low survival. Even lower concentrations of colchicine were lethal for direct exposure of callus. Since callus is a mass of loosely arranged single cells, penetration of colchicine into these cells will be more compared to the cells of seeds/embryos which are more compactly arranged. Callus induction from leaf explants of *Dionaea muscipula* soaked in different concentration of colchicines was attempted and they observed that callus induction rate decreased when colchicine concentration was high [10]. In the second treatment where colchicine was incorporated in the culture medium, there was no browning of the colchicine treated callus, instead the callus first turned white and watery and later, upon transfer to callus proliferation medium, new friable embryogenic calli emerged. Whitening and browning of the callus after colchicine treatment was also observed in case of winter rose where effect of different concentration of colchicine was experimented [11]. Colchicine concentration and its duration of application is another factor influencing the success. The exact concentration and time of exposure needs to be standardized in each crop and with different explants. A lot of changes happen in the cellular level with the application of colchicine. During cell division, colchicine is reported to arrest the spindle fiber formation

345 leading to doubling of chromosome number. Higher concentrations above the optimum will
346 result in abnormalities and lead to low success. Similarly exposure for longer durations is
347 also detrimental. Varying stress symptoms are shown by the explants exposed to colchicine
348 treatment. Callus growth was repressed at higher concentrations and/or longer treatment
349 duration with colchicines [12]. Most of the cells after colchicine treatment died, owing to
350 persistent lethality of colchicine.

351 It has been observed that the concentration as well as the duration of colchicine
352 treatment influenced callus proliferation. The proliferation rate of the callus was different for
353 each treatment. Out of the different treatments tried, highest callus proliferation of 73 % was
354 obtained in cultures treated with 1.25 μM colchicine for 3 days. At higher concentrations of
355 colchicine the proliferation rate was low. Similar observations were made in mulberry, where
356 the callus treated with 0.025 % colchicine for 3 days showed the highest percentage of
357 survival (76 %) [3]. Mortality rate is very important while applying colchicine to explants for
358 *in vitro* induction of polyploids [13]. In the present study with *Hevea* embryogenic callus,
359 survival rate decreased with increased concentrations of colchicine. 1.25 μM colchicine was
360 identified as an optimum concentration for the induction of tetraploids without compromising
361 the survival and further proliferation.

362 The culture conditions and basal media for somatic embryogenesis have already been
363 standardized by many workers in *Hevea* [14]. 0.2 mg/l NAA was effective for embryo
364 induction from immature anther [15]. In the present study 48 % embryo induction was
365 obtained in presence of 2.3 μM GA₃ and 1.8 μM BA in MS medium. The role of GA₃ (4.35
366 μM) and BA (8.84 μM) in differentiation of friable embryogenic calli derived from root
367 explants into somatic embryos were demonstrated in *Hevea* [14]. Embryo maturation in
368 *Hevea* is generally induced by a hormonal stress [15]. Hence most maturation media do not
369 contain auxins or cytokinins. But in our study 30 % embryo maturation in modified MS
370 medium supplemented with 4.7 μM Kin and 1.9 μM ABA was obtained. Normal
371 maturation of somatic embryos needs an ABA treatment in walnut [16]. Similarly
372 maturation of somatic embryos of Fraser fir was not observed on medium lacking ABA for
373 both precotyledonary and cotyledonary embryos [17]. 80 μM ABA was most effective in
374 producing cotyledonary stage embryos in *A. fraseri*. The role of phytigel in embryo
375 maturation is well established in *Hevea*, a drastic increase in the embryo maturation
376 frequency was noticed when phytigel was increased from 0.2 to 0.5 % [18]. Raising the
377 concentration of phytigel in the medium from 0.4 % to 0.8% improved the maturation of
378 somatic embryos of *Larix eurolepis* [19]. The use of high concentrations of phytigel as
379 gelling agent reduces water availability. Similarly 0.5 % phytigel was effective in the
380 maturation of anther derived embryos of coconut [20].

381 Earlier reports are there indicating the beneficial effect of GA₃ on germination in
382 *Hevea* [21]. Also a combination of BA (0.3 mg l⁻¹) and GA₃ (0.3 mg l⁻¹) has been reported to
383 favour germination of rescued zygotic embryos in *Hevea* [22]. However in our result embryo
384 germination (30 %) from colchicine treated callus was obtained in MS basal medium fortified
385 with IBA (7.3 μM) and BA (6.6 μM). Similar to our results, a combination of BA and IBA
386 was used in mulberry [3], for the induction of shoot and root from the colchicine treated
387 callus. Embryo culture to enhance efficiency of colchicine induced polyploidization in grape
388 fruit reported that even a low level (0.03mg l⁻¹) of colchicine was lethal for embryos towards
389 germination [23]. But in our study such lethality was not observed in the embryos obtained
390 from callus treated with 1.25 μM colchicine for 3 days. Lower concentrations of colchicine
391 (0.1%, 0.2% and 0.3%) did not affect the rooting behavior [24]. However, the number of
392 roots developed was considerably decreased in the progeny treated with higher concentration
393 of colchicine (0.4% and 0.5%). In our study, root growth was not affected in the embryos

394 raised from cultures exposed to 1.25 μM colchicine. Embryos germinated with well-
395 developed root system. Plant regeneration frequency of 20 % was obtained in WPM medium
396 fortified with IAA (2.8 μM), BA (4.4 μM) and GA₃ (5.7 μM). Similar basal medium (WPM)
397 fortified with GA₃ (2.9 μM) and BA (8.8 μM), brought about 60 % plant regeneration
398 frequency from root explants of *Hevea* [25].

399 Ploidy determination of the callus and regenerated plants through cytological and
400 flow cytometry analyses revealed tetraploidy ($4n= 72$). Now a day's flow cytometer is used
401 for ploidy determination since it is time saving and because of its easiness to predict result.
402 The DNA content in the colchicine treated callus was doubled when compared with the
403 control having the highest peak at 400 channel.

404 **Conclusion**

405 Development of tetraploids through chromosome doubling of diploid callus using
406 colchicine has been achieved. Embryo induction, germination and plant regeneration have
407 been obtained. Ploidy determination of the callus and regenerated plants through cytological
408 and flow cytometry analyses revealed tetraploidy ($4n= 72$). Acclimatization and field
409 establishment needs to be accomplished for further evaluation of the tetraploid plants.

410 In vitro induction of ploidy variation in *Hevea* has been accomplished through the
411 development of tetraploids, using appropriate techniques. The ploidy was confirmed using
412 cytological and flow cytometric analyses. The tetraploid plantlets can be further utilized in
413 *Hevea* breeding for enhancing the vigor and productivity. The breakthrough achieved in the
414 development of tetraploids could pave the way for developing future strategies for exploiting
415 the benefits of ploidy variation towards crop improvement in *Hevea brasiliensis*.

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420 **Competing Interests**

421 Authors have declared that no competing interests exist

422 **Authors contribution**

423 Author 'SKS' designed the study. Author 'SKS' and 'DUK' performed the statistical
424 analysis, wrote the protocol, and wrote the first draft of the manuscript. 'Author 'DUK'
425 managed the analyses of the study as well as literature searches. "All authors read and
426 approved the final manuscript."

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