Enhanced in vitro Shoot Regeneration and Biochemical Properties of Stevia rebaudiana using Chitosan

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ABSTRACT

Stevia rebaudiana is a herbaceous perennial plant with great global demand due to its beneficial steviol glycosides (SGs) content. Current conventional breeding technique is unable to cater the need for more S. rebaudiana planting materials. Therefore, an improved in vitro shoot regeneration protocol was developed for S. rebaudiana by using chitosan. The highest fresh weight of plant (0.586 ± 0.176 g/explant), dry weight of plant (0.056 ± 0.02 g/explant) and plant height (4.94 ± 1.17 cm/explant) with maximum number of leaves (25.33 ± 6.95/ explant) were observed on explants grown in optimun treatment of MS basal medium supplemented with 1.0 mg/L of 6-benzyaminopurine (BAP) and 60 mg/L of low molecular weight (MW) chitosan after 4 weeks of culture. Scanning electron microscopy (SEM) analysis showed that new shoot primordia and shoot bud formation can be seen as early as day 3 and 5 of cultures on optimun treatment. Further biochemical assays showed that total phenolic acid content, total protein content and total hydrolyzed sugar content were recorded higher in explants cultured in optimun treatment as compared to control media. In contrast, total chlorophyll content and total flavonoids content were reduced in optimum treatment. Meanwhile, no significant difference in antioxidant activity was observed. All cultures from the optimal treatment were successfully regenerated, acclimatized and grew well with 100% survival rate. Taken together, an enhanced and efficient shoot regeneration protocol of S. rebaudiana was successfully developed which will be useful for rapid and large-scale micropropagation in future.

Keywords: Chitosan; micropropagation; scanning electron microscope (SEM); stevia; tissue culture

ABSTRAK

Stevia rebaudiana ialah sejenis tumbuhan herba saka yang mempunyai permintaan antarabangsa yang tinggi dengan kandungan steviol glikosid (SGs) yang berfaedah. Kaedah biakbaka konvensional tidak dapat memenuhi permintaan anak pokok S. rebaudiana yang semakin meningkat. Oleh itu, satu protokol penambahbaikan penjanaan semula pucuk S. rebaudiana secara in vitro dengan menggunakan kitosan telah dibangunkan. Tumbuhan dengan berat basah maksimum (0.586 ± 0.176 g/eksplan), berat kering maksimum (0.056 ± 0.02 g/eksplan), ketinggian maksimum (4.94 ± 1.17 cm/ eksplan) dan bilangan daun tertinggi (25.33 ± 6.95/ eksplan) telah diperhatikan dalam media bes MS yang ditambah dengan 1.0 mg/L 6-benzilaminopurina (BAP) yang ditambah dengan kitosan berat molekul (MW) rendah, 60 mg/L, selepas 4 minggu dari tempoh kultur. Analisis mikroskop elektron pengimbas (SEM) telah menunjukkan pertumbuhan pucuk primordium baru serta tunas pucuk telah wujud seawal 3 hari dan 5 hari selepas rawatan optimum kitosan. Selanjutnya, asai biokimia telah menunjukkan bahawa kandungan jumlah asid fenolik, kandungan protein dan kandungan gula terhidrolisis adalah lebih tinggi di dalam eksplan yang tumbuh di dalam media optimum berbanding dengan media kawalan. Namun begitu, kandungan klorofil dan flavonoid di dalam eksplan adalah berkurangan dalam media optimum. Tiada perbezaan ketara dapat dilihat di dalam kandungan antioksidan. Semua kultur daripada rawatan optimum telah berjaya ditumbuh semula, diaklimatiti dan tumbuh dengan sihat pada kadar kelangsungan hidup 100%. Kesimpulannya, satu protokol penjanaan semula S. rebaudiana yang ditambahbaik serta berkesan telah berjaya dibangunkan dan berguna untuk pembiakan dengan cepat dan besar-besaran pada masa hadapan.

