An easy and reliable method for establishment and maintenance of tissue cultures of *Nicotiana tabacum* cv. TAPM 26

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**Abstract**

*In vitro* regeneration of *Nicotiana tabacum* was efficiently achieved using growth regulators combination supplemented into semi-solid MS medium using leaves from 1-month-old plant as explants. The growth regulators used were NAA for auxin (0.0, 1.0, 2.0, and 3.0 mgL\(^{-1}\)) and BAP (0.0, 0.5, 1.0, 2.5, and 5.0 mgL\(^{-1}\)). The effects of growth regulator combination were assessed based on a number of callus formation, shoots formation and fresh callus weight. The maximum number of callus formation was 100% at five hormone combination, observed at the fourth week after culture. Maximum number of shoots produced per explant was 21.4 shoots at 1.0 mgL\(^{-1}\) BAP + 3.0 mgL\(^{-1}\) NAA, after 6 weeks of culture. The maximum callus fresh weights were obtained at 0.5 mgL\(^{-1}\) BAP + 1.0 mgL\(^{-1}\) NAA after 6 weeks (9.92 g). The best combination for shoots regeneration of *Nicotiana tabacum* was 1.0 mgL\(^{-1}\) BAP + 3.0 mgL\(^{-1}\) NAA. And the best combination for heaviest callus production was 0.5 mgL\(^{-1}\) BAP + 1.0 mgL\(^{-1}\) NAA.

**Abbreviations** BAP, 6-benzyl-aminopurine; IAA, indole-3-acetic acid; MS, Murashige and Skoog medium (1962); NAA, naphthaleneacetic acid

1. Introduction

Plant tissue culture covers the development of selected plant tissue and grown aseptically for indefinite duration on a nutrient medium under controlled conditions (Mohammed, 2020). Tissue culture usually applied as a medium for micropropagation, creation of virus-free plants (Arvas et al., 2018), genetics transformation (Kutty et al., 2011; Kaya et al., 2013) and it is also demonstrated more effective in the creation of secondary metabolites,
for examples, phenolics (Ozyigit, 2008), Juglone (Kocacaliskan et al., 2018) and Anthocyanins (Marchev et al., 2020) etc. The most regular technique in tissue culture is micropropagation which defined as a technique to procreate genetically clonal plantlets by using tissue culture methods and it supports in creating pathogen-free stock plants (Arvas et al., 2018) or genetically superior clones that cannot be propagated by seeds or plant that with low propagation efficiency in conventional vegetative propagation (Kutty et al., 2010). Moreover, plant cell exhibits its totipotency that every single cell can regenerate into a whole new plant.

Shoot regeneration for tobacco from an explant has achieved attention in the past and has promising application in the area of plant biotechnology (Deo et al., 2010). In addition, Kaya (2010) showed that adventitious regeneration of tobacco plant has a higher regeneration potential via embryogenesis when explants derived from cotyledons were used. Thus, the aim of the research is to establish an easy and reliable technique for maintenance of tissue cultures of Nicotiana tabacum cv. TAPM 26.

2. Materials and Methods

2.1. Culture Conditions

The seed of Nicotiana tabacum cv. TAPM 26 was provided by National Tobacco Board Kota Bharu Kelant. The study was achieved in the Department of Biosciences, Faculty of Science, Universiti Teknologi Malaysia. In vitro plant regeneration was pursued at 25 ± 2°C in either dark or light with a 16 h photoperiod and 6 h dark distributed by cool white fluorescent tubes. The tissue culture medium was heated to melting solutes and then distributed in tissue culture tubes for seed germination of tobacco plant and tubes for regeneration.

2.2. Basic Media Preparation

Basic media was prepared based on Murashige and Skoog (1962) containing mineral salts, micronutrients and vitamins, supplied with 20 gL⁻¹ sucrose, 0.1 gL⁻¹ myo-inositol and 3.2 gL⁻¹ phytigel. The pH of the plant culture media was adjusted to 5.7 with either 1N NaOH (sodium hydroxide) before sterilization. The tissue culture media were sterilized at 121°C.
for 21 min in an autoclave. After autoclaving, the medium was left to cool down to 40 to 45°C before combination of growth hormone (BAP and NAA) were added.

