Plant Growth Regulator Effects on In Vitro Propagation and Stevioside Production in Stevia rebaudiana Bertoni

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Abstract. Stevia rebaudiana is of great importance due to its steviol glycosides (SGs) which are natural sweeteners used by the food industry as well as having medicinal purposes. In the present study, the effect of plant growth regulators (PGRs) and explant types on in vitro propagation and shoot growth of S. rebaudiana were studied, the effect of PGRs on SGs production was determined. For this purpose, nodal explants and shoot tip explants were cultured on woody plant medium (WPM) supplemented with cytokinins [6-benzyladenine (BA), kinetin (Kn), or thidiazuron (TDZ)] or cytokinins + auxins combinations [BA + indoleacetic acid (IAA); BA + naphthaleneacetic acid (NAA); Kn + IAA; Kn + NAA]. Although, the best shoot proliferation was obtained on WPM supplemented with BA + NAA combinations, shoots grown on PGR-containing media produced callus at the base of the shoots and showed chlorosis and necrosis. Additionally, shoots showed at all concentrations of TDZ, and at higher concentrations of BA, morphological changes such as malformed leaves and poor shoot growth. In contrast to PGR-containing media, on the PGR-free control medium, the development of shoots and roots occurred simultaneously and healthy and well-developed plantlets were obtained. Thus, we developed an economical viable means of in vitro propagation by minimizing the micropropagation steps and removing the requirement of PGRs. According to the high-performance liquid chromatography (HPLC) results, PGR-free control medium (WPM) led to considerably higher stevioside content in the leaves compared with the PGR(s)-containing media and the highest stevioside [34 mg g⁻¹ dry weight (DW)] and rebaudioside A content was only detected on the control medium without PGRs. Steviolbioside, rubusoside, and dulcoside A were detected qualitatively in the leaves of shoots grown on WPM supplemented with 2.27 μM TDZ, 4.54 μM TDZ, 2.22 μM BA + 2.69 μM NAA, 2.22 μM BA + 5.37 μM NAA, 2.32 μM Kn + 5.71 μM IAA, or 2.32 μM Kn + 2.69 μM NAA.

Stevia rebaudiana Bertoni belongs to the family Asteraceae and is a perennial that is primarily native to eastern Paraguay (Brandle et al., 1998). S. rebaudiana accumulates mainly in its leaves several noncaloric sweet-tasting SGs (Ceunen and Geuns, 2013). The two main SGs produced in the leaves are stevioside and rebaudioside A (Kinghorn, 2002). The sweetening properties of Stevia leaves have already been used since the sixteenth century by the native Paraguayan populations in the Amambay region (Soejarto, 2002). In the early 1970s, Japan started the commercial cultivation of S. rebaudiana, and sometime later the first Stevia products have been offered on the Japanese market. Since then, Stevia has been introduced in several countries and can be used as food additive and for therapeutic purposes (Brandle et al., 1998; Lemus-Mondaca et al., 2012).

In general, Stevia plants are propagated by seeds or vegetative cuttings. Since the Stevia seeds are very small and infertile, they have poor germination rate. Therefore, large-scale production of Stevia via seeds is not efficient (Brandle et al., 1998; Gantait et al., 2015). Besides this, the seeds show a wide variation with regard to the SGs content as well as the morphological features (the shape and color of leaves) (Tamura et al., 1984). Clonal propagation via vegetative cuttings is practical for small-scale production, but it is not suitable due to the small number of individuals and economically viable because of high labor costs for large-scale production (Brandle et al., 1998; Gantait et al., 2015). A genetically homogeneous population, which produces high yields of the desired SGs, can be achieved by in vitro propagation of a selected plant (Gantait et al., 2015; Lemus-Mondaca et al., 2012; Tamura et al., 1984). The use of apical and axillary meristems, shoot tips, or nodal explants for in vitro propagation, enables the conservation of the genotype during clonal propagation. In addition, in vitro plant propagation using axillary buds was reported to be a simple and economic method to obtain many genetically uniform true-to-type plants within a short period (Faisal et al., 2007).

