ALTERNATION USING BACTERIAL SUPERNATANTS IN DATE PALM MICROPROPAGATION

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ABSTRACT

The present investigation was aimed to use phytohormones produced by some bacterial strains like Azospirillum brasilense, Azotobacter chroococcum, Bacillus megaterium and Klebsiella pneumoniae instead artificial hormones as a protocol in date palm micropropagation through somatic embryogenesis induced from shoot tip or leaf primordial explants, which were isolated from offshoot apex. In order to test the possibility to use supernatant of the strains, which are auxin over producers and/or cytokinin over producers in culture medium composition according to development stages for maximizing yield and to study the effect on each stage, i.e., callus initiation, embryo formation, shooting and rooting stages. In general, results showed that Klebsiella and Azotobacter have an auxin effect but Azospirillum and Bacillus were more related to cytokinin nature.

INTRODUCTION

Several soil microorganisms are capable of producing physiologically active quantities of plant growth regulators (PGRs), which may have a pronounced effect on plant growth and development. A diverse array of bacteria, including species of Azospirillum, Azotobacter, Bacillus and Klebsiella have mechanisms by which these rhizobacteria enhance plant growth through producing substances such as auxins, GA3 and cytokinins (phytohormones) and mineralize organic phosphorus and enhance minerals uptake (Grayston et al. 1990 and Joo et al. 2004). So that, using these natural substances in vitro in culturing economic plants become necessary to overcome increasing artificial substances and that is of variable dangerous on healthy.

Because, date palm is the most important fruit crop planted and as ornamental plants in the Middle East and Arabian lands (FAO, 1984 and Moursy and Saker, 1996). Date palm is a monocot and dioecious fruit tree, which are difficult to study due to their long life. Therefore, in vitro techniques appear to be promising tools to study palm growth and development compared to field and greenhouse experimentation (Tisserat, 1983).

The concentration and type of hormones especially, auxins and cytokinins are considered the most critical factors in stimulating callus production and subsequent embryogenesis (Reynolds and Murashige, 1979 and Tisserat, 1979).

Generally, Auxins play a role in many development processes, including cell elongation, swelling of tissue, apical dominance, adventitious root formation and somatic embryogenesis (Trigiano and Gray, 2000). In
addition, cytokinins are required for inducing cell division, callus formation, cell extension, shoot proliferation and shoot morphogenesis (Pan and Van Staden, 1998) Naphthaleneacetic acid (NAA) and kinetin at levels from 1-10 mg/L induced callus growth (Khan et al., 1983). An embryo can be defined as the earliest recognizable multicellular stage of an individual that occurs before it has developed the structures or organs characteristic of a given species (Trigiano and Gray, 2000). Somatic embryogenesis is a process by which somatic (non-gametic) cells undergo differentiation to form a bipolar structure containing both root and shoot axes. These somatic embryos are similar to zygotic embryos and can mature and germinate (Smith, 2000).

Abdel-Hamid et al. (2001) found that in date palm micropropagation, the highest rate of shoot multiplication induced from shoot tip explants was obtained through callus cultured on medium containing 3 mg/L 2iP. Also, Zaid (2003) showed that in date palm micropropagation that its culture medium was supplemented with 0.5 mg/L kinetin was more effective than that supplemented with 0.5 mg/L 2iP on stimulating the produced shoots number and shoot length.

MATERIALS AND METHODS

This work was carried out in the Central Laboratory for Research and Development of Date Palm, Agricultural Research Center (ARC), Ministry of Agriculture, Giza, Egypt and Plant Biotechnology Department, Genetic Engineering and Biotechnology Research Institute (GEBRI), Minufiya University during the period 2003-2006.