2.3. Preparation of Explants Material

Approximately 30 seeds of *Nicotiana tabacum* was placed in a test tube supplied with 2 mL of sterilizing solution (70 % ethyl alcohol). It was mixed gently for 20 min. Then the sterilising solution removed and washed all the seeds with double sterile water for 3 times. The seeds were put onto a sterile petri dish for drying and then the seeds were transferred to germination medium. Only 5 seeds were put onto each petri dishes. After a few days the germinated seeds will be ready to be used as a young plantlet that will be the source of explants for plant tissue culture work. The leaves of plantlet were sterile as they grow in the tissue culture systems under controlled parameters. In 4 weeks, the old plantlets were used as sources of explant. Callus induction and shoots regeneration from callus was obtained by using MS medium modified with auxin and cytokinin. The callus was induced from leaf explant on media contained MS (Murashige and Skoog, 1962) salts, vitamins, 30 gL⁻¹ sucrose, 8 gL⁻¹ agar powder, 6-benzylaminopurine (0.0, 0.5, 1.0, 2.5 or 5.0) and 1-naphthalene acetic acid (0.0, 1.0, 2.0 or 3.0 mgL⁻¹).

The pH was adjusted during each experiment to 5.7 ± 1 with 1 N NaOH or 1 N HCl using an electronic pH indicator. All the operations and inoculations were carried out under aseptic conditions in laminar airflow cabinet.

2.4. Data Analysis

All the assays were carried out in triplicates and the data were subjected to statistical analysis of one-way ANOVA using SPSS 15.0 for Windows. A value of P<0.05 was considered to be significant.

3. Results and Discussion

The explants produced callus on the one month after cultivation on all combination of NAA and BAP at room temperature and 16 h light under plant growth chamber. The results achieved in the aspects of callus weight and percentage of callus from hypocotyl explants.
The first step in the start of plant tissue culture represents the disinfection of the explant or seed. The efficiency of the surficial sterilization technique has an important effect on the development and growth of the plant for tissue culture. Standard surficial sterilization technique for starting a tissue culture of members of the tobacco plants comprise the application of alcohol, sodium hypochlorite (Steyn et al., 1996; Ganapathi et al., 2004).

3.1. Effect of BAP/NAA on Callus Fresh Weight

In Figure 1, the maximum average weights of callus form were detected at 0.5 mgL\(^{-1}\) BAP + 1.0 mgL\(^{-1}\) NAA. The maximum mean weight produces was 9.92 g while the minimum mean weight was 0.127 g observed at the presence of zero growth regulator. The product of experiments was analyzed using one-way ANOVA and the results were significant for a value \(P<0.05\). From the previous researches, the maximum mean weight produced was 1.81 g at 0.2 mgL\(^{-1}\) BAP and 3.0 mgL\(^{-1}\) NAA (Ali et al., 2007). This showed that lower concentration of BAP and NAA created heavier and bulkier callus but too low or too high concentration may retard the growth of the callus. Figure 2 showed the best callus formed in terms of fresh callus weight as described in Figure 1.

![Figure 1. The average weight of callus on different combination of growth regulator](image)
3.2. Effect of BAP/NAA on Callus Formation

At 3.0 mgL⁻¹ NAA concentration, the callus formation was lower than others (Table 1). The same result was observed for the maximum concentration BAP which only formed 20% of callus compared to other BAP and NAA combinations. This showed that explants need both growth regulators combination to form callus.

Table 1. Percentage of callus formed for each growth regulator combinations

| Hormone combination mgL⁻¹ | BAP  | 0.0% | 0.5 | 1.0 | 2.5 | 5.0 |
|---------------------------|------|------|-----|-----|-----|-----|
| NAA                       |      |      |     |     |     |     |
| 0.0                       |      | 33.3%| 86.7| 80.0| 60.0| 20.0|
|                           |      | callus formation | callus formation | callus formation | callus formation | callus formation |
| 1.0                       |      | 53.3%| 86.7| 86.7| 93.3| 33.3|
|                           |      | callus formation | callus formation | callus formation | callus formation | callus formation |
| 2.0                       |      | 53.3%| 100 | 100 | 100 | 73.3|
|                           |      | callus formation | callus formation | callus formation | callus formation | callus formation |
| 3.0                       |      | 13.3%| 100 | 100 | 73.3| 66.7|
|                           |      | callus formation | callus formation | callus formation | callus formation | callus formation |

In Figure 3, the maximum number of callus induction percentage was 100% at MS
media containing 0.5 mgL\(^{-1}\) BAP + 2.0 mgL\(^{-1}\) NAA, 1.0 mgL\(^{-1}\) BAP + 2.0 mgL\(^{-1}\), 2.5 mgL\(^{-1}\) BAP + 2.0 mgL\(^{-1}\) NAA, 0.5 mgL\(^{-1}\) BAP + 3.0 mgL\(^{-1}\) NAA and 1.0 mgL\(^{-1}\) BAP + 3.0 mgL\(^{-1}\) mgL\(^{-1}\) NAA. Maximum concentration of hormone combination gives a medium percentage of calli formation, about 66.7%. This indicated that too low or too high concentration of growth regulator may inhibit the formation of calluses.