Kata kunci: Kitosan; kultur tisu; mikroperambatan; mikroskop elektron pengimbas (SEM); stevia
INTRODUCTION

**Stevia rebaudiana** is a perennial herb that belongs to the Asteraceae family and originated from northern regions of South America (Chiew et al. 2019). The leaves are rich with sweetening agents known as SGs, which are estimated to be 100 - 300 times sweeter than sucrose. SGs are mainly composed of stevioloside and rebaudioside A (Reb-A) (Chiew et al. 2016; Gauchan et al. 2014). Over the years, the demand for *S. rebaudiana* has been increasing rapidly in the global market (Future Market Insights 2014). This is because SGs are zero in calories and suitable to be consumed by diabetes patient (Ummi et al. 2014). Hence it can be widely used to replace artificial sweetening agent in food and beverage industry.

In general, *S. rebaudiana* plantlet is propagated through conventional plant breeding including stem cutting, seedling and cross-pollination methods. However, these conventional methods are low in germination rate (below 10 %), slow in regeneration efficiency, and higher variation in offspring (Tadhani et al. 2006; Ummi et al. 2014). Often, seed germination does not produce a homogenous population of offsprings, resulting in variations in SGs level and composition. In addition, the qualities of the plantlets produced are unstable and variable. Most importantly, conventional methods rely heavily on human capitals for stem cutting and maintenance. Thus, an alternative route for propagation of *S. rebaudiana* is through in vitro micropropagation techniques. Several studies had reported the use of vegetative tissues such as nodal segment and axillary bud (Das et al. 2011), shoot (Ummi et al. 2014), and hypocotyl (Ramirez-Mosqueda & Iglesias-Andreu 2016) for in vitro micropropagation of *S. rebaudiana*. Despite these reported studies, there is a need to improve the existing formulation for more efficient micropropagation of *S. rebaudiana*. This is because the efficiency of in vitro micropropagation is greatly influenced by factors such as plant genotypes, type of explant, medium composition, and growth additives.

Chitosan is a growth enhancer, widely used in agricultural industry. It is a biopolymer deacetylated form of chitin (poly [β-(1->4)-2-amino-2-deoxy-D-glucopyranose]) that present in outer shell of organisms, such as shrimp, crab, and crustaceans (Chawla et al. 2015). Chitosan is bio-degradable, safe, and environmental friendly. Generally, chitosan can be divided into low (50 - 190 kDa), medium (190 - 310 kDa) and high molecular weight (MW) (310 - 375 kDa) with a deacetylation degree of 75 - 85% (Grobler & Perchyonok 2018). Low MW chitosan shows higher solubility and low viscosity in water at physiologically acceptable pH values as compared to medium and high MW chitosans. Previously, chitosan had been reported to use as growth enhancer in plant species such as *Oryza sativa* (Channammanoontham et al. 2015) and orchid (Pornpienpakdee et al. 2010; Sopalan et al. 2010). Besides, chitosan also used to enhance secondary metabolite production in ginseng (Jeong & Park 2005) and *Psoralea corylifolia* (Ahmed & Baig 2014). In addition, it also enhances plant disease resistance towards pathogens attack (Uthairatanakij et al. 2007) and various environmental stresses (El Hadrami et al. 2010). However, the efficiency of chitosan is greatly dependent on the type and concentration used (Sivanandhan et al. 2013). Therefore, a proper optimization and manipulation of chitosan are required in order to achieve the desire outcomes.

To date, the effects of chitosan on *S. rebaudiana* regeneration remain largely unknown despite it has been proven to be effective in other plant species. Hence, the aim of this study was to enhance the shoot regeneration protocol of *S. rebaudiana* through optimization of suitable types and concentration of chitosan used. Furthermore, biochemical properties of *S. rebaudiana* cultured on control and optimum treatment media were also analyzed. Ultimately, the improved shoot regeneration protocol with growth enhancer can be used to speed up the mass micropropagation of *S. rebaudiana*.