Source of bacterial strains:

Representative bacterial strains used throughout this study were Klebsiella pneumoniae and Azospirillum brasilense, which were isolated through a previous study carried out at the Environmental Studies and Research Unit (ESRU) Department of Agricultural Microbiology, Faculty of Agriculture, Cairo University, Giza by Farrag(2000). The other two strains Bacillus megaterium and Azotobacter chrooccum were kindly obtained from the Culture Collection of Microbiological Resource Center, Faculty of Agriculture, Ain Shams University, Cairo, Egypt.

Purification and maintenance of the isolates:

The selected strains were purified by streaking on agar plates containing the proper selective medium for each bacterium type as the following:

I. Nitrogen deficient medium (NFDM) for Klebsiella (Dixon et al., 1980)
II. Semi–solid malate medium for Azospirillum (Dobereiner and Day 1976)
III. Nitrogen – deficient medium of Ashby for Azotobacter (Hegazi and Niemela, 1976)
IV. Bacillus medium (Allen, 1959).

which were surface inoculated with single strain and incubated at 28±2°C for 48hr to 7 days (5-7 times or more) until pure colonies were obtained.

Characteristics of colonies for each tested strains developed on agar plates of every culture medium were carefully examined, which was having different colors.
Maintenance of the selected strains was carried out by sub-culturing on several selective agar media as slants periodically, which were then kept in refrigerator at (4°C). Strains were separately grown in liquid medium for preparation batch cultures by inoculating 10 mL of 10^5-10^6 cells mL^-1 in selective culture media and incubated in a rotary shaker at 100 rpm and temperature was controlled at 30°C to reach a population density of >10^7 cfu mL^-1 (colony forming/ unit). PH was adjusted to 7.0.

**Plant material:**

Selected young offshoots, 2-3 years old, were grown in El- Wahaat El- Baharia in Egypt. Offshoots were carefully separated from the mother date palm trees (*phoenix dactylifera* cv. Sewi). The selected offshoots were 6-9 Kg in weight, about 1-1.5 m in length and 20-30 cm in diameter.

**Isolation and sterilization of explants:**

The selected young offshoots were immediately transferred to the laboratory. Two types of explants were separated from the offshoots, i.e., shoot tip and leaf primordial. For excising both types of explants, the green leaves of the offshoots were carefully removed by using sharp knife. The separation of leaves was carried out starting from the base and acropetaly continued upwards until the white soft leaves, surrounding the shoot tip appeared. However, cork and other remaining tissues of the offshoot were discarded.

The plant materials including the shoot tip region (10 – 12 cm in length and 5 cm in diameter) were sterilized as follows:

- Plant materials were washed several times with liquid soup and water, then placed under running tap water for 2 hours. The plant materials were then surface sterilized by soaking for 3 min in ethyl alcohol (70%).
- Again, plant materials were surface sterilized by immersing in mercuric chloride solution (0.1%) containing two drops of tween 20 for 5 min. After removing all affected tissues, the remaining part was sterilized by soaking for 20 min in commercial Clorox 70 % (v/v) (sodium hypochlorite 5.25%) containing two drops of tween 20 as a surfactant. It was then rinsed three times with sterilized distilled water. Additional injured leaves were removed from the shoot tip material. The shoot tip material was again rinsed with sterilized distilled water and soaked for 20 min. in filter sterilized solutions consisted of citric acid (150 mg/L) and ascorbic acid (100 mg/L) as an anti-oxidant. The shoot tip materials were again soaked for 20 min. in clorox 50% (v/v) (sodium hypochlorite 5.25 %) containing two drops of tween 20.
- Under aseptic conditions, all affected primordial leaves were removed and shoot tip materials were again rinsed three times with autoclaved distilled water followed by soaking in filter sterilized solution containing citric acid (150 mg/L) and ascorbic acid (100 mg/L).
- The shoot tip regions including the shoot apex plus the associated primordial leaves were used. Shoot apical meristem plus few primordial leaves (4-5 leaves) were excised together, and would be considered as shoot tip explant. The shoot tip explants was about 1 cm in length and 0.5-0.8 cm in diameter. The shoot tip was divided longitudinally with a sharp cuter into 4
equal quarters. The leaf primordial explants were taken from the rest primordial leaves surrounding the meristem tip.

