Effect of growth regulators on callus morphology of Rice anther culture

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ABSTRACT

The present investigation was undertaken to know the effect of various combination of auxin and cytokinin on callus morphology in japonica rice. As the callus induction is a prerequisite for anthers culture and finally development of haploids certain callus morphology criteria empirically identified. Among the colour of callus, this was either white or yellow. The white colour was preferred. In addition to callus growth determine empirically low, medium and high define as 1, 2 and 3 respectively. Callus texture was also identified as compact and friable but compact texture was accepted as more promising. Callus morphology is an empirical which can be use to predict the regeneration ability of calli.

1. INTRODUCTION

Rice (Oryza sativa L.) is the world most important food supplier cereal crop after wheat and maize. It provides half of total dietary carbohydrate, especially in Asian countries and it is suitable diet for more than three billion people, supplying 50-80% of their daily calorie intake [1]. However, traditional rice breeding methods are not sufficient to fulfill the demands of growing population. The production of haploids via anther culture represents an alternative biotechnology tool for crop improvement programs. Several advantages of haploid culture technique have been reported in the breeding program [2, 3 and 4]. Anther culture is an important technique to develop homozygous line by shortening of the breeding cycle of new varieties and allows early expression of recessive genes. Beside the advantages of using the techniques, anther culture has disadvantages and constraints, i.e., low efficiencies of callus production, low frequency of plant regeneration, and high proportion of albino plants [5, 6]. Several factors are effecting the callus texture derived from in vitro anther culture of rice are types of growth regulator along with its concentrations, genotype and microspore developmental stages [7]. Texture and colours of the calli reflect their green plantlet redifferentiation competences. The embryogenic calli which were milky white in colour and compact in texture had excellent regeneration ability. However, friable calli had poor plant regeneration ability or did not respond at all. These results are indicated that the callus induction medium has an influence on the morphogenic competence of the induced callus, determining its regeneration capability [8]. It was reported that there are conditions in which genotypes show high callus induction has displayed poor regeneration ability and vice versa [9]. Application of higher dose of auxin sources can significantly increase the callus induction efficiency, however such calli are less in embryogenic and poor in green plant regeneration. Anthers of three rice cultivars viz, BR-3, BR-10 and BRRI Dhan 29 produced friable and compact callus texture with white in colour in Z2 media containing 2 mg/L 2, 4-D + 2.5 mg/l NAA + 0.5 mg/l Kinetin [10]. Many an embryogenic and nonembryogenic callus with multiple colours (white, yellow and brown) reported in rice cultivar Swat II on MS media containing different concentration of auxin and kinetin with Tryptophan [11]. The rice cultivars chinigura, kalijira, Radhuni pagal, modhumala, kataribog and mohonbhog produce compact callus texture with white in colour in media containing different concentration of 2, 4-D and NAA [12]. Anthers of BC3F2 of Oryza sativa L. × Oryza rufipogon regenerate double haploid lines through compact callus texture with light green colour in N6 media containing 2 mg/l NAA + 0.5 mg/l Kinetin + 2 mg/l 2, 4-D [13]. Different type of callus texture with multiple callus colour reported in rice cultivar Swarna on N6 media containing 2, 4-D, 2, 4-D + Kinetin, 2, 4-D + BAP and 2, 4-D + NAA [14]. With this background the present study was aimed to know effect of growth regulators on morphological characters of callus (texture and colour) produced by japonica rice varieties through anther culture.
2. MATERIALS AND METHODS

Seeds of two *japonica* rice varieties viz. Azucena and Moroberekan were grown under standard agronomic practices. The panicles from the plants of each variety were harvested between 6.00 to 9.00 a.m. on sunny days. Panicles were harvested when the distance between collar of flag leaf and penultimate leaf was about 10-15 cm. Panicles were wrapped in aluminum foil with a moisturized cotton plug at the base and sealed in polypropylene bags separately. Surface sterilized panicles were kept for cold pre-treatment at 4°C for one week [15]. The anthers with mid-uninucleate stages were first determined by cytological test using acetocarmine staining technique. Cold treated panicles were sterilized by immersing them in 70% ethanol for 20 seconds followed by immersing in 0.2% of HgCl₂ for 10 min and washed thoroughly with sterile water [16]. The spikelets were picked up with their open ends down by a forcep and tapped on the rim of a bottle, causing anthers in the spikelets to vibrate and fall onto surface of callus induction medium. The N6 medium supplemented with various concentration of 2, 4-D, NAA and Kinetin with 3% maltose and 0.8% agar. The cultures were placed in the dark at 23±2°C with relative humidity 50-60% for callus induction.

The cultures were subculture at weekly intervals for 10-20 weeks. The observations were recorded as number of anthers responded, calli texture with colour and callus growth.