MATERIALS AND METHODS

PLANT MATERIALS PREPARATION

One month old potted *S. rebaudiana* plant was bought from Jacky Lim Nursery, Sungai Buloh, Malaysia (3.223257, 101.589001). Young shoot tips were collected and nodes measuring 2 cm were chosen as explants from potted *S. rebaudiana* plant. Explants were cut and washed thoroughly under running tap water containing two drops of surfactant Tween-20 solution for 15 min. In aseptic condition, the explants were surfaced sterilized with 70% (v/v) ethanol for 3 min and rinsed with sterilized distilled water for 3-5 min, followed by 10% (v/v) chlorox solution for 15 min and rinsed with distilled water again. Explants were dried with sterile blotting paper and cut into 2 cm in sterilized petri dish.

MEDIA PREPARATION ON SHOOT INDUCTION

Excised explants of *S. rebaudiana* were transferred and cultured on Murashige and Skoog (MS) basal medium (Murashige & Skoog 1962) supplemented with 30 g/L sucrose and solidified with 7.5 g/L of agar (Gelrite) and incorporated with 1 mg/L 6-benzylaminopurine (BAP). The pH of the medium was adjusted to 5.70 with 0.1 M NaOH or HCl before autoclaving at 121 °C for 15 min. The cultures were incubated at 24 ± 2 °C with a
Sixteen hours photoperiod. The regenerated shoots were cut and individual shoots were placed on MS medium supplemented with 1 mg/L BAP for further multiple shoots induction.

CHITOSAN TREATMENT
Nodal explants from four weeks old *S. rebaudiana* shoot tips (~1 cm) cultured from MS medium supplemented with 1.0 mg/L BAP were cut and individual explant was cultured on MS medium containing different MW (low, medium and high) and concentration (0, 20, 40, 60, 80, and 100 mg/L) of chitosan supplemented with 1.0 mg/L of BAP. Excised explants cultured on MS medium supplemented with 1.0 mg/L of BAP only (without chitosan) was used as control. Data for plant height, number of leaves, fresh weight (FW) and dry weight (DW) for all treatments were collected at four week-old culture.

SEM
For scanning electron microscope examination, day 3 and day 5 *S. rebaudiana* samples cultured in optimum concentration of chitosan medium were used. Scanning electron microscopy was carried out according to protocol by Lai et al. (2011) with slight modifications. Firstly, samples were cut into 1 cm² slices and put into separate vials according to different day of culture. Then, samples were fixed in fixative which contained 4% glutaraldehyde for 2 days. After 2 days of primary fixation, samples were washed with 0.1 M sodium cacodylate buffer for 3 times of 30 min each and post-fixed in 1% osmium tetroxides for 2 h at 4 °C. Next, samples were washed again with 0.1 M sodium cacodylate buffer for 3 times (30 min each) before dehydration process was done. After fixation and washing process was done, samples were subsequently dehydrated in a series of acetone (35, 50, 75, and 95% once for 30 to 45 min at each step, and then immersed in 100% of acetone thrice for 1 h each). Then, the samples were transferred from absolute acetone to critical point drying apparatus. By using critical point drying (CPD) method, specimens were transferred into specimen basket and put into critical dryer for half an hour. Specimens dried by CPD method were transferred and mounted to specimen stub using colloidal silver. A number of gold layers were coated over the specimen to increase the conductivity of the secondary electrons. After the coating process, specimens were observed under scanning electron microscope (JEOL JSM-600, Japan) in Institute of Bioscience (IBS), Universiti Putra Malaysia (UPM).

REGENERATION OF S. rebaudiana
Four weeks old of well-grown *S. rebaudiana* explant cultured at optimum media was transferred to rooting media with MS basal medium supplemented with 0.4 mg/L of NAA growth regulator (Thiyagarajan & Venkatachalam 2012). After 3 weeks of rooting treatment, well-rooted plantlet with multiple shoot was acclimatized and transferred to pot with sand and soil (1:1) for adaptation.