**Culture medium:**

The MS basal medium containing macro and microelements as well as vitamins (Murashige and Skoog, 1962) was used in this study.

The control MS basal medium was supplemented with 200 mg/L glutamine 100 mg/L myo-inositol, 3% sucrose and 0.6% agar. Medium was adjusted to pH 5.8 ± 0.1.

The medium was distributed into the culture jars (250 mL); each jar contains 25 ml of MS medium. Jars were capped with polypropylene closure and autoclaved at 121°C and 1.1 kg/cm² for 20 minutes. The jars were transferred to the culture cabinet to get cool. The different explants were cultured into MS medium supplemented with bacterial supernatants to obtain cultured medium containing all components as natural medium to study its effect on the yield of each studied stage.

**Callus initiation stage:**

Twenty-five offshoots were considered as a source of explants. The control medium was MS culture medium 10 mg/L 2,4-D 3 mg/L 2iP, 30 g/L sucrose and all B. Ss (bacterial supernatants) were added to the MS medium without growth regulators at 150 mL/L concentration as recommended by Farrag (2000) in treatments A, a, B, b, C, c and D, d as the following:

**A)** Azotobacter chroococcum + tryptophane (as a producer for 41µg/mL GA3 → 6.2ML/150mL),

**a)** Azoto. Cultured in free trypto. Medium (non significant production),

**B)** Klebsiella pneumoniae + tryptophane (IAA over producer 47µg/mL → 7mL/150mL, also GA3 66µg/mL → 10mL/150mL),

**b)** Kleb. Cultured in free trypto. Medium (non significant production),

**C)** Azospirillum brasilense + tryptophane (cytokinin over producer 5.4µg/mL → 0.81mL/150mL),

**c)** Azosp. Cultured in free trypto. medium (non significant production),

**D)** Bacillus megaterium + tryptophane (non significant production),

**d)** B. megaterium Cultured in free trypto. medium (cytokinin producer 5µg/mL → 0.75mL/150 mL),

These bacterial strains ascertainment in the considered treatments were chosen according to their ability to produce phytohormones from previous studies of Farrag et. al. (2007). According to Zaid and Tisserat (1983) cultures were transferred to fresh media every 6 weeks, and incubated in complete darkness for 24 weeks.

**The data recorded as follows:**

1. Callus formation degree.
2. Browning degree.
3. Swelling degree at the end of each subculture (4 subculture).

The compact callus, which formed from the previous culture were placed on media containing 10 mg/L 2,4-D and 3 mg/L 2iP to produced friable callus (control). Data were calculated visually as scores (Pottino, 1981) from 1- 4 as the following:

Negative results (-) = 1
Below average results (+) = 2
Average results (++) = 3
Good results ( +++ ) = 4

**Embryogenic Callus Formation:**

Friable callus was transferred to the following media:

- MS + free growth regulators (control) was (0).
- MS + 150 mL/L B.Ss. A, a, B, b, C, c and D, d.

For embryogenesis, each treatment consisted of 3 replicates and each replicate contains about 1x1 cm friable callus. Cultures were incubated in total darkness for 3 months. Data were recorded as growth vigor and number of embryo.

**Shoot formation stage:**

Individual embryos were separated from previous stage and cultured on the same media. The data of growth and development were recorded as:
1. Number of shoots / plant.
2. Shoot length (cm).

**Root formation stage**

Individual shoots were used as explants in this stage and they were cultured in MS supplemented with NAA (control) 0.1 mg/L and the same treatments. All culture media of each treatment were distributed into culture tubes. The culture tubes were generally placed in light condition for 16 h illumination of 3000 lux light intensity (white fluorescent tubes) at 27 ± 2°C. The data of this stage were recorded as:
1. Number of roots/plant.
2. Root length (cm).

