The effect of cytokinins on \textit{in vitro} shoot length and multiplication of \textit{Hymenocallis littoralis}

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This study was performed to determine the effects of different cytokinins at various concentrations on the \textit{in vitro} shoot formation of \textit{Hymenocallis littoralis}. The bulb scales of \textit{H. littoralis} were used as explants to establish the cultures. The explants were grown on semi-solid Murashige and Skoog (MS) medium supplemented with 2iP (2-isopentenyladenine), TDZ (thidiazuron) and Zeatin respectively at six different concentrations (2.25, 4.50, 9.00, 13.50, 18.00, and 22.50 µM). Two subcultures were performed at 30 days interval after the initial \textit{in vitro} culture establishments. It was found that 2iP and concentration 13.50 µM respectively was the best for shoot elongation which was measured in length (cm). However, the three cytokinins and the six tested concentrations were recorded to have no significant difference in terms of shoot multiplication. Highest total chlorophyll content was observed in shoots grown on semi-solid MS medium supplemented with 2.25 µM of Zeatin.

Key words: \textit{Hymenocallis littoralis}, \textit{in vitro}, cytokinins.

INTRODUCTION

\textit{Hymenocallis littoralis} (Jacq.) Salisb. is a bulbous, herbaceous plant from the family Amaryllidaceae. It is also known as \textit{Hymenocallis panamensis} Lindl., \textit{Pancratium americanum} Mill., \textit{Pancratium littorale} Jacq. (Ioset et al., 2001; Ocampo and Balick, 2009). The plant is distributed by the sea and in swamps in tropical, subtropical, and temperate regions throughout the world (Ji and Meerow, 1985).

Other than its application as an ornamental plant, \textit{H. littoralis} was discovered to exhibit various medicinal properties (Backhaus et al., 1992). Indeed, many plants from the same family (Amaryllidaceae) as \textit{H. littoralis} had served as cures of various illnesses for decades. In particular, the bulbs of \textit{Hymenocallis Americana} were made into poultice for varicose veins, sores and swellings. \textit{Hymenocallis amancaes} is commonly employed as an ornamental plant and used in cosmetic preparations.

Applying the mixture of oil and crushed bulbs onto the face has been found to treat freckles and blemishes (Ocampo and Balick, 2009). Hence, it is obvious that the various medicinal applications of plants in the genus \textit{Hymenocallis} come primarily from their bulbs. In fact, almost all of the studies on \textit{H. littoralis} were done on its seemingly most valuable part, the bulb (Ioset et al., 2001). Throughout the history of \textit{H. littoralis}, several alkaloids have been discovered from its bulb. The first alkaloid, lycorine was proven to have antineoplastic, cytotoxic and antiviral properties (Ioset et al., 2001). Another compound, pancratistatin has been proven to be effective against U.S. National Cancer Institute’s panel of 60 human cancer cell lines demonstrating greatest effectiveness against melanoma, brain, colon, lung and renal cancers (Backhaus et al., 1992; Pettit et al., 1993). Littoraline alkaloid was then discovered with inhibitory activity on HIV reverse transcriptase (Lin et al., 1995). As the compounds isolated from \textit{H. littoralis} have been proven to demonstrate various medicinal properties (Lin et al., 1995; Idso et al., 2000), it acts as the driving force behind the current study to micropropagate the plant using plant tissue culture techniques.

Micropropagation is the \textit{in vitro} asexual multiplication of
genetically identical individuals using plant tissue culture technology. Cytokinin is one of the plant hormones crucial for plant growth and development and is known to promote cell division. Various types of cytokinins can also stimulate lateral bud growth and thus causing multiple shoot formation by breaking shoot apical dominance (Trigiano and Gray, 2005). Cytokinins are categorically divided into naturally-occurring and synthetic. Out of the three cytokinins used in the present study, 2iP (2-isopentenyl adenine) and Zeatin are naturally-occurring while TDZ (thidiazuron) is synthetic and highly active (Schmulling, 2004). The specific objectives of this study are (i) to determine the growth of *H. littoralis* in semi-solid MS medium supplemented with three types of cytokinins namely 2iP (2-isopentenyl adenine), TDZ (thidiazuron) and Zeatin at six different concentrations (2.25, 4.50, 9.00, 13.50, 18.00, and 22.50 µM); (ii) to select the most effective cytokinin and its concentration for *in vitro* micropropagation of *H. littoralis* in terms of shoot growth (length) and shoot multiplication, and (iii) to select the best cytokinin and its concentration, in terms of highest total chlorophyll content, for the *in vitro* micropropagation of *H. littoralis*.

**MATERIALS AND METHODS**

**Bulb preparation**

The explant used in this study was the bulb of *H. littoralis*. The plants were ordered from a nursery. In order to obtain the bulb, the outer black polyethylene cover and the soil covering the bulb were removed. Then, without damaging the bulb structure, the leaves and roots of the bulb were carefully cut off.