The presence of diterpene glycosides was investigated in callus culture, suspension culture (Bondurev et al., 2001; Mathur and Shekhawat, 2013), etiolated in vitro regenerants, callus culture grown in the light, etiolated heterotrophic callus (Ladygin et al., 2008), hairy root culture (Pandey et al., 2016; Yamazaki and Flores, 1991), in vitro root culture (Reis et al., 2011), in leaf-derived callus cultures (Sivaram and Mukundan, 2003), salts (NaCl and Na₂CO₃), proline or polyethylene glycol–treated callus, and suspension culture (Gupta et al., 2014, 2015). However, there is little information about the effect of PGRs on SGs of in vitro-propagated Stevia shoots. The synthesis of SGs is initiated in the chloroplasts and takes place mainly in the leaves (Brandle and Telmer, 2007; Ladygin et al., 2008). The SGs production varies from clone to clone as well as depending on the physical conditions under which Stevia plants are regenerated. It is of particular interest to develop new S. rebaudiana varieties with a high SGs content and to obtain sufficient propagating and planting material which possesses this quality characteristic. The extracts with homogeneous and high secondary metabolite contents can be achieved in vitro under controlled physical conditions, using a suitable growth media. Furthermore, it is possible to achieve within a considerably short time a large number of, with regard to development and growth, uniform Stevia plantlets by in vitro propagation. Therefore, the purpose of the present study was to develop an efficient and economical in vitro propagation protocol for a selected clone line of S. rebaudiana. Furthermore, to determine the effect of some PGRs on the shoot growth and SGs production.

Materials and Methods
Stevia rebaudiana shoot cultures
In vitro clonally propagated plantlets, which were obtained from single seed descent seedlings of S. rebaudiana, were used as the...
plant material. To obtain a sufficient number of plants, nodal segments and shoot tips (with one axillary bud) of 4-week-old plantlets were transferred to glass tubes each containing 10 mL of WPM (Lloyd and McCown, 1980) supplemented with 3% (w/v) sucrose, and solidified with 0.65% (w/v) plant agar (Duchefa Biochemie B.V., Haarlem, The Netherlands). The pH of all media was adjusted to 5.8 before the addition of the gelling agent, and they were autoclaved at 121 °C at 1.04 kg·cm⁻² for 15 min. For plant multiplication, they were transferred every 4 weeks to fresh solidified medium.

**PGR application and culture conditions**

To investigate the effect of different PGRs on shoot proliferation and growth as well as on SGs production, two different explant types as specified above, were used. The explants were cultured in glass tubes each containing 10 mL of WPM supplemented with various concentrations of different PGRs, 3% (w/v) sucrose, and 0.65% (w/v) agar (Table 1).

The cultures were incubated in a growth room under 26 ± 1 °C in a cool white fluorescent light (50 μmol·m⁻²·s⁻¹) and a 16-h photoperiod for 4 weeks.

### Acclimatization

Acclimatization of the in vitro-rooted plantlets was carried out in a climate chamber under 25 ± 1 °C, 70% humidity, 50 μmol·m⁻²·s⁻¹ irradiance, and 16-h photoperiod conditions. Plantlets, 8–10 cm long with 10–14 leaves and well-developed roots, were carefully removed from the culture vessels, and their roots were washed with water. Each plantlet was transferred to a 5-cm diameter pot containing a 1:3 peat:loam mixture and was completely covered with perforated transparent bags to retain humidity. These plants were ventilated for 10 min by removing the bags from the plants once a day for a period of 2 weeks. The acclimation bags were completely removed at the end of the 2-week period. The plants were then transferred from the climate chamber to the greenhouse conditions 3 weeks after the beginning of acclimatization and were maintained there for a period of 5 weeks. Completely acclimatized plants were transferred to field conditions at the end of 8 weeks.

**SGs analysis**

**Chemicals.** HPLC grade acetonitrile and methanol were purchased from Merck (Darmstadt, Germany). Milli-Q water, used for the analysis, was obtained from an in-house ultrapure water system (Sartorius Arium 611; Sartorius Stedim Biotech, Göttingen, Germany). Steviol glycosides were isolated from the leaves of *S. rebaudiana* by our research group with a purity of > 90%.