**Statistical analysis:**

Experiments were arranged in factorial complete design and date were statistically analyzed according to method described by Snedecor and Cochran (1972).

**RESULTS AND DISCUSSION**

**callus initiation:**

Data presented in Table (1) showed callus formation, browning and swelling degree as affected by adding bacterial supernatants. Results revealed that the highest score value of callus formation degree was observed with Klebsiella supernatant in the presence of tryptophan (B) that is (5.3) followed by Azotobacter supernatant(A), Azospirillium supernatant(C) with tryptophan and control(0), which recorded the same score (3). These treatments were the best in the swelling (3.25) except for the control (2.25). On the other hand, treatment (C) was the lowest in the browning degree (0.5), in the opposite, the highest value of browning degree was (2.5) in (A) and (D) treatments but the lowest values of callus degree was in Azospirillium supernatant free tryptophan (c) and Bacillus supernatant free tryptophan (d) which their score were (1). In this respect, Chung and Tzeng (2004) showed that the production of IAA in culture media was solely dependent on the presence of tryptophan. Moreover, One et al. (2005) stated...
that varying the concentration of tryptophan in batch experiments has an effect on both growth of *Azospirillum* and IAA synthesis. This results are in agreement with the obtained results which, in the present study, which gave positive effect by using bacterial supernatants from medium containing tryptophan as precursor for production of auxin.

Table(1) . Effect of bacterial supernatant added to MS medium on callus development and differentiation of date palm(*Phoenix dactylifera cv.Sewi*) after 18 weeks from culturing

<table>
<thead>
<tr>
<th>Treatments</th>
<th>Growth vigor</th>
<th>Number of embryo</th>
</tr>
</thead>
<tbody>
<tr>
<td>Azotobacter chroococcum (A)</td>
<td>3.0</td>
<td>15</td>
</tr>
<tr>
<td>Azoto. free tryptophan (a)</td>
<td>2.5</td>
<td>10</td>
</tr>
<tr>
<td>Klebsiella pneumoniae (B)</td>
<td>3.5</td>
<td>20</td>
</tr>
<tr>
<td>Klebsiella free tryptophan (b)</td>
<td>1.5</td>
<td>6.0</td>
</tr>
<tr>
<td>Azospirillium brasilense (C)</td>
<td>4.0</td>
<td>30</td>
</tr>
<tr>
<td>Azosp. free tryptophan (c)</td>
<td>3.0</td>
<td>17</td>
</tr>
<tr>
<td>Bacillus megaterium (D)</td>
<td>3.5</td>
<td>26</td>
</tr>
<tr>
<td>Bacillus free tryptophan (d)</td>
<td>4.0</td>
<td>25</td>
</tr>
<tr>
<td>Control</td>
<td>3.0</td>
<td>18</td>
</tr>
<tr>
<td><strong>L.S.D. at 5%</strong></td>
<td>1.01</td>
<td>4.4</td>
</tr>
</tbody>
</table>

Moreover, Reuveni and Kipnis (1974) showed that auxins were found to be the most critical component in the media for callus production and Tisserat (1982) mentioned that (MS) medium supplemented with high level of auxin resulted in high percentage of callus. In this concern, Brackpool (1988) reported that in date palms micropropagation, high concentrations of the artificial auxins naphthaleneacetic acid (NAA) and 2,4-dichloroenoxyacetic acid (2,4-D) are needed to induce callus formation. While (Trigiano and Gray 2000) reported that auxins play a role in many development processes, including cell elongation, swelling of tissue, apical dominance, adventitious root formation and somatic embryogenesis.