**Figure 3.** The percentage of callus formed in different growth regulators combination

**3.3. Effect of BAP/NAA on Shoots Induction**

In Figure 4, it demonstrationed that shoots need 2.0 mgL\(^{-1}\) and 3.0 mgL\(^{-1}\) NAA concentrations. The highest average number of shoots was 21.4 ± 2.13 per explant. At 0.0 mgL\(^{-1}\) and 1.0 mgL\(^{-1}\) of NAA concentration, there was no shoot formed. At all BAP concentrations, there were shoots formed and it varied upon the combination. The highest average number of shoots per explant was obtained from the combination of 1.0 mgL\(^{-1}\) BAP + 3.0 mgL\(^{-1}\) NAA. The results were analysed using one-way ANOVA and the results were significant for a value P<0.05. The lowest number of shoots for per explants was observed on MS medium supplemented with 2.0 mgL\(^{-1}\) NA (1.067 shoots per explants). Roots initiation was not detected on all explants after 6 weeks culture. This showed that the NAA concentration should be higher to achieve rooting or it takes longer time to regenerate. Figure
5 shows the shoot callus for the best combination growth regulator used as described in Figure 4.

![Figure 4](image)

**Figure 4.** The average number of shoots formed

![Figure 5](image)

**Figure 5.** Shoots regenerated at 1.0mgL⁻¹ BAP+ 3.0mgL⁻¹ NAA. (Scale bar = 0.5cm)

From Table 2, it was indicated that different sizes of callus formed at different combination of growth regulator. From the observation, the combination of 0.5 mgL⁻¹ BAP and 2.0 mgL⁻¹ NAA showed the best biggest callus size formed.
Table 2. Different sizes of callus formed regenerates based on varies growth regulator concentration after treatment for 6 weeks

| Hormone combination | BAP concentration in mgL⁻¹ |
|---------------------|---------------------------|
|                     | 0.0 | 0.5 | 1.0 | 2.5 | 5.0 |
| 0.0                 | ![Image](image1.jpg)       |
| 1.0                 | ![Image](image2.jpg)       |
| 2.0                 | ![Image](image3.jpg)       |
| 3.0                 | ![Image](image4.jpg)       |

The tobacco plant is created of several repetitive modules in its vegetative part, successively generated by meristems. Tissue culture methods via calli have become an essential tool for plant biotechnology such as *Gossypium hirsutum* L. (Ozyigit et al., 2007), *Impatiens balsamina* (Taha et al., 2009), *Lycopersicon esculentum* Mill. (Jawad et al., 2020), *Oryza sativa* (Kaya and Karakutuk, 2018) *Tagetes minuta* (Latifian et al., 2018), *Citrullus lanatus* cv. Round Dragon (Ganasan and Huyop, 2010) and many others. The explant type and hormones are also significant influences for *in vitro* plant regeneration (Kumar and Reddy, 2010). Explant was a critical parameter when optimizing tissue culture methods (Kumar et al., 2011). Therefore, choice of appropriate explants is an important factor of tissue culture of *Nicotiana tabacum* (Kaya, 2010). Growing the explant of *Nicotiana tabacum* under different concentrations of plant growth hormones such as BAP, NAA as very essential (Table 1). The combination of plant growth hormone was crucial in determining the tissue culture performance resulting in plant regeneration seen in shoot cultures. The present study (Figures 3 and 4) established the good combination of hormone in the media preparation where the shoot formation from regeneration of callus were established.

In conclusion, the results obtained showing the success of tissue culture of *Nicotiana tabacum* with the best hormone combination would be MS + 1.0 mgL⁻¹ BAP + 3.0 mgL⁻¹
NAA for production of maximum number of callus and also the highest shoots formation. The callus started to form at fourth weeks after cultivation and the shoots started to initiate two weeks later. Suitable growth regulator combination promote the optimum growth explants and also induced the callus and shoots formation. It also helps to shorten the time taken to initiate callus and shoots. The present method demonstrated that *Nicotiana tabacum* TAPM 26 was successfully propagated by organogenesis and can be applied for further research such as transgenic *Nicotiana tabacum* TAPM 26.

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