**Bulb surface sterilization and dissection**

The brown coating of the bulb was peeled off, revealing a whole “clean” white bulb. It was then placed in a beaker and washed with tap water added with 2 to 3 drops of Teepol for an hour to thoroughly remove all of the attached dirt, debris, and soil.

The bulb was then dissected into four parts equally. Before further dissection, surface sterilization procedures were conducted on the divided tiller shoots. Firstly, they were immersed in 70% ethanol for 2 min and transferred to 50% sodium hypochlorite solution (Clorox 5.5%) for 20 min. Upon completion of the sodium hypochlorite wash, subsequent steps were carried out in aseptic conditions in the laminar flow hood. A second immersion in 70% ethanol for another 2 min is followed by rinsing the tissues three times in distilled water. The tissues were blot dried on a filter paper prior transferring under aseptic condition.

**Culture establishment**

To initiate a culture, the basal vascular portion of explant was planted into the semi-solid agar medium. At every concentration out of the six concentrations (2.25, 4.50, 9.00, 13.50, 18.00, and 22.50 µM) for 2iP, TDZ and Zeatin, four explants were planted on the semi-solid MS medium (Murashige and Skoog, 1962) in a single culture jar. It was then repeated for three times which made up to (4 explants × 4 jars) a total of 16 replicates. All of the cultures were placed on a white wooden rack in the culture room at 25 ± 2 °C with continuous cool white fluorescent light. The cultures were grown for 30 days which was the interval for a subculture. Two subcultures had been performed in this study.

**Shoot length and multiplication**

After a period of 30 days from the initial culture establishment date, all of the plantlets were taken out, one culture jar after another, and placed on a sterilized Petri dish under which a graph paper was adhered to. The morphology of all the plantlets was carefully observed and compared to each other. Then, the number of shoots and their respective lengths were measured and recorded. All of the procedures were done under strict aseptic techniques and conditions as the contamination of any culture would require it to be discarded. After the measurements were completed, all of the plantlets were transferred to a culture jar containing new semi-solid MS medium according to their respective cytokinin type and concentration. From here, the first subculture was done and the steps were repeated after another 30 days for the second subculture.

**Chlorophyll content**

When the second subculture was completed, the plantlets were used to determine their total chlorophyll content using Harborne method (Harborne, 1973).

**RESULTS AND DISCUSSION**

**Shoot length**

*H. littoralis* shoots start to form within one week from the day of culture establishment. The success rate of shoot formation was at an average of 93.4%.

Figure 1 shows mean shoot length (cm) against the six concentrations used for 2iP, TDZ and Zeatin after the first and the second subcultures. The cytokinin concentration of 13.50 µM was found to be the best for stimulating shoot length followed by 18.00, 22.50, 9.00, 2.25 and 4.50 µM. Nevertheless, based on Dunnett’s test (2-sided) of multiple mean comparisons, only 13.50 and 18.00 µM were significantly different from other concentrations tested in terms of mean shoot length. While the mean shoot length increased from 2.25 µM up to 13.50 µM (the highest), it started to decrease when the cytokinin concentration was further increased. This phenomenon is actually similar to a research on micropropagation of different banana cultivars using scalps where the shoot length was increased with higher BAP (benzyl amino purine) level until 22.2 µM after which the shoot length also began to fall (Shirani et al., 2010).

Figure 2 shows mean shoot length (cm) against the type of cytokinins used in the study; 2iP, TDZ and Zeatin. Both figures recorded 2iP to produce the highest mean shoot length followed by TDZ and Zeatin. After the first subculture, both TDZ and Zeatin were significantly different from 2iP. Meanwhile after the second subculture, only Zeatin was significantly different from 2iP. This observation is similar to a research on the regeneration of...
Figure 1. Mean shoot length produced by *H. littoralis* under different concentrations of various cytokinins (2iP, TDZ and Zeatin) on semi-solid MS medium. (A) After 30 days of cultivation. (B) After 60 days of cultivation. Mean with the same alphabet on top of every bar is not significantly different [Dunnett’s test (2-sided), α = 0.05].

Figure 2. Mean shoot length produced by *H. littoralis* under different cytokinin treatments of various concentrations (2.25, 4.50, 9.00, 13.50, 18.00, and 22.50 µM) on semi-solid MS medium. (A) After 30 days of cultivation. (B) After 60 days of cultivation. Mean with the same alphabet on top of every bar is not significantly different [Dunnett’s test (2-sided), α = 0.05].

