To Study the Effect of Activated Charcoal, Ascorbic Acid and Light Duration on In Vitro Micropropagation of Aloe vera L

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ABSTRACT: The research was conducted to determine the effect of ascorbic acid (50 mg/l-1, 100 mg/l-1, 200 mg/l-1) and activated charcoal (0.5 gl-1, 1 gl-1, 2 gl-1) independently with different light duration (darkness for 4 weeks, and 16 hours light for 4 weeks followed by 2 weeks in 16 hours light) on shoot regeneration. Explants of Aloe vera were planted on MS basal media supplemented with 1.6 mg/l-1 IAA, 4.0 mg/l-1 BAP and cultured for 4 weeks. After 4 weeks, degree of explant browning was evaluated. Explants were then cut vertically into two pieces and planted on shoot regeneration media. After 4 weeks more number of shoot developed on shoot regeneration medium, i.e. MS medium supplemented with 1.6 mg/l IAA + 4.0 mg/l BAP + 100 mg/l ascorbic acid + 0.5 g/l activated charcoal. Browning of explants was minimized in this medium and elongation of micro-shoots and the growth of the plantlets were also better. Further elongation and rooting of micro-shoots were obtained when sub-cultured on to MS basal medium containing 0.5 g/l activated charcoal and 100% of the survival of rooted plantlets was observed after acclimatization. Therefore, the above protocol could be effectively used in rapid micropropagation of elite plants of Aloe vera.

KEYWORDS: Aloe vera, Explant browning, Micropropagation, Growth hormones, Light duration, Culture media, Phenolic compounds.

I. INTRODUCTION

Aloe vera (Liliaceae), is a succulent plant indigenous to Northern Africa and Mediterraneante countries and has become naturalized almost in all parts of India (Klein et al. 1988). The plant has stiff gray-green lance-shaped leaves containing clear gel in a central mucilaginous pulp. Aloe vera has been used for several thousands of years in folk medicine in many cultures from ancient Egypt, Greece, and Rome to China and India (Kemper and Chiou 1999). Some of the most important pharmacological activities of Aloe vera are antiseptic (Capasso et al. 1998), anti-tumor (Winter et al. 1981), anti-inflammatory (Yagi et al. 1998), wound and burn healing effect (Heggers et al. 1993), anti diabetic (Rajasekaran et al. 2006) and as an adjunct to current AIDS therapy (Mc Daniel et al. 1990).

Aloe vera propagates vegetatively in its natural state. However, propagation rate is very slow because a single plant can produce only three to four lateral shoots in a year. Moreover, the production of Aloe leaves is insufficient to meet the industry demand in India (Aggarwal and Barna 2004) and the production of cosmetics, foods and pharmaceuticals containing A. vera has experienced a slow increase due to limited availability of raw material with high quality (Campestrin et al. 2006). Therefore, there is a need to develop suitable, an alternative method for traditional propagation of A. vera.

In vitro techniques using micropropagation and tissue culture offer a great possibility to overcome this problem. Micropropagation using stem and lateral shoot pieces of Aloe vera has already been proved successful (Natali et al. 1990; Roy and Sarkar 1991; Mayer and Staden 1991; Aggarwal and Barna 2004). However, source of explants, their sterilization procedure, media composition, culture conditions, phenolic browning of explants and media discoloration greatly affect shoot regeneration from different genotypes of the same species. Aloe vera exudes lot of phenolic...
substances into the culture media which could decrease the survival of explants (Roy and Sarkar 1991). Concentration of phenolic compounds may vary in different genotypes of the same species (Glynn et al. 2004), and also those were grown under different climatic conditions (Kjaer et al. 2001).

Problem often encountered in mass micropropagation of Aloe vera is the explant browning. Explant browning caused by oxidation of phenolic compounds resulting from injuries during the isolation of explant[1]. It leads to the death of explant and failure of shoot regeneration[2]. This is due to quinones produced by oxidation of phenolic compounds[3] are toxic and it diffuses into culture media causing tissue necrosis and death of explant[4]. The concentration and combination of auxin and cytokinin in culture media is a key factor which determines successful shoot regeneration[5]. Besides growth hormone, other compounds such as ascorbic acid and activated carbon can be added to culture media. Both compounds are able to reduce the oxidation of phenolic compounds that can prevent death due to explant browning and increase shoot regeneration[6]. Ascorbic acid is an antioxidant that are able to prevent or inhibit oxidation process[7]. Besides its role as an antioxidant, ascorbic acid is involved in cell division and elongation[8]. Phenolic browning of explant, media composition and culture conditions greatly affect shoot regeneration[9]. Activated charcoal is an essential component of plant tissue culture media. It is a strong adsorbent that can absorb toxic substances[10].