**HPLC sample preparation.** Approximately 75 mg of leaf samples were sonicated three times with 5 mL methanol for 20 min. The clear extracts were combined and diluted with methanol to 20 mL. Before HPLC analysis, the samples were filtered with a 0.45-μm polytetrafluoroethylene filter (Sartorius AG, Göttingen, Germany) to remove nondissolved particles.

**HPLC-evaporative light-scattering detector conditions.** HPLC-evaporative light-scattering detector (ELSD) analyses were performed with a Thermo Surveyor Plus HPLC (Thermo Fisher Scientific, Waltham, MA) equipped with a quaternary pump, an autosampler, and a SofTAg HPLC–evaporative light-scattering detector (ELSD) system (SofTAg Corporation, Westminster, CO). As the analytical column, an apHlera NH₂ column (150 mm × 4.6 mm, particle size 5 μm, Sigma Aldrich, Germany) was used. An acetonitrile/water mixture (75:25 v/v) served as isotropic mobile phase. The total run time was 10 min at a flow rate of 1 mL·min⁻¹ and column temperature was 40 °C. LC-MS experiments were performed in electrospray ionization (ESI) (+) mode with capillary temperature set to 320 °C, vaporizer temperature set to 350 °C, and spray voltage set to 3000 V. Pressures of sheath and aux gas of probe were set to 50 and 25 psi, respectively. All the samples were scanned between 600 and 1200 amu to qualitatively identify *Stevia* glycosides. Existence of *Stevia* glycosides were examined by scanning their molecular mass spectrum.

### Table 1. Effects of different types and concentrations of PGRs and explant types on the shoot proliferation in *Stevia rebaudiana*.

| Concentrations of PGRs (μM) | Mean number of shoots/explant ± se | Mean length (cm)/explant ± se | Mean number of nodes/explant ± se |
|----------------------------|-----------------------------------|--------------------------------|----------------------------------|
| BA | Kn | TDZ | IAA | NAA | Nodal explant | Shoot tip explant | Nodal explant | Shoot tip explant |
| 2.22 | — | — | — | — | 2.00 ± 0.00 c | 1.20 ± 0.03 h | 7.88 ± 0.06 d | 12.44 ± 0.02 ab |
| 2.44 | — | — | — | — | 6.51 ± 0.08 bc | 6.00 ± 0.06 c | 3.57 ± 0.00 f | 4.03 ± 0.04 efg |
| 8.88 | — | — | — | — | 4.98 ± 0.13 d | 6.04 ± 0.06 c | 2.28 ± 0.03 fgh | 2.23 ± 0.07 hi |
| 17.76 | — | — | — | — | 4.49 ± 0.08 d | 5.40 ± 0.16 c | 1.61 ± 0.04 gh | 1.67 ± 0.05 i |
| 9.29 | — | — | — | — | 2.22 ± 0.06 e | 2.82 ± 0.04 efgh | 11.36 ± 0.11 b | 11.30 ± 0.18 b |
| 18.59 | — | — | — | — | 1.91 ± 0.01 c | 1.20 ± 0.02 h | 7.22 ± 0.06 d | 6.43 ± 0.07 d |
| 2.27 | — | — | — | — | 2.69 ± 0.05 e | 2.44 ± 0.12 efg | 3.19 ± 0.12 d | 3.06 ± 0.06 gh |
| 4.54 | — | — | — | — | 2.30 ± 0.03 e | 2.64 ± 0.01 ef | 2.78 ± 0.04 gh | 2.46 ± 0.03 hi |
| 9.08 | — | — | — | — | 1.98 ± 0.01 e | 1.98 ± 0.04 fgh | 1.87 ± 0.04 gh | 1.95 ± 0.05 i |
| 18.16 | — | — | — | — | 2.00 ± 0.01 e | 1.99 ± 0.09 fgh | 1.38 ± 0.04 gh | 1.99 ± 0.06 h |
| 2.22 | — | — | 2.85 | — | 6.27 ± 0.03 c | 6.16 ± 0.01 c | 5.39 ± 0.03 c | 4.47 ± 0.06 ef |
| 2.22 | — | — | 5.71 | — | 1.91 ± 0.01 c | 2.24 ± 0.04 efg | 11.36 ± 0.11 b | 11.30 ± 0.18 b |
| 2.22 | — | — | 2.69 | — | 6.29 ± 0.03 c | 10.02 ± 0.13 a | 5.24 ± 0.06 efg | 5.47 ± 0.06 ef |
| 2.22 | — | — | 5.37 | — | 8.00 ± 0.09 a | 9.33 ± 0.09 a | 5.13 ± 0.11 c | 5.09 ± 0.09 e |
| 2.32 | — | — | 2.85 | — | 7.51 ± 0.10 ab | 6.42 ± 0.07 bc | 5.17 ± 0.12 e | 7.17 ± 0.20 cd |
| 2.32 | — | — | 5.71 | — | 4.69 ± 0.03 c | 4.09 ± 0.12 d | 7.36 ± 0.24 d | 7.86 ± 0.30 c |
| 2.32 | — | — | 2.69 | — | 1.82 ± 0.03 fgh | 12.70 ± 0.05 a | 13.16 ± 0.02 a | 13.07 ± 0.03 a |
| 2.32 | — | — | 5.37 | — | 1.89 ± 0.04 c | 12.54 ± 0.19 a | 13.07 ± 0.03 a | 13.07 ± 0.03 a |