### Table 1: Morphological characteristics of the androgenic callus obtained from Azucena and Moroberekan.

<table>
<thead>
<tr>
<th>Treatments</th>
<th>No. of anthers inoculated*</th>
<th>No. of calli obtained</th>
<th>Colour of callus</th>
<th>Type of callus</th>
<th>Callus Growth</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Azucena</td>
<td>Moroberekan</td>
<td>Azucena</td>
<td>Moroberekan</td>
<td>Azucena</td>
</tr>
<tr>
<td>a) 2,4-D + Kinetin</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>T₀ (Control)</td>
<td>28.33</td>
<td>38.66</td>
<td>0</td>
<td>0</td>
<td>-</td>
</tr>
<tr>
<td>T₁ (1mg/l + 0.5 mg/l)</td>
<td>34.00</td>
<td>25.00</td>
<td>2</td>
<td>0</td>
<td>White</td>
</tr>
<tr>
<td>T₂ (1mg/l + 1 mg/l)</td>
<td>56.66</td>
<td>52.66</td>
<td>9</td>
<td>0</td>
<td>White</td>
</tr>
<tr>
<td>T₃ (2 mg/l + 0.5 mg/l)</td>
<td>69.00</td>
<td>38.66</td>
<td>2</td>
<td>3</td>
<td>Yellow</td>
</tr>
<tr>
<td>T₄ (2 mg/l + 1 mg/l)</td>
<td>27.66</td>
<td>36.66</td>
<td>13</td>
<td>7</td>
<td>Yellow</td>
</tr>
<tr>
<td>b) NAA + Kinetin</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>T₀ (Control)</td>
<td>28.33</td>
<td>38.66</td>
<td>0</td>
<td>0</td>
<td>-</td>
</tr>
<tr>
<td>T₁ (1mg/l + 0.5 mg/l)</td>
<td>38.66</td>
<td>47.66</td>
<td>4</td>
<td>3</td>
<td>White</td>
</tr>
<tr>
<td>T₂ (2 mg/l + 0.5 mg/l)</td>
<td>53.00</td>
<td>45.00</td>
<td>3</td>
<td>1</td>
<td>Yellow</td>
</tr>
<tr>
<td>T₃ (2 mg/l + 1 mg/l)</td>
<td>35.66</td>
<td>47.00</td>
<td>4</td>
<td>5</td>
<td>White</td>
</tr>
<tr>
<td>c) 2,4-D + NAA + Kinetin</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>T₀ (Control)</td>
<td>28.33</td>
<td>38.66</td>
<td>0</td>
<td>0</td>
<td>-</td>
</tr>
<tr>
<td>T₁ (1mg/l + 0.5 mg/l)</td>
<td>38.33</td>
<td>54.66</td>
<td>0</td>
<td>5</td>
<td>Yellow</td>
</tr>
<tr>
<td>T₂ (1mg/l + 1 mg/l)</td>
<td>47.00</td>
<td>38.33</td>
<td>1</td>
<td>1</td>
<td>White</td>
</tr>
<tr>
<td>T₃ (1mg/l + 2 mg/l + 0.5 mg/l)</td>
<td>38.66</td>
<td>37.00</td>
<td>7</td>
<td>18</td>
<td>White</td>
</tr>
<tr>
<td>T₄ (1 mg/l + 2 mg/l + 1 mg/l)</td>
<td>36.00</td>
<td>41.66</td>
<td>2</td>
<td>2</td>
<td>Yellow</td>
</tr>
<tr>
<td>T₅ (2 mg/l + 1 mg/l)</td>
<td>63.66</td>
<td>43.33</td>
<td>2</td>
<td>17</td>
<td>Yellow</td>
</tr>
<tr>
<td>T₆ (2 mg/l + 1 mg/l + 0.5 mg/l)</td>
<td>33.33</td>
<td>43.33</td>
<td>4</td>
<td>7</td>
<td>Yellow</td>
</tr>
<tr>
<td>T₇ (2 mg/l + 2 mg/l + 0.5 mg/l)</td>
<td>53.66</td>
<td>46.00</td>
<td>5</td>
<td>8</td>
<td>White</td>
</tr>
<tr>
<td>T₈ (2 mg/l + 2 mg/l + 1 mg/l)</td>
<td>57.00</td>
<td>51.66</td>
<td>7</td>
<td>12</td>
<td>White</td>
</tr>
</tbody>
</table>

Legend: * Average of 3 replication; - No callus; + low callus growth; ++ Medium growth; +++ High callus growth.

Fig.1: Callus morphology of Azucena (a, b) and Moroberekan (c, d).
3. RESULTS AND DISCUSSION

Rice (Oryza sativa L., 2n=24) is classified under the tribe Oryzeae, subfamily Oryzoideae of the grass family Poaceae [17]. Among cereals, it is one of the most studied, serving as staple food of more than half of the world’s population [18]. Rice is composed of two subspecies: indica rice, described as tropical long grain, accounts for approximately 80% of cultivated rice, and japonica rice, with rounded and shorter grain, is more adapted to temperate climates [19].