Sugarcane using various cytokinins (2iP, Benzyl adenine [BA], Kinetin, TDZ, and Zeatin) where 2iP had been reported to be the most effective for shoot elongation (Chengarayan and Gallo-Meager, 2001). The low mean shoot length of TDZ in the first subculture was because TDZ had been found to reduce shoot elongation (Bates et al., 1992; Murthy et al., 1998). However, if the culturing period was increased, the shoots would eventually
A cytokinins demonstrated similar descending pattern, 3A). Dunnett’s test (2-sided) of multiple mean comparisons indicated that 9.00 and 22.50 μM were significantly different from 2.25 μM. Since explants usually require a period of adaptation to in vitro environment, this may be the cause of significantly lower mean shoot number at concentrations 9.00 and 22.50 μM from 2.25 μM after the first subculture (Carelli and Echeverrigaray, 2002). However, after the second subculture (Figure 3B), statistical analysis revealed that there was no significant difference between the six concentrations tested. This may be because the cytokinin concentrations used were not high enough to effectively stimulate rapid shoot multiplication. Previous studies involving other plants noted that relatively high concentrations of cytokinins should be present to observe high multiplication rates. In the case of Gerbera cultivars multiplication, it was reported that the supplementation of 10 mg/L (44.40 μM) benzyl adenine (BA), twice the maximum cytokinin concentration used in this study, increased adventitious bud formations in all cultivars during the initial and the multiplication stages (Rael and Tönis, 2001).

The relationship between mean shoot number and the cytokinins used is shown in Figure 4. There was no significant difference in terms of mean shoot number among the three cytokinins. Further statistical analysis indicated that the effect of the three cytokinins on mean shoot number was not significantly different between the first and the second subculture periods. Cytokinin alone may not be able to significantly induce multiple shoot formation in H. littoralis. Some studies demonstrated that the addition of auxin in synergy with cytokinin promotes better shoot proliferation (Chengalrayan and Gallo-Meagher, 2001). Another suggestion would be to supply combinations of cytokinins. In a study concerning Bauhinia vahlii, a leguminous plant, the combination of TDZ and Kinetin recorded significant increase in shoot numbers lasting for as long as 4 subculture periods (Bhatt and Dhar, 2000).

**Chlorophyll content**

For TDZ and Zeatin, the overall pattern of total chlorophyll content was seen to be descending from the lowest concentration (2.25 μM) to the highest concentration (22.50 μM) (Figure 5). This was translated as lower cytokinin concentration stimulating higher chlorophyll content and vice versa. Although these two cytokinins demonstrated similar descending pattern,
Figure 4. Mean shoot number produced by *H. littoralis* under different cytokinin treatments of various concentrations (2.25, 4.50, 9.00, 13.50, 18.00, and 22.50 µM) on semi-solid MS medium. (A) After 30 days of cultivation. (B) After 60 days of cultivation. Mean with the same alphabet on top of every bar is not significantly different [Dunnett's test (2-sided), α = 0.05].

Figure 5. Total chlorophyll content of *H. littoralis* under different concentrations (2.25, 4.50, 9.00, 13.50, 18.00, and 22.50 µM) for 2iP, TDZ and Zeatin on semi-solid MS medium after 60 days of cultivation.
Zeatin deposited higher total chlorophyll contents than TDZ in all of the six tested concentrations. This situation occurred because Zeatin was able to induce chloroplast differentiation but TDZ was known to inhibit the process (Oliveira et al., 2008). 2iP registered a fluctuating pattern between the total chlorophyll content of 2.00 and 5.00 across the six tested concentrations. The fluctuation can be explained by comparing the Figure 5. Before the first subculture, no plantlet in 2iP, TDZ or Zeatin demonstrated root growth. However, 2iP stimulated root growth especially at lower concentrations, prior to the second subculture. One of the characteristics of in vitro plantlets is the tendency to assume heterotrophic rather than autotrophic mode of nutrition as the nutrients are abundantly supplied in the medium. Hence, instead of depositing chlorophyll to synthesize food, the regenerated roots in 2iP plantlets directly absorbed nutrients from the medium resulting in unpredictable total chlorophyll contents across different concentrations (Hazarika, 2006). Nevertheless, explants treated with 13.50 µM of 2iP recorded the highest total chlorophyll content.

In conclusion, the effect of cytokinins 2iP, TDZ and Zeatin at various concentrations on the micropropagation of Hymenocallis littoralis using bulb scale discovered that 2iP and 13.50 µM respectively to be the best for shoot elongation. The types of cytokinins and their concentrations did not demonstrate any significant difference on the shoot multiplication of H. littoralis. The highest chlorophyll content was found to be in explants treated with 2.25 µM of Zeatin due to its ability to stimulate chloroplast differentiation. When the chlorophyll content analysis was considered specifically for 2iP, 13.50µM registered the highest total chlorophyll content out of the total six concentrations. This finding indeed strengthened the conclusion that 2iP and 13.50 µM were the best for shoot elongation as shoots are one of the main depositories of chlorophyll for photosynthesis.

REFERENCES