Effect of light duration on regeneration shoots is also evaluated. According to[11] light plays an important role in the regeneration shoots. The main objective of this research was to investigate the effect of ascorbic acid and activated charcoal in culture media independently and light duration on shoot regeneration on Aloe vera in vitro culture. Thus, present study aimed to develop a rapid and high frequency shoot regeneration protocol for elite plants of Aloe vera suitable for mass propagation by improving culture media while controlling phenolic browning of explants.

II. MATERIALS AND METHOD

2.1 Source of Explant: Shoots with young leaves was collected from the elite plants. The extra leaves were removed and the explants were prepared by removing roots and brown colored tissues and extending leaf portions to give an average size of 3-4cm.

2.2 Sterilization Procedure: The explants were washed thoroughly with running tap water for about 10 minutes till all soil and other foreign materials washed off. Sets of twenty explants were then washed with tap water containing a few drops of Tween 20 and rinsed in 70% ethanol for 30 seconds followed by initial soaking in sodium hypochlorite approximately for 10 minutes and then in freshly prepared mercuric chloride solution (0.1 %) for 10 minutes. Finally they were washed 3-4 times with sterile distilled water before culturing.

2.3 Initiation Media: Surface sterilized explants were planted on Murashige and Skoog (1962) basal media supplemented with growth regulator hormone BAP 4 mg l-1 and IAA 1.6 mg l-1. Ascorbic acid and activated charcoal were added into media independently. The concentration of ascorbic acid was 50 mg l-1(A-1), 100 mg l-1 (A-2), 200 mg l-1 (A-3) and activated charcoal 0.5 g l-1 (K-1), 1 g l-1 (K-2), 2 g l-1 (K-3). pH media was 5.7-6.1. 50ml of nutrient medium was dispensed into 300ml glass jars that were capped with polycarbonate caps.

2.4 Culture Condition: Explants were incubated at –

1) Darkness for 4 weeks (D)
2) Light 16 hours for 4 weeks (L)
3) Darkness for 2 weeks followed by light 16 hours for 2 weeks (DL).

All culture conditions were at room temperature 27± 2°C under cool white fluorescent tubes (20-30 E.m-2.s-1), using Phillips lamps of 20 watt which were placed in 20 cm above bottles. Total sample of each treatment was twenty.

After 4 weeks on initiation media, explant browning was evaluated using degree of explants browning (Figure 1). Then explants were cut vertically into two pieces and planted on shoot regeneration media which its concentration is same as initiation media. Cutted explants were incubated at the same condition as mentioned (c).
III. RESULT AND DISCUSSION

3.1 Effect of Ascorbic Acid and Light Duration on Regeneration of Shoot:

Explants on ascorbic acid free media (A-0) produced a higher number of shoots than those on media with addition of ascorbic acid in concentration of 50 mg/l-1 (A-1), 100 mg/l-1 (A-2) and 200 mg/l-1 (A-3) (Figure 2). Ascorbic acid is known to decay rapidly in plant tissue culture media. Ascorbic acid is oxidized by reactions catalysed by Cu (II) and Fe (III), both of which are component of Murashige and Skoog media[12]. Light and pH accelerated the decay[12]. In darkness, ascorbic acid might be preserve. Ascorbic acid was most stable at pH 4.5[12]. In this research, media culture used at pH 5.9 - 6.1. Since ascorbic acid in an ephemeral component of culture media, it is quite possible that none exist when explant is planted.

The result showed explants of Aloe vera produced higher number of shoots at dark condition for 4 weeks (D) than at 16 hours light for 4 weeks (L). Even at dark for 2 weeks followed by light 16 hours for 2 weeks (DL) number of shoots produced higher than at 16 hours light for 4 weeks (L). Dark conditions enhanced higher number of shoots than light conditions suggesting that Aloe vera in vitro culture is a photomorphogenically process. Although light may be essential for plant development, darkness is also beneficial for plant morphogenesis.