Tissue culture technique is known as a novel means to create genetic variability and has been proposed as an excitant supplementary technique for plant which can accelerate the breeding programs through the use of new expended genetic variability. Successful application of tissue culture method involved the establishment of a more or less de-differentiated cells or tissue under defined culture condition, proliferation of a number of cells and the subsequent regeneration of plants [20]. Many new rice cultivars have been developed through biotechnological techniques like anther culture, embryo rescue and somaclonal variation [21 and 22].

3.1 Callus texture and colour

In the present study results obtained on morphology of callus was presented in Table 1. The cultured anthers were started turning brown after two weeks of inoculation. The first indication of callus initiation was swelling of the anther wall followed by emergence of microcalli from anther lobes. Later, callus appeared from the cut ends. This result was not surprising because it also reported changing of anther colours which is possibly due to the transition of gametophytic phase to sporophytic phase during androgenesis [23].

Irrespective of varieties, among 2, 4-D + Kinetin growth regulator combination of treatments in Azucena calli were compact and white colour obtained in only one treatment (T1) and yellow with compact calli were obtained in the two treatments (T2, T3) however, similar results obtained on (T3, T5) and (T2) in Moroberekan respectively. Yellow calli with compact type reported in rice genotypes on N6 media containing 2, 4-D + Kinetin [24]. Friable callus texture with cream colour found in rice cultivar MR 219 on N6 media containing 2, 4-D and Kinetin either alone or in combination which was contradictory of the present study [25]. Yellow and white colour callus reported in three japonica rice varieties Hayahishiki, Fujisaka 5 and Nipponbare on N6 media containing 2, 4-D [26].

Combination of 2, 4-D + Kinetin was more effective in producing embryogenic callus compared to non-embryogenic callus [27]. Embryogenic calli with nodular structures appeared on the surface of the non-embryogenic callus. These calli tend to be light yellow to whitish in color which is slightly different from the non-embryogenic callus. Nonembryogenic calli with white, wet and friable characters were found predominantly under dark condition. It was documented that embryogenic callus displayed higher frequency of plant regeneration than the non-embryogenic one [28]. Therefore, media modifications should target the production of embryogenic callus with good regeneration ability rather than simply inducing prolific callusing, from which regeneration would not be possible [29]. Among NAA + Kinetin combination of growth regulator treatments in Azucena white calli with compact type was obtained in three treatments (T8, T9, T10) and yellow calli with compact type was obtained in the one treatment (T2) (Fig. 1a,b). This finding agrees with the earlier observations [30]. However, in Moroberekan white and compact type of calli were obtained in two treatments (T6, T9) and yellow calli with friable type were observed in one treatment (T7) (Fig. 1c, d). Yellow calli with compact and friable type reported in BC1F1 anthers of (KDML105//IRBB5/KDML105) on N6 media containing NAA + Kinetin [31]. The combination of auxin and cytokinin growth regulators: 2, 4-D + NAA + Kinetin treatments in Azucena calli were white with compact obtained in four treatments (T1b, T11, T15, T16) and two treatments (T12, T13) were produced yellow calli with compact and one treatment (T1s) yellow with friable type calli were obtained. In Moroberekan, white calli with compact type was produced in four treatments (T1b, T12, T14, T1s) and three treatments (T7, T13, T16) was obtained yellow calli with compact type and only one treatment (T11) was produced yellow calli with friable type. These results also supported to the earlier studied [32]. The white friable callus might be due to higher doses of hormone in the callus induction medium [33]. It was reported that hormone requirement is genotype specific and optimum level of auxin in the callus induction media required some degree of compromise between callus induction and regeneration frequency [34].

3.2 Callus growth

Among the 2, 4-D + Kinetin growth regulator combination that induced callus in the two varieties, one treatment in Azucena (T2) obtained high callus growth and rest of the treatment (T1, T3) induced poor callus growth. However, in Moroberekan, poor callus growth was induced in treatments (T2, T3, T4). Anthers of Kyoto Asahi rice variety produced good callus growth at N6 media containing 2, 4-D + Kinetin [35]. However, NAA + Kinetin growth regulator combination that induced poor callus growth in both the varieties. Among the 2, 4-D + NAA + Kinetin growth regulator combination induced poor callus growth in seven treatments (T11, T12, T13, T14, T15, T16) in Azucena, whereas in Moroberekan highest (T11) and moderate callus growth was obtained in treatments (T13) and rest of the treatments (T9, T10, T12, T14, T15, T16) was induced poor callus growth. Callus growth reported in japonica rice variety Azucena at N6 media containing 2, 4-D + NAA + Kinetin with similar grading of callus growth [36]. In conclusion embryogenic calli with compact type was predominant in both Azucena and Moroberekan varieties. Growth regulator along with its concentration has ability to produce different types of callus, colour of callus and callus growth. Therefore, types of growth regulator along with its concentration need to be considered in order to achieve high embryogenic calli and callus growth.
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