**Embryogenesis:**

The effect of bacterial supernatants on callus development is shown in Table (2). Results revealed that the greatest number of individual embryos was (30) in MS medium supplemented with *Azospirillum* supernatant with tryptophan (C) followed by (26) formed by using *Bacillus* supernatant with tryptophan (D)and (25) in *Bacillus* supernatant from medium free tryptophan (d) this results may due to the presence of cytokinin in this supernatants. In addition, the best growth vigor was due to (C) and (D) which recorded a score (4) that may be owed to presence other substances like gibberellins and auxins. These results are in harmony with Gonzalez - Lopez et al., (1986) who observed that the culture supernatant of *A. chroococcum & A. vinelandii* contained at least three gibberellins-like substances. Mahmoud et al. (1984) investigated the ability of different isolates belonging to genera *Bacillus and Azotobacter* to synthesize plant growth substances. The supernatant of the bacterial cultures showed the presence
of a compound, which correspond to indole acetic acid together with unidentified derivatives. These organisms also secreted gibberellins.

Table (2). Effect of bacterial supernatant on callus formation of date palm (Phoenix dactylifera cv. Sewi) after 18 weeks from culturing (three subculture)

<table>
<thead>
<tr>
<th>Treatments</th>
<th>Callus Formation</th>
<th>Browning</th>
<th>Swelling</th>
</tr>
</thead>
<tbody>
<tr>
<td>Azotobacter chrooccum (A)</td>
<td>3.0</td>
<td>2.5</td>
<td>3.25</td>
</tr>
<tr>
<td>Azoto. free tryptophan (a)</td>
<td>2.0</td>
<td>1.5</td>
<td>2.0</td>
</tr>
<tr>
<td>Klebsiella pneumoniae (B)</td>
<td>3.5</td>
<td>1.0</td>
<td>3.25</td>
</tr>
<tr>
<td>Klebsiella free tryptophan (b)</td>
<td>1.5</td>
<td>1.0</td>
<td>1.25</td>
</tr>
<tr>
<td>Azospirillium brasiliense (C)</td>
<td>3.0</td>
<td>0.5</td>
<td>3.0</td>
</tr>
<tr>
<td>Azosp. free tryptophan (c)</td>
<td>1.0</td>
<td>1.0</td>
<td>2.0</td>
</tr>
<tr>
<td>Bacillus megaterium (D)</td>
<td>2.5</td>
<td>2.5</td>
<td>3.25</td>
</tr>
<tr>
<td>Bacillus free tryptophan (d)</td>
<td>1.0</td>
<td>2.0</td>
<td>2.25</td>
</tr>
<tr>
<td>Control</td>
<td>3.0</td>
<td>2.0</td>
<td>2.75</td>
</tr>
<tr>
<td>L.S.D. at 5%</td>
<td>1.2</td>
<td>0.12</td>
<td>0.93</td>
</tr>
</tbody>
</table>

At the same time, Omar (1988) investigated that ovular callus was noticed after 6 weeks on a medium containing 10 mg/L 2,4-D and 2 mg/L kinetin. While, Sharon et al. (1998) mentioned that white friable callus was initiated from leaf primordia of date palm cv. Yakubi cultured on modified MS medium supplemented with 50 mg/L 2,4-D, 1 mg/L kinetin and 0.5 mg/L 2iP and Ibrahim (1999) showed that, embryogenic callus was obtained from shoot tips and leaf primordial explants cultured on modified MS medium containing either 10 mg/L NOA, 5 mg/L 2,4-D, 1 mg/L kinetin and 1 mg/L 2iP or 100 mg/L 2,4-D, 3 mg/L kinetin and 3 mg/L 2iP.

Shoot formation:

Results in Table (3) showed that best growth vigor was in Klebsiella supernatant in the presence of tryptophan (B) that is (4) and the highest shoot was (7 cm) in the same treatment followed by Azotobacter supernatant with tryptophan (A), which gave growth vigor (3.5) and shoot length (6.5 cm). The largest number of shoot/plant was (5) due to culturing in MS medium containing Azospirillium supernatant with tryptophan (C) these results are in agreement with those found by Kapulink et al. (1985) who found that Azospirillium has the ability to produce plant growth promoting substances such as auxins and cytokinins. El-Khawas et al. (1996) found that Azospirillium and other groups of PGPR were produce and release a broad spectra of plant growth regulators (auxins, several gibberrellins and cytokinins). Garcia de Salamone et al. (2001) stated that five plant growth promoting rhizobacteria (PGPR) strains produced the cytokinin dihydrozeatin riboside (DHZR) in pure culture. In addition to, Saker et al. (1998) demonstrated that in in vitro tissue culture of date palm, only 2iP containing media was found to be satisfactory for shoot induction after a phase of callus formation. The highest percentage of shoot proliferation occurred when embryos were cultured on MS medium supplemented with 3 mg/L 2iP, 0.1 mg/L NAA and 3 g/L charcoal.
Table (3). Effect of bacterial supernatant added to MS medium on shoot formation of date palm (*Phoenix dactylifera* cv. Sewi) after 18 weeks from culturing

<table>
<thead>
<tr>
<th>Treatments</th>
<th>Growth vigor</th>
<th>Shoot number</th>
<th>Shoot length</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>Azotobacter chrooccum</em> (A)</td>
<td>3.5</td>
<td>1.0</td>
<td>6.5</td>
</tr>
<tr>
<td>Azoto. free tryptophan (a)</td>
<td>1.0</td>
<td>2.0</td>
<td>4.5</td>
</tr>
<tr>
<td><em>Klebsiella pneumoniae</em> (B)</td>
<td>4.0</td>
<td>2.0</td>
<td>7.0</td>
</tr>
<tr>
<td><em>Klebsiella</em> free tryptophan (b)</td>
<td>2.0</td>
<td>1.0</td>
<td>3.5</td>
</tr>
<tr>
<td><em>Azospirillum brasiliense</em> (C)</td>
<td>2.5</td>
<td>5.0</td>
<td>5.5</td>
</tr>
<tr>
<td>Azosp. free tryptophan (c)</td>
<td>1.5</td>
<td>2.0</td>
<td>4.0</td>
</tr>
<tr>
<td><em>Bacillus megaterium</em> (D)</td>
<td>2.5</td>
<td>3.0</td>
<td>6.0</td>
</tr>
<tr>
<td>Bacillus free tryptophan (d)</td>
<td>3.0</td>
<td>4.0</td>
<td>4.0</td>
</tr>
<tr>
<td>Control</td>
<td>3.0</td>
<td>3.0</td>
<td>4.5</td>
</tr>
<tr>
<td>L.S.D. at 5%</td>
<td>0.95</td>
<td>1.35</td>
<td>1.68</td>
</tr>
</tbody>
</table>

Moreover Bekheet and Saker (1998) reported that in date palm micropropagation, the presence of cytokinins such as BA and 2iP in the culture media was considered as an important factor for bud differentiation. Shoot induction was confined to media containing 4 mg/L BA, 4 mg/L 2iP and 0.5 mg/L NAA. Also, Zaid (2003) showed that in date palm micropropagation the culture medium supplemented with 0.5 mg/l kinetine was more effective than 0.5 mg/l 2iP on stimulating the produced shoots number and shoot length.

**Rooting formation:**

Results in Table (4) revealed that, significant differences were observed among treatments, while *Klebsiella* supernatant in the presence of tryptophan (B) gave the best growth vigor value (4), the greatest root number (6) per plant and the tallest root (6.5cm) followed by *Azotobacter* supernatant with tryptophan (A), which recorded growth vigor value (3.5), root number (4) per plant and root length (5cm). These results and all the previous ones are in the same line with Hafeez et al. (2004) who conducted different experiments to determine the growth promotion activity of various rhizobacteria. The results showed that the root growth was depending on the ability of *Azotobacter* strains for IAA production.

Table (4). Effect of bacterial supernatant added to MS medium on root formation of date palm (*Phoenix dactylifera* cv. Sewi) after 18 weeks from culturing

<table>
<thead>
<tr>
<th>Treatments</th>
<th>Growth vigor</th>
<th>Root number</th>
<th>Root length</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>Azotobacter chrooccum</em> (A)</td>
<td>3.5</td>
<td>4.0</td>
<td>5.0</td>
</tr>
<tr>
<td>Azoto. free tryptophan (a)</td>
<td>1.5</td>
<td>2.0</td>
<td>3.5</td>
</tr>
<tr>
<td><em>Klebsiella pneumoniae</em> (B)</td>
<td>4.0</td>
<td>6.0</td>
<td>6.5</td>
</tr>
<tr>
<td><em>Klebsiella</em> free tryptophan (b)</td>
<td>2.5</td>
<td>1.0</td>
<td>2.0</td>
</tr>
<tr>
<td><em>Azospirillum brasiliense</em> (C)</td>
<td>3.5</td>
<td>3.0</td>
<td>4.0</td>
</tr>
<tr>
<td>Azosp. free tryptophan (c)</td>
<td>3.0</td>
<td>1.0</td>
<td>2.5</td>
</tr>
<tr>
<td><em>Bacillus megaterium</em> (D)</td>
<td>1.0</td>
<td>3.0</td>
<td>4.5</td>
</tr>
<tr>
<td>Bacillus free tryptophan (d)</td>
<td>3.0</td>
<td>1.0</td>
<td>2.0</td>
</tr>
</tbody>
</table>

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Recently, Abd El-Kader(2007) cited that an isolate belongs to genus *Klebsiella* was IAA over producers. While, (Thorpe, 1981) reported that Auxin alone or with a very low concentration of cytokinin is important for the induction of root primordial in plant tissue culture. Bekheet and Saker (1998) reported that in date palm tissue culture, rooting of proliferated shoots was achieved upon supplementation of MS culture medium with 1 mg/L NAA. Moreover, Gadalla (2003) demonstrated that in mass propagation of dry cultivars of date palm, the highest significant value of rooting percentage was observed when MS culture medium was supplemented with 3 mg/L of either NAA or IBA.

**REFERENCE**


Farrag, Hala, M.A. et al.


### تعاقب استخدام الراشحات البكتيرية في زراعة أنسجة نخيل البلح

هالة محمد أنور فرج – وداد النتهائي عويضه – عادل ابادنور جرجس – عاطف محمد أبراهيم – أبراهيم عبد المنعم عبد الودود اليا – إبراهيم عبد العطوسد

المعمل المركزى للابحاث - جامعة شمس – معهد الهندسة الزراعية - جامعة المنوفية

* بكلية الزراعة - جامعة عين شمس

** معهد الهندسة الزراعية و التكنولوجيا الحيويةـ جامعة المنوفية

يدفع هذا البحث إلى استخدام الهرمونات المنتجة طبيعيا بواسطة بعض السلالات البكتيرية مثل *Azospirillum brasilense*, *Azotobacter chroococcum*, *Bacillus megaterium* and *Klebsiella pneumoniae* بدلا من منظمات النمو المصنعة في زراعة أنسجة نخيل البلح عن طريق الأنبوب البكترى الذي قامت به إعداد الأوراق المصبوغة من مسارات النخيل في تلك السلالات. واستنادا لذلك لدراسة هذه السلالات قام الباحثان باستنباط النتائج التالية: 

- من خلال مرحلة زراعة الأوراق الساقطة من نخيل البلح، و نخيل الباي، و نخيل الكالس، و نخيل الأوراق الساقطة تم قياس النتائج لتقييم الأموث النباتية لتمكين النبات من إنتاج النباتات البالغة، و النباتات الساقطة، و النباتات التي تميزها في تأثيرها

وردر أوكسسين قوي بينما تتمي في تأثيرها *Bacillus megaterium*, *Azospirillum brasilense* أكثر إلى السيتوكينين.