BA = 6-benzyladenine; IAA = indoleacetic acid; Kn = kinetin; NAA = naphthaleneacetic acid; PGR = plant growth regulator; TDZ = thidiazuron.

Values represent mean ± se of three replications, with 15 explants per replicate. Values (mean ± se) sharing the same letter in the same column is not significantly different at P ≤ 0.01, according to least significant difference test. Data were recorded after 4 weeks of culture.
All PGR treatments suppressed the root formation and induced callus formation at the base of explants (Fig. 2B), whereas on the PGR-free control medium, no callus formation occurred and normal roots were developed. The percentage of explants which showed callus formation ranged between 45.56% (18.16 μM TDZ) and 100% (all concentrations of BA and combinations of BA with auxins, as well as 4.65 μM Kn and 2.32 μM Kn + 2.85 μM IAA). The callus diameter ranged between 0.42 cm (9.08 μM TDZ) and 1.06 cm (2.22 μM BA + 2.85 μM IAA) (Table 2).

PGR-supplemented media caused the formation of significant chlorosis and necrosis (Fig. 2C and D). WPM supplemented with 2.32 μM Kn + 2.85 μM IAA or 5.37 μM NAA induced the highest percentage of chlorosis, and WPM supplemented with 4.65 μM Kn or 8.88 μM BA caused the highest necrosis formation, respectively. In general, the presence of TDZ and combinations of cytokinin with auxin in the media induced a high percentage of chlorosis (Table 2).

Shoots grown on the PGR-free control medium proved to be the most suitable for in vitro propagation, since they showed, in general, a better development with respect to important criteria such as shoot length, length of internodes, and leaf size. In addition, no callus was formed at the base of the explants and root as well as shoot formation occurred simultaneously. Furthermore, neither necrosis nor chlorosis was detected (Fig. 1A).

The plantlets (8–10 cm long with 10–14 leaves and well-developed roots) (Fig. 3A) obtained from PGR-free control medium (WPM) were transferred to soil and were acclimatized over a period of 5 weeks (Fig. 3B and C). Completely acclimatized plants were transferred to field conditions at the end of 8 weeks (Fig. 3D).
short and thin with smaller and abnormal leaves (Fig. 1D). Kataeva et al. (1991) reported that the addition of PGRs to the medium might alter the hormone content and/or balance in explants, promoting abnormal plant growth.