BIOCHEMICAL ASSAY ANALYSIS
Biochemical assays were carried out using leaves of 4 weeks old *S. rebaudiana* cultured on MS medium supplemented with 1.0 mg/L of BAP (Control) and optimum treatment media (MS medium supplemented with 1.0 mg/L BAP and 60 mg/L of low MW chitosan). Biochemical assay performed included total chlorophyll assay (Talreja 2011); total phenolic acid assay (John et al. 2014); total flavonoid assay (John et al. 2014); total hydrolysed sugar assay (Van Handel 1985); total protein assay (Bradford 1976) and DPPH radical scavenging assay (Büyükün cel et al. 2014).

STATISTICAL ANALYSIS
Each of the growth parameters were carried out in triplicates with 3 samples per replicate. Data were analyzed using one-way ANOVA (analysis of variance) in a completely randomized design. The differences among means were determined by S-N-K (Student-Newman-Keuls) Test at 95% significance level. Similarly, each of the biochemical assay experiment was carried out in triplicates. Data were analyzed using independent sample of T-test in a completely randomized design. All statistical analyses were performed at 95% (p ≤ 0.05) significant level using SPSS (SPSS Inc. USA version 24).

RESULTS AND DISCUSSION
EFFECT OF CHITOSAN ON SHOOT REGENERATION
Previous studies had shown that chitosan can act as growth enhancer in plant due to its ability as an elicitor which could stimulate the production of endogenous phytohormone (Chamnanmanoontham et al. 2015). However, usage of chitosan as a growth enhancer should be studied properly because its efficiency is known to be highly dependent towards plant species, concentration and molecular weight (Dzung et al. 2017; Singh 2016). In our study, 60 mg/L low MW chitosan was found to be the optimum treatment in promoting the growth of *S. rebaudiana* (Figure 1). In MS media supplemented with 60 mg/L of low MW of chitosan, the highest plant height (4.94 cm), number of leaves (25.33), fresh weight (0.586 g) and dry weight (0.056 g) were recorded (Figure 1). Optimum concentration (60 mg/L of low MW of chitosan)
also induced early leaf formation which could be seen on the week 1 (Figure 3(c)). According to Rahman et al. (2018), chitosan could speed-up the formation of leaf due to changes in programming of protein metabolism in plant growth. Application of chitosan will promote the number of leaf formation had also been reported in the works done by Agbodjato et al. (2016) and Mondal et al. (2013) on maize plants. Their results showed that increasing concentration of chitosan increased the number of leaves until optimum concentration is achieved. However, other concentrations (20, 40, 80, and 100 mg/L) were not significantly enhanced the growth parameters of *S. rebaudiana* (Figure 1). High concentration of chitosan (80 and 100 mg/L) showed significant inhibition of the plant growth. This could be due to high concentration of chitosan, which is harmful to the plant (Singh 2016). Similarly, Asghari-Zakaria et al. (2009) also reported that high concentration of chitosan used in potato would eventually inhibit the plant growth. In addition, our study demonstrated that high MW of the chitosan resulted in distortion of leaf development (curly leaves) as shown in Figure S1. On the other hand, low concentrations of chitosan (20 and 40 mg/L) were found to be insufficient to induce the plant growth signalling pathway which will not enhance plant growth (Singh 2016).

**SEM ON SHOOT BUD FORMATION**

SEM was used to further observe the morphological changes and formation of shoot bud at the stem nodal culture on optimum treatment media (60 mg/L of low MW chitosan). SEM image showed that there was a formation of new shoot primordia in the culture sample which gave rise to shoot formation on day 3 of culture (Figure 2). Formation of new shoot primordia in a short period of time indicates a good regeneration efficiency of shoot bud in stevia plants. This result was positively correlated with the work done by San José et al. (2014) which showed that formation of shoot primordia was due to cell division and led to cell differentiation. Meanwhile, SEM image of day 5 culture sample with optimum treatment (Figure 2(b) - 2(d)) showed formation of shoot bud. The shoot primordia form shoot bud which will then give rise to shoot and leaf formation in the explant.