According to[9], addition of ascorbic acid to the surface of culture media not only prevented the development of lethal browning but also greatly increased the number of shoot produced. The result showed increasing of ascorbic acid concentration in media did not increase the number of shoot produced (Figure 2). The effectiveness of ascorbic acid to induce regeneration shoots suggests that adding ascorbic acid onto the surface of the media should be tried.
3.2 Effect of Light Duration and Degree of Explant Browning on Number of Shoots Produced by Explant on Media with Addition of Ascorbic Acid:

Explants with degree of browning level 1, 2 and 3 on ascorbic acid free media and media with addition of ascorbic acid produced shoots. No shoots produced by explant with degree of explant browning level 4 (Figure 3). The explant became necrotic and die. Number of shoots produced depends on degree of explant browning. The higher degree of explant browning, the lesser number of shoots produced.

At all degree of explant browning, increasing of ascorbic acid concentration could not induced more shoots. In ascorbic acid free media, more shoots produced at all degree of explant browning, especially in darkness for 4 weeks (D). It showed that darkness condition is beneficial for phenolics contents of explant incorporated with the use of an unstable ascorbic acid.

**Figure 3.1:** Effect of Light Duration and Degree of Explant Browning on Number of Shoots Produced on Media without Ascorbic Acid

**Figure 3.2:** Effect of Light Duration and Degree of Explant Browning on Number of Shoots Produced on Media with 50mg/l Ascorbic Acid
3.3 Effect of Activated Charcoal and Light Duration on Regeneration of Shoot:

The result showed that explants on activated charcoal-free media (K-0) produced a higher number of shoots than those on media with addition of activated charcoal in concentration of 0.5 gl-1 (K-1), 1 gl-1 (K-2) and 2 gl-1 (K-3) (Figure 4). Activated charcoal is a strong adsorbent [10]. It adsorbs not only toxic substances, but also nutrients in media. Use of activated charcoal as an adsorbent was not an appropriate option. The development of a activated charcoal - free media is an alternative.

Figure 4 showed that explant on media with addition of activated charcoal in concentration of 2 gl-1 (K-3) produced more shoots than activated charcoal in concentration 0.5 gl-1 and 1 gl-1. Activated charcoal stimulate growth and differentiation during culture regeneration [9]. However its addition to regeneration media may have adverse effects on growth and development as activated charcoal is able to adsorb high concentration of growth regulators [13]. The non-selective adsorption of growth regulators may result in inhibitory effects of growth in vitro [14]. It will reduce their effectiveness in tissue culture.
3.4 Effect of Light Duration and Degree of Explant Browning on Number of Shoots Produced by Explant on Addition of Activated Charcoal Media:

Explants with degree of browning level 1, 2 and 3 on media with addition of activated charcoal produced shoots. Even on free activated charcoal media. No shoots produced by explant with degree of browning level 4 (Figure 5). The explant became necrotic and die. Number of shoots produced depends on degree of explant browning. The higher degree of explant browning, the lesser number of shoots produced.

In producing number of shoot, there is a big difference between explant on activated charcoal free media and media with addition of activated charcoal. In all degree of explant browning, increasing of activated charcoal concentration could not induced more shoots. Combination of BAP and IAA was crucial for shoot regeneration. It showed that degree of explant browning incorporated with adsorption properties of activated charcoal result in variations in plant growth regulator levels in media.
Figure 5.2: Effect of Light Duration and Degree of Explants Browning on Number of Shoots Produced by Explants on Addition of 0.5 g/L of Activated Charcoal

Figure 5.3: Effect of Light Duration and Degree of Explants Browning on Number of Shoots Produced by Explants on Addition of 1g/L of Activated Charcoal

Figure 5.4: Effect of Light Duration and Degree of Explant Browning on Number of Shoots Produced by Explant on Addition of 2g/l of Activated Charcoal
IV. CONCLUSION

The research described an efficient protocol for shoot regeneration of *Aloe vera* in vitro culture. Higher number of shoots produced on media without addition of ascorbic acid and activated charcoal. Dark conditions enhanced higher regeneration shoot than light condition. The optimal collaboration of a dark incubation period together with growth regulator hormone increased regeneration shoot. Although light may be essential for plant development, darkness is also beneficial for plant morphogenesis, mainly at its initial stage of development in vitro. Degree of explant browning greatly affects shoot regeneration. The higher degree of explant browning the lesser number of shoots produced.

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REFERENCES