On WPM medium containing 2.22 μM BA + 2.69 μM NAA, the number of shoots per shoot tip explant was 8.35-fold higher compared with the number obtained on the PGR-free control medium. The positive effect of BA + NAA combinations on shoot proliferation of
Table 2. Effects of different types and concentrations of PGRs on the in vitro shoot growth of *Stevia rebaudiana*.

| Concentrations of PGRs (μM) | Mean callus diam (cm/explant ± SE) | Percentage of explants producing callus at base ± SE | Percentage of explants showing chlorosis ± SE | Percentage of explants showing tip and leaf necrosis ± SE |
|---------------------------|----------------------------------|-----------------------------------------------|-----------------------------------------------|--------------------------------------------------|
| BA  Kn  TDZ  IAA  NAA | 0.00 ± 0.00 b  0.00 ± 0.00 g | 0.00 ± 0.00 l | 0.00 ± 0.00 k | 0.00 ± 0.00 k |
| 2.22  —  —  —  — | 1.01 ± 0.00 a  100.00 ± 0.00 a | 75.12 ± 0.97 c | 3.94 ± 0.20 fghij | |
| 4.44  —  —  —  — | 1.02 ± 0.01 a  100.00 ± 0.00 a | 20.72 ± 0.53 j | 17.63 ± 0.95 b | |
| 8.88  —  —  —  — | 0.94 ± 0.02 a  100.00 ± 0.00 a | 20.78 ± 0.78 j | 22.61 ± 1.15 a | |
| 17.76  —  —  —  — | 0.80 ± 0.03 a  100.00 ± 0.00 a | 30.86 ± 0.59 h | 15.48 ± 0.79 b | |
| —  —  —  —  — | 0.45 ± 0.00 ab  88.88 ± 0.19 c | 7.48 ± 0.47 k | 1.73 ± 0.00 jk | |
| —  —  —  —  — | 0.46 ± 0.00 ab  100.00 ± 0.00 a | 32.93 ± 0.42 h | 24.48 ± 0.08 a | |
| —  —  —  —  — | 0.76 ± 0.01 a  96.67 ± 0.67 b | 23.74 ± 1.43 ij | 7.30 ± 0.98 cdfg | |
| —  —  —  —  — | 0.65 ± 0.01 ab  96.67 ± 0.33 b | 9.31 ± 1.41 k | 6.16 ± 0.26 cddefghij | |
| —  —  —  —  — | 0.77 ± 0.01 a  76.67 ± 0.58 d | 90.25 ± 0.49 ab | 6.90 ± 0.42 cdefghij | |
| —  —  —  —  — | 0.65 ± 0.01 ab  78.89 ± 1.02 d | 82.97 ± 0.75 cd | 5.85 ± 0.63 c | |
| —  —  —  —  — | 0.42 ± 0.01 ab  52.22 ± 0.69 e | 86.43 ± 0.44 bc | 0.00 ± 0.00 k | |
| —  —  —  —  — | 0.45 ± 0.01 ab  45.6 ± 0.84 f | 82.68 ± 1.32 cd | 6.67 ± 0.38 cdefghij | |
| —  —  —  —  — | 0.46 ± 0.00 ab  100.00 ± 0.00 a | 69.76 ± 0.95 f | 4.14 ± 0.8 b | |
| —  —  —  —  — | 2.22 ± 0.01 a  78.89 ± 0.01 g | 27.98 ± 0.49 h | 5.77 ± 0.44 cdefghij | |
| —  —  —  —  — | 2.22 ± 0.01 ab  52.22 ± 0.69 e | 86.43 ± 0.44 bc | 0.00 ± 0.00 k | |
| —  —  —  —  — | 2.22 ± 0.01 ab  100.00 ± 0.00 a | 75.12 ± 0.97 c | 2.97 ± 0.93 hij | |
| —  —  —  —  — | 2.22 ± 0.01 ab  52.22 ± 0.69 e | 86.43 ± 0.44 bc | 0.00 ± 0.00 k | |
| —  —  —  —  — | 2.22 ± 0.01 ab  75.12 ± 0.97 c | 2.97 ± 0.93 hij | |
| —  —  —  —  — | 2.22 ± 0.01 ab  52.22 ± 0.69 e | 86.43 ± 0.44 bc | 0.00 ± 0.00 k | |

**BA =** 6-benzyladenine; **IAA =** indoleacetic acid; **Kn =** kinetin; **NAA =** naphthaleneacetic acid; **PGR =** plant growth regulator; **TDZ =** thidiazuron.