**REGENERATION OF *S. rebaudiana***

A well regenerated potted *S. rebaudiana* was observed after 4 weeks of acclimatization in pot (Figure 3). In this study, we successfully regenerated *S. rebaudiana* in 3 months by using chitosan as an enhancer. Similar works done by Thiyagarajan and Venkatachalam (2012) on regeneration on *S. rebaudiana* without using growth enhancer took approximately 4 months. Thus, mass propagation of *S. rebaudiana* with addition of chitosan as growth enhancer is recommended in order to shorten its regeneration time.

**BIOCHEMICAL ASSAYS**

The plant growth can be determined by total chlorophyll content as it is linked with the production of food sources in plant *via* photosynthesis (Muley et al. 2019). However, our study showed that total chlorophyll content in *S. rebaudiana* explant using control media was 55% higher than in optimum treatment media (Figure 4(a)). As reported previously, no significant difference in chlorophyll content was observed in okra plants treated with chitosan (Bistgani et al. 2017a; Mondal et al. 2012). It was suggested that the chitosan may not have direct effects on the total chlorophyll content, but it enhanced the photosynthetic activity which was driven by upregulated expression level of rubisco in carbon fixation (Zhang et al. 2017). In addition, SGs are synthesized via methylerthritol phosphate (MEP) pathway which also involved synthesis of chlorophyll pigments (Kumar et al. 2012; Totté et al. 2000). This suggests that there will be competition in energy consumptions between SGs synthesis and chlorophyll synthesis. Another study done by Zheng et al. (2019) on overexpression of MEP pathway responsive gene, *SrDXS* in *S. rebaudiana* plant also resulted in no difference in chlorophylls and carotenoids levels, suggesting that excess precursors from MEP pathway may divert biosynthesis on other secondary metabolites instead of chlorophylls. Therefore, similar activity may be applied in this study.

The total hydrolyzed sugar contents recorded in optimum treatment (0.102 mg/g) was significantly higher compared to control (0.077 mg/g) (Figure 4(b)). This result was in-line with the works done by He et al. (2018) and Zou et al. (2015) on strawberry and maize plants. In addition, study done by Chamnanmanootham et al. (2015) also showed altered expression of carbohydrate metabolism in rice due to chitosan treatment *via* increased expression level of glyceraldehyde-3-phosphate dehydrogenase, enolase, and phosphoglucomutase which indicate higher energy production level in rice. Thus, plant growth enhancer by chitosan might involve in regulation of carbon metabolism which lead to sugar accumulation (Chamnanmanootham et al. 2015; Zhang et al. 2017). As reported previously, derivatives of sugar is important as a precursor in SGs biosynthesis pathways (Ceunen & Geuns 2013; Yoneda et al. 2017). In addition, Sun et al. (2019) reported that increased of sugar accumulation lead to enhance of SGs production. Hence, 60 mg/L of low MW chitosan treatment could stimulate SGs production in *S. rebaudiana*. 
On the other hand, the protein concentration in *S. rebaudiana* plantlets grown in optimum treatment was higher compared to control treatment (Figure 4(c)). The result was similar with the study recorded by Anusuya and Sathiyabama (2016) on effect of chitosan on turmeric plants. As reported previously, chitosan could increase the protein content in the plant via enhancement of nitrogen and carbon assimilation (Zhang et al. 2017). The protein could be used to enhance plant metabolism for growth improvement (Suriyaprabha et al. 2017).

Phenolics are recognised as the largest phytochemicals that play an important role in plant defense mechanism against biotic and abiotic stresses (Samanta et al. 2011). Most of the phenolic compounds found in plants can be classified into phenolic acids while flavonoids (Hannan et al. 2016). Phenolic compounds also show antioxidant activity in plant (Rahman et al. 2018; Shukla et al. 2009). Phenolic compounds also show antioxidant activity in plant (Rahman et al. 2018; Shukla et al. 2009). As reported previously, chitosan could increase the protein content in the plant via enhancement of nitrogen and carbon assimilation (Zhang et al. 2017). The protein could be used to enhance plant metabolism for growth improvement (Suriyaprabha et al. 2017).

In this study, 60 mg/L of low MW chitosan had caused reduction in flavonoids while antioxidants content remained unchanged in *S. rebaudiana* (Figure 4(e) - 4(f)). However, 13% increment of phenolic acid was observed in chitosan treated plant (Figure 4(d)). As reported previously, chitosan treatment could impose additional stress to the plant (Lopez-Moya et al. 2017). Upon stress induction, plants are required to produce more phenolic compounds to maintain the cellular homeostasis (Jamalian et al. 2013).

In this study, it was found that the signalling pathway of phenolic acid was activated to counteract with the stress imposed by chitosan in *S. rebaudiana*. With the presence of phenolic acids to counteract with the stress, excessive free radicals will no longer be produced. Thus, the antioxidant capacity remained unchanged between control and chitosan treated plants. Other studies done by Hajihashemi and Geuns (2013) and Shukla et al. (2009) also suggested that phenolics compound may be the major contributors to antioxidants activity in *S. rebaudiana*. In addition, works done by Álvarez-Robles (2016) also suggesting that phenols biosynthesis pathway of *S. rebaudiana* will lead more on production of hydroxycinnamic acid derivatives than flavonoids production. Previous study done on thyme plants with chitosan treatment also showed similar result in which phenolic content was increased while flavonoids content remains almost unchanged (Bistgani et al. 2017b). Rahman et al. (2018) also reported that chitosan treatment successfully induced the production of phenolic acid in strawberry plants. Increased of phenolic acid together with enhanced plant growth supported the alternative concept on interlinked processes between growth and immunity instead of trade off relationship between plant growth and immunity (Kliebenstein 2016; Rahman et al. 2018).

**FIGURE 1.** Effect of different types and concentrations of chitosan on (a) average plant height, (b) average number of leaves produced, (c) average fresh weight, and (d) average dry weight of *S. rebaudiana* explant. All results were obtained after 4 weeks of culture. Result indicates the mean (± SE) of 3 independent experiments with 3 replicates for each treatment. Bars with different letters represent significant difference at $P \leq 0.05$. 
FIGURE 2. SEM observation on formation of shoot bud using optimum concentration 60 mg/L of low MW chitosan on the *in vitro* growth of *S. rebaudiana* using shoot explants after 3 days and 5 days culture sample: (a) SEM image of day 3 culture sample, and (b - d) SEM image of day 5 culture sample. White asterisk represents formation of new shoot primordia while white arrow represents formation of shoot bud.

FIGURE 3. Stages of *in vitro* plant regeneration of *S. rebaudiana*: (a) initiation (day 0) of *S. rebaudiana* stem nodal culture, (b) shoot bud formation after 5 days of culture in MS basal medium with 1.0 mg/L of BAP hormone and supplemented with 60 mg/L of low MW chitosan, (c) leaves formation after 1 week of culture in MS basal medium with 1.0 mg/L of BAP hormone and supplemented with 60 mg/L of low MW of chitosan, (d) development of shoots after 4 weeks of culture, (e) development of root of *in vitro* *S. rebaudiana* explant using root medium after 3 weeks of culture, and (f) well-grown regenerated *S. rebaudiana* plant after 4 weeks of planting. Bar a-e = 1 cm each, f = 5 cm.
An improved in vitro *S. rebaudiana* shoot regeneration protocol has been successfully established by using 60 mg/L of low MW of chitosan. Chitosan also able to enhance the production of proteins and hydrolysed sugars which will lead to better plant growth.

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