IN VITRO RELATIONSHIP BETWEEN ORGANOGENESIS FROM *Datura metel* L. ANther AND CATALASE ACTIVITY

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**ABSTRACT**

The aim of the present study was to investigate the effect of different concentrations of plant growth regulators on *in vitro* induction of callus and plantlets regeneration from anthers culture of *Datura metel* L. The activity of catalase (CAT) as one of the important antioxidant enzymes was measured during different stages of callus and organogenesis under all combinations of hormones. Anthers were cultured under aseptic conditions on Murashige and Skoog (MS) medium supplemented with benzyladenine (BA) (0.0, 3.0, 5.0 and 7.0 mg/l) in combination with 2,4-D (2,4-dichlorophenoxy acetic acid) (0.0, 0.01 or 0.05 mg/l) for callus induction and organogenesis. Results showed that MS containing different combinations of BA and 2,4-D induced calli with percentage ranging between 20-100%. High concentrations of growth regulators enhanced the highest callus percentage and promoted plantlets regeneration by indirect organogenesis reached to higher numbers of regenerated plantlets (4.5 and 7.4) at 5.0 mg/l BA + 0.05 mg/l 2,4-D and 7.0 mg/l BA + 0.05 mg/l 2,4-D, respectively. Results revealed that catalase activity increased significantly in organogenic callus with regenerated plantlets (23.5 and 47.3 unit) at higher concentrations of BA and 2,4-D (5.0 mg/l BA + 0.05 mg/l 2,4-D and 7.0 mg/l BA + 0.05 mg/l 2,4-D, respectively).

**Key words:*** Organogenesis, anther culture, *Datura metel* L., catalase activity, *in vitro* culture.

**INTRODUCTION**

The Solanaceae family contains many important plants. *Datura metel* L. is one of the pharmaceutical plants belonging to this family that produces secondary metabolites such as alkaloids (Sato et al., 2001; Murch et al., 2009). Plant tissue culture technique induce plants to produce secondary metabolites under controlled conditions (Gumuscu et al., 2008), or as an efficient method for propagation of many plants using various explants by either shoot proliferation and multiplication from shoot tip and nodal explants (Shahsavari et al., 2010; Ibrahim et al., 2013; Martins et al., 2015), or organogenesis in many plant species (Sudhersan and Hussain, 2003; Radhika et al., 2006; Selvaraj et al., 2006; Mannan et al., 2013). Anther culture technique is one of these methods, which has been successful in many plants. Through this method, the production of homozygous doubled haploids can be achieved in short period that made it useful in plant breeding (Nurhidayah et al., 1996; Kadota and Niimi, 2004).

Plant cells secreted many antioxidant enzymes which are different in their nature and function according to the type of oxidation. These enzymes can play an important role in controlling of reactive oxygen species (ROS) that causing damages to plant cells (Blokhina, 2000). Catalase (CAT) is one of these important defending enzymes in plant tissues, it works on hydrogen peroxide ($H_2O_2$) that resulted from the cellular metabolism processes and converts it to water and oxygen molecules (Willekens et al., 1997).

The aim of this study is to induce callus and organogenesis from *Datura metel* L. anther and to survey the changes in Catalase activity during these processes.

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MATERIALS AND METHODS

Preparation, Sterilization and Culture of Explants

Flower buds were collected from *Datura metel* L. plants. In the laboratory, they were immersed in ethanol (96%) for 10 minutes for surface sterilization, then rinsed with sterilized distilled water for 3 minutes. The buds were transferred to a sterile Petri dish and were split open using a blade and the anthers were excised under aseptic conditions without damage and the filaments were removed, then anthers were cultured horizontally (Fig. 1) on Murashige and Skoog (MS) medium (Murashige and Skoog, 1962), supplemented with BA (0.0, 3.0, 5.0 or 7.0 mg/l) alone or in combination with 2,4-D (0.0, 0.01 or 0.05 mg/l) to induce callus and organogenesis. Cultures were incubated in growth room under controlled conditions (25 ± 2°C and 1000 Lux for 16 hr., photoperiod). Ten replicates were used for each treatment (in each replicate, three anthers were cultured). Experiments were repeated two times. The frequency of callus induction (%) from anthers was measured according to Zaidi *et al.* (2006) formula:

\[
\text{Frequency of callus induction} (\%) = \frac{\text{No. of anthers induced callus}}{\text{No. of cultured anthers}} \times 100
\]

The number of regenerated plantlets were calculated after 45 days from culture.

Estimation of Catalase activity

The reaction solutions were prepared according to Whitaker (2003) as follows:

a) Phosphate buffer (50 mM): (1) Dissolve 6.81g KH \(_2\)PO\(_4\)_2H\(_2\)O in water and make to 1L. (2) Dissolve 8.90g Na\(_2\)HPO\(_4\)_2H\(_2\)O in water and make to 1L. Mix solutions (1) and (2) in proportion of 1:1.5 (V/V), pH 7.0.

b) Hydrogen Peroxide (H\(_2\)O\(_2\) 30 mM): Dilute 0.34 ml of 30% H\(_2\)O\(_2\) (freshly opened) with phosphate buffer, pH 7.0 (above) to 100 ml (smaller volumes of reagents can be prepared by keeping the concentrations constant as given).

c) Sample of catalase : prepared in the 50 mM , pH 7.0 Phosphate buffer.

Catalase activity was assayed from the hydrogen peroxide decomposition rate as measured by the decrease of absorbance at 240 nm, and the activity of enzyme calculated according to the following equation (Frary *et al.*, 2010):

\[
\Delta\text{Abs/min} \times \text{reaction volume}
\]

\[
\text{Catalase activity (unit)} = \frac{\text{reaction volume}}{0.001}
\]

Where :

\(\Delta\text{Abs} : \text{the difference in absorbance.}\)

\(\text{reaction volume: final volume of reaction sample in cuvette.}\)

\(\text{min : reaction time.}\)

0.001: is the enzyme quantity that causing change in absorbance in 1 min.

Statistical Analysis

Data were analyzed using completely randomized design (CRD) and means were compared using the least significant difference (LSD) at 0.05 (GentSas, 2007).

![Fig. 1. Process of excised and cultured of *Datura metel* L. anthers. (A) flower buds. (B,C) excised of anthers. (D) anthers cultured on MS medium](image-url)
RESULTS AND DISCUSSION

Callus Induction

Results in Table 1 and Fig. 2 show differences in callus induction frequency from anthers between BA and 2,4-D concentrations. BA gave the best callus induction percentages (76.7 and 83.3%) at 5.0 and 7.0 mg/l, respectively. Results show also that MS supplemented with 2,4-D at 0.01 and 0.05 mg/l was most effective in callus induction (85.0 and 90.0%, respectively) as compared with the control. The percentage of anthers that forming callus was significantly higher (100%) on MS medium supplemented with BA 5.0 mg/l + 0.01 mg/l 2,4-D, 5.0 mg/l BA + 0.05 mg/l 2,4-D and 7.0 mg/l BA + 0.05 mg/l 2,4-D as compared with control (since the MS basal medium alone did not form callus on anthers). Previous studies indicated that plant hormones played an important role in callus induction from different explants, and the best response different according to the original plant species and the type of explants that used in tissue culture (Brasileiro et al., 1999; Alagumanian et al., 2004; Ibrahim et al., 2012; Mannan et al., 2013). Plantlets Regeneration from Callus

After the formation of callus on anthers under different concentrations of growth regulators; indirect organogenesis from callus was occurred in the same combinations of BA and 2,4-D (Table 2). Within 45 days of culture, green swellings appeared on the surface of callus, then, these swellings were regenerated to plantlets (Fig. 3). As results in Table 2, the maximum mean number of plantlets (4.83) was obtained on medium containing 7.0 mg/l BA that was significantly higher compared with other BA concentrations. The presence of cytokinins is necessary for regeneration, so the treatments that lack of BA did not induce any regeneration response. Researchers found that addition of cytokinins in suitable concentrations could be stimulate shoot or bud formation from callus (Kaviani et al., 2011).

Plantlets regeneration was occurred also in 2,4-D concentrations in which the highest significant number of regenerated plantlets (3.80 plantlets) was recorded with 0.05 mg/l as compared with other concentrations. There were significant differences in regeneration when MS medium containing different combinations of BA and 2,4-D. In general, the maximum number of regenerated plantlets (7.40) was obtained in MS + 7.0 mg/l BA + 0.05 mg/l 2,4-D comparing with other interactions whereas the control, MS + 0.01 mg/l 2,4-D and MS + 0.05 mg/l 2,4-D did not induce any morphogenic responses and they grown in randomly pattern. Mishra et al. (2006) reported that non-organogenic callus grows in unorganized manner, but the organogenic callus grows very fast and gives rise to multiple shoots or buds. Cytokinins and auxins are usually known to promote the formation of callus and shoots in many excited and in vitro cultured explants. Proper type and concentration of these growth regulators are different for each plants and explants (Jain and Ochatt, 2010; Mannan et al., 2013).

Catalase Activity

Catalase activity changes according either to the physiological status of plant cells or to the composition of medium specially hormones. Results in Table 3 show that there was gradual increasing in the enzyme activity with the increasing of BA concentration reaching to 20.0 units at 7.0 mg/l BA. Significant increasing in activity was occurred with 2,4-D concentrations in which the highest concentration (0.05 mg/l) gave the maximum activity of enzyme which was 20.9 units.

The interaction results show that the organogenic callus with regenerated plantlets on MS medium supplemented with 7.0mg/l BA+ 0.05 mg/l 2,4-D, 5.0 mg/l BA + 0.05 mg/l 2,4-D, 7.0 mg/l BA + 0.01 mg/l 2,4-D and 3.0 mg/l BA + 0.05 mg/l 2,4-D gave the maximum catalase activities (47.3, 23.5, 12.1 and 12.5 units, respectively) followed by organogenic callus only as compared with other interactions, whereas the control showed the lowest activity due to no callus was induced on it. The changes in catalase activity may be due to the synthesis of amino acids that required for the synthesis of this important antioxidant enzyme and certain proteins that required for callus proliferation and to initiate shoots or buds or roots formation (Palai et al., 2000 ; Tian et al., 2003; Konieczy et al., 2014).
# Table 1. Influence of different concentrations of BA and 2,4-D (mg/l) in MS medium on callus induction frequency (%) from anthers of *Datura metel. L.*

<table>
<thead>
<tr>
<th>2,4-D (mg/l)</th>
<th>0.0</th>
<th>0.01</th>
<th>0.05</th>
<th>Mean of BA</th>
</tr>
</thead>
<tbody>
<tr>
<td>BA (mg/l)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>0.0</td>
<td>0.0</td>
<td>60.0</td>
<td>70.0</td>
<td>43.3</td>
</tr>
<tr>
<td>3.0</td>
<td>20.0</td>
<td>80.0</td>
<td>90.0</td>
<td>63.3</td>
</tr>
<tr>
<td>5.0</td>
<td>30.0</td>
<td>100.0</td>
<td>100.0</td>
<td>76.7</td>
</tr>
<tr>
<td>7.0</td>
<td>50.0</td>
<td>100.0</td>
<td>100.0</td>
<td>83.3</td>
</tr>
<tr>
<td>Mean of 2,4-D</td>
<td>25.0</td>
<td>85.0</td>
<td>90.0</td>
<td></td>
</tr>
<tr>
<td>LSD (0.05)</td>
<td>BA = 17.89</td>
<td>2,4-D = 15.50</td>
<td>BA * 2,4-D = 30.99</td>
<td></td>
</tr>
</tbody>
</table>

Fig. 2. Formation of callus on anthers cultured on MS medium supplemented with different concentrations of BA and 2,4-D.

# Table 2. Influence of different concentrations of BA and 2,4-D (mg/l) in MS medium on number of regenerated plantlets from callus anthers of *Datura metel. L.*

<table>
<thead>
<tr>
<th>2,4-D (mg/l)</th>
<th>0.0</th>
<th>0.01</th>
<th>0.05</th>
<th>Mean of BA</th>
</tr>
</thead>
<tbody>
<tr>
<td>BA (mg/l)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>0.0</td>
<td>0.0</td>
<td>0.0</td>
<td>0.0</td>
<td>0.0</td>
</tr>
<tr>
<td>3.0</td>
<td>0.1</td>
<td>1.5</td>
<td>3.3</td>
<td>1.63</td>
</tr>
<tr>
<td>5.0</td>
<td>0.2</td>
<td>2.4</td>
<td>4.5</td>
<td>2.37</td>
</tr>
<tr>
<td>7.0</td>
<td>2.8</td>
<td>4.3</td>
<td>7.4</td>
<td>4.83</td>
</tr>
<tr>
<td>Mean of 2,4-D</td>
<td>0.77</td>
<td>2.05</td>
<td>3.80</td>
<td></td>
</tr>
<tr>
<td>LSD (0.05)</td>
<td>BA = 0.97</td>
<td>2,4-D = 0.84</td>
<td>BA * 2,4-D = 1.68</td>
<td></td>
</tr>
</tbody>
</table>
Fig. 3. Plantlets regenerated from anther cultures, (A) green swellings on callus, (B,C) plantlets regeneration through callus (arrows) on medium with 7.0 mg/l BA+0.05 mg/l 2,4-D and 5.0 mg/l BA+ 0.05 mg/l 2,4-D.

Table 3. Influence of different concentrations of BA and 2,4-D (mg/l) in MS medium on catalase activity (unit) of different stages of callus and organogenesis in anthers cultures of *Datura metel* L.

<table>
<thead>
<tr>
<th>BA(mg/L)</th>
<th>2,4-D(mg/L)</th>
<th>0.0</th>
<th>0.01</th>
<th>0.05</th>
<th>Mean of BA</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.0</td>
<td>0.1</td>
<td>0.2</td>
<td>0.3</td>
<td>0.2</td>
<td></td>
</tr>
<tr>
<td>3.0</td>
<td>0.3</td>
<td>6.1</td>
<td>12.5</td>
<td>6.3</td>
<td></td>
</tr>
<tr>
<td>5.0</td>
<td>0.4</td>
<td>8.3</td>
<td>23.5</td>
<td>10.7</td>
<td></td>
</tr>
<tr>
<td>7.0</td>
<td>0.5</td>
<td>12.1</td>
<td>47.3</td>
<td>20.0</td>
<td></td>
</tr>
<tr>
<td>Mean of 2,4-D</td>
<td>0.3</td>
<td>6.7</td>
<td>20.9</td>
<td></td>
<td></td>
</tr>
<tr>
<td>LSD (0.05)</td>
<td>BA= 15.50</td>
<td>2,4-D= 13.42</td>
<td>BA * 2,4-D= 26.85</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

The increasing in catalase activity in organogenic callus with regenerated plantletes is in agreement with Gupta and Datta (2003) and Meratan et al. (2009) who referred in their studies that the greatest increasing in catalase activity of regenerated shoots than in callus and regenerated roots may suggest the enzyme effective scavenging mechanism to remove hydrogen peroxide that produced in regenerated shoots.

**Conclusion**

The present study revealed that the callus formation and organogenesis from callus cultures was dependent on plant growth regulators concentrations. Catalase may be used as a marker for organogenesis in *Datura metel* L. anther cultures.

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العلاقة بين نشوء الأعضاء من منك الداتورة

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تهدف الدراسة الحالية إلى التحري عن تأثير التركيزات المختلفة لمنظومات النمو النباتية في استجابة الكلاسك واستعادة
تكوين النباتات من مزارع طائر الداتورة (Datura metel L.) (معملًا، كما تم قياس فعالية إزيم الكاتلز كواحد من أهم
الإحترامات للكاذبة خلال مراحل الكلاسك وتكوين الأعضاء لجميع التوليفات الهرمونية المستعملة، زعزعة المتك
بطرمو معقم على سطح موراستيج واسكيت المغنيسيوم ملء بالنبيذ الأدنى (0.0، 0.3 و 5.0 و 7.0 ملجم/لتر)
بالتداخل مع (2 و 4- دارو كلورفينوكسي أستيك أسد) (0.0 و 0.01 و 0.05 ملجم/لتر) لإنتاج الكلاسك وتكوين
الأعضاء، بينما النتائج أن سطح موراستيج واسكيت المتناثر مختلف التوليفات من بنزال أدنين و2 و4- دارو
كلورفينوكسي أستيك أسد قد ساعد على استجابة الكلاسك بعمل تراوح بين 20-60%، وعززت التركيز عالية من
منظومات النمو وجدت نسبة عالية من الكلاسك مع تفعيل استجابة تكون النباتات عن طريق التكوين غير المباشر ليصل إلى
أعلى معدل لعدد النباتات بلغ 4.5 و 4.7 عند التركيز (5.0 ملجم/لتر بنزال أدنين + 0.5 ملجم/لتر بنزال أدنين)
دارو كلورفينوكسي أستيك أسد (7.0 ملجم/لتر بنزال أدنين + 0.05 ملجم/لتر بنزال أدنين + 0.05 ملجم/لتر بنزال أدنين +
دارو كلورفينوكسي أستيك أسد و7.0 ملجم/لتر بنزال أدنين + 0.05 ملجم/لتر بنزال أدنين + 0.05 ملجم/لتر بنزال أدنين +
دارو كلورفينوكسي أستيك أسد و7.0 ملجم/لتر بنزال أدنين + 0.05 ملجم/لتر بنزال أدنين + 0.05 ملجم/لتر بنزال أدنين +
دارو كلورفينوكسي أستيك أسد + 0.05 ملجم/لتر بنزال أدنين + 0.05 ملجم/لتر بنزال أدنين + 0.05 ملجم/لتر بنزال أدنين + 0.05 ملجم/لتر بنزال أدنين)
(استيك أسد على التوالي).

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