Values represent mean ± SE of three replications, with 15 explants per replicate. Values (mean ± SE) sharing the same letter in the same column is not significantly different at P = 0.01, according to least significant difference test. Data were recorded after 4 weeks of culture.

*S. rebaudiana* is consistent with other studies (Bondarev et al., 2003; Giridhar et al., 2010; Soliman et al., 2014). Bondarev et al. (2003) achieved a 1.5-fold higher shoot number on MS medium supplemented with 0.1 mg·L⁻¹ BA and NAA compared with the number obtained on the PGR-free control medium.

Up to now, there are a lot of reports regarding in vitro propagation of *Stevia* using stem tips with a few leaf primordia, nodal or leaf explants (Aman et al., 2013; Sivaram and Mukundan, 2003; Tamura et al., 1984; Thiyagarajan and Venkatachalam, 2012; Yang et al., 1981). The highest number of shoots per explant obtained from micropropagation studies of *S. rebaudiana* varies considerably such as 40 shoots/explant (Tamura et al., 1984), 11.2 shoots/explant (Sivaram and Mukundan, 2003), 23.4 shoots/explant (Hwang, 2006), 8.75 shoots/explant (Ahmed et al., 2007), 83.2 shoots/explant (Sairkar et al., 2009), 28 shoots/explant (Giridhar et al., 2010), 40.54 shoots/explant (Satpathy and Das, 2010), 15.69 shoots/explant (Thiyagarajan and Venkatachalam, 2012), 60 shoots/explant (Lata et al., 2015). In the present study, the highest number of shoots per explant was 10.02. This variety among the results may be due to used explant, basal medium and PGRs and the endogenous level of PGRs in used explants.

In previous publications, it is stated that for rooting of in vitro regenerated shoots of *S. rebaudiana* an additional rooting medium is necessary. For this purpose, different media were used: Linsmaier and Skoog medium with 0.1 mg·L⁻¹ NAA (Tamura et al., 1984), half-strength MS medium with 4.90 μM indole-3-butyric acid (IBA) (Sivaram and Mukundan, 2003), MS medium with 0.1 mg·L⁻¹ IAA (Ahmed et al., 2007), MS medium with 2.0 mg·L⁻¹ IBA (Hwang, 2006). We were able to develop a one-step in vitro propagation protocol. On the PGR-free control medium (WPM) the development of shoots and roots occurred simultaneously and healthy and well developed plantlets were obtained. The plantlets showed no callus formation at the base of the explants and also no chlorosis and necrosis. At the same time, a good rooting was achieved (Fig. 3A). This result is in accordance with the results reported by Ibrahim et al. (2008). Regeneration in the absence of PGRs could both reduce the costs and make the process more efficient as the risk of somaclonal variations is reduced (Ibrahim et al., 2008).

The occurrence of necrosis and chlorosis to a considerable extent (Fig. 2C and D) and
the formation of callus instead of roots (Fig. 2B) were significant adverse effects, when PGRs containing media were used. Cassells and Curry (2001) associated the occurrence of necrosis with basal callus formation and reported that callus formation at the basal part of the explants caused the loss of the apical dominance of the shoots. The formation of chlorosis and necrosis in the shoots might be explained in a similar manner. The lack of root formation, which is one of the most effective factors in endogenous cytokinin biosynthesis, might lower the internal cytokinin levels, ceasing cell division and causing necrosis in the apical meristem (Kataeva et al., 1991). In the present study, the explants formed shoots as well as callus tissue at the basal part of the explants, therefore the minerals and PGRs in the medium were needed for the growth of two different tissue formations and thus could have led to an imbalance in their distribution. Furthermore, besides an imbalanced use of minerals and PGRs for the callus formation, has probably also obstructed the transport of minerals to the shoots, which on the other hand, most likely induced the formation of chlorosis and necrosis. Tamura et al. (1984) reported that callus formation at the basal parts of shoots was prevented by culturing the stem tips of shoots in medium containing only cytokinins, whereas on media containing both Kn and NAA, callus formation was promoted. The aforementioned result does not support our findings. As the usage of cytokinins individually did not prevent callus formation in the basal part of in vitro shoots. The other result agrees with the results of the present study. Since Kn + NAA combination caused callus formation in the basal part of in vitro shoots.

**Effect of PGRs on production of SGs.** Internal factors such as PGRs were reported to affect the secondary metabolite production along with external factors such as temperature, light, pH and salt concentration. Consequently, secondary metabolite production might be promoted by the addition of suitable PGRs to the medium, modification of the composition of the growth medium and by alteration of the physical conditions of the culture. Therefore, we investigated the effect of various cytokinins and various cytokinin–auxin combinations on the secondary metabolite production of *Stevia rebaudiana*. Previous investigations regarding the content of secondary metabolites of *Stevia*, focused on the determination of SGs in the donor plant and the in vitro plant material, regenerated from

![Fig. 3. Acclimatization of *Stevia rebaudiana* plantlets: rooted plantlets on woody plant medium (WPM) (A); acclimatized plantlets (B); root development in 5-cm pots containing a 1:3 peat to loam mixture (C); hardened plants in the field (D).](image)

Table 3. Effects of different types and concentrations of PGRs on the stevioside and rebaudioside A content (mg·g⁻¹ dry weight) in leaves of in vitro grown *Stevia rebaudiana*.

| Concentrations of PGRs (µM) | Stevioside (mg·g⁻¹ dry wt) | Rebaudioside A (mg·g⁻¹ dry wt) |
|-----------------------------|-----------------------------|-------------------------------|
| BA | Kn | TDZ | IAA | NAA | | |
| — | — | — | — | — | — | 34.00 | 12.20 |
| 2.22 | — | — | — | — | — | 8.40 | ND |
| 4.44 | — | — | — | — | — | 9.50 | ND |
| 8.88 | — | — | — | — | — | 10.80 | ND |
| 17.76 | — | — | — | — | — | 4.60 | ND |
| — | 2.32 | — | — | — | — | 13.60 | ND |
| — | 4.65 | — | — | — | — | 10.10 | ND |
| — | 9.29 | — | — | — | — | 9.80 | ND |
| — | 18.59 | — | — | — | — | 7.00 | ND |
| — | — | 2.27 | — | — | — | 20.60 | ND |
| — | — | 4.54 | — | — | — | 26.40 | ND |
| — | — | 9.08 | — | — | — | 13.80 | ND |
| — | — | 18.16 | — | — | — | 7.20 | ND |
| 2.22 | — | — | 2.85 | — | — | 7.50 | ND |
| 2.22 | — | — | 5.71 | — | — | 9.80 | ND |
| 2.22 | — | — | — | 2.69 | — | 16.20 | ND |
| 2.22 | — | — | — | 5.37 | — | 22.30 | ND |
| — | 2.32 | — | 2.85 | — | — | 14.30 | ND |
| — | 2.32 | — | 5.71 | — | — | 15.60 | ND |
| — | 2.32 | — | — | 2.69 | — | 21.10 | ND |
| — | 2.32 | — | — | 5.37 | — | 13.20 | ND |

BA = 6-benzyladenine; IAA = indoleacetic acid; Kn = kinetin; NAA = naphthaleneacetic acid; ND = not determined; PGR = plant growth regulator; TDZ = thidiazuron.
the donor plants. Bondarev et al. (2001) found in different plant material of a *S. rebaudiana* clone, following stevioside and rebaudioside-A contents (mg g⁻¹ dry matter): 24.9 and 12.0 mg g⁻¹ in leaves of intact plants, 3.4 and 1.0 mg g⁻¹ in leaves of in vitro plants, 0.015 mg g⁻¹ and a trace amount in material of a suspension culture, 0.060 and 0.023 mg g⁻¹ in morphogenic callus and 0.387 and 0.112 mg g⁻¹ in shoots, regenerated in vitro from callus. They also established a callus culture on MS medium supplemented with 1 mg L⁻¹ NAA and 0.5 mg L⁻¹ 6-benzyl aminopurine in continuous darkness; however, they found no stevioside and rebaudioside-A in this culture. SGs are thought to be produced in the chloroplasts and therefore it comes as no surprise that callus cultures grown in the dark, by Bondarev et al., were found not to contain SGs. We found in leaves of shoots grown in vitro on WPM (in light), containing the same PGRs in the same concentration as indicated by Bondarev et al. (2001) a stevioside content of 22.3 mg g⁻¹ DW. Ladygin et al. (2008) found a positive correlation between the development and activity of the photosynthetic apparatus in *Stevia* leaves and the biosynthesis of diterpenoid SGs. The initiation of the bio-synthesis of SGs is reported to take place in chloroplasts (Brandle and Telmer, 2007; Ladygin et al., 2008). Heterotrophic callus cells grown in the dark held almost no pigments and they did not synthesize SGs (Ladygin et al., 2008). Yamazaki and Flores (1991) suggested that SGs are synthesized in the leaves not in the roots of *S. rebaudiana*. Reis et al. (2011) established successfully an adventitious root culture of *S. rebaudiana* in a roller bottle system, but no stevioside and other major SGs were synthesized in this culture. Aman et al. (2013) investigated the effect of some PGRs and different concentrations of agar on the dulcoside-A, stevioside and rebaudioside-A production in in vitro shoots of *S. rebaudiana*. They reported that the highest content of dulcoside-A (71.8 µg g⁻¹ DW), stevioside (82.48 µg g⁻¹ DW) and rebaudioside A (12.35 µg g⁻¹ DW) were observed in shoots grown on medium, containing a combination of BA and Kn (3.0 mg L⁻¹) with 3.5 g L⁻¹ agar, a combination of BA and Kn (3.0 mg L⁻¹) with 7.0 g L⁻¹ agar and on medium containing only BA (1.0 mg L⁻¹) with 7.0 g L⁻¹ agar, respectively. Even though the addition of BA to the culture medium stimulated the multiple shoot production, inhibited the biosynthesis of stevioside. Similar results regarding the effect of BA on the SGs formation were reported (Bondarev et al., 2003; Sivaram and Mukundan, 2003). In the present study, Kn limited both multiple shoot induction and secondary metabolite production in the leaves of *S. rebaudiana*. The addition of all PGRs, PGR combinations and PGR concentrations tested, resulted compared with the content achieved on the PGR-free control medium, in a considerable lower stevioside content in the leaves. The conversion of stevioside into rebaudioside A is the terminal step in the metabolic pathway of the SGs (Brandle and Telmer, 2007). The addition of BA and Kn to the culture media had a negative effect on the stevioside formation and the conversion of stevioside into rebaudioside A was completely inhibited. Since BA promotes the multiple shoot formation and Kn the growth, it can be assumed that in response to the addition of these PGRs, an active primary metabolism is prioritized by the plant and consequently the formation of secondary metabolites is reduced. Among the all PGRs applied, the highest stevioside content (26.40 mg g⁻¹ DW) were detected in shoots grown on WPM supplemented with 4.54 µM TDZ. There has been no published data regarding the effect of TDZ on the stevioside A content in *S. rebaudiana*. Callus tissue, as it was formed at the base of shoots, grown on TDZ-containing media, can obstruct the vascular connection between shoot and medium and therefore result in a reduction or cessation of passageways for minerals and PGRs toward the shoots. With the time this leads to a decrease of the internal hormone level, which in turn inhibits the apical cell division and limits the plant growth. As a result chlorosis followed by necrosis might occur, as it was observed. At this stage, the plant growth stops due to the stress. Therefore, the remaining resources might be supplied to the secondary metabolism and thus lead to an increased formation of secondary metabolites. This might explain the high contents of rebaudioside A, as they were found. In the first step of SGs pathway, steviol convert into steviolmonoside and in the second step, steviolmonoside convert into steviolbioside or rubusoside. After the glycosylation of steviolbioside, stevioside, and rebaudioside A, stevioside, rebaudioside A, and rebaudioside D occur, respectively (Prakash et al., 2014). It is also reported that there is a wide range of variation in the four main glycosides and a positive correlation between dulcoside A and stevioside and a negative correlation between stevioside and rebaudioside A. These correlations result from the biosynthetic relationships between the individual glycosides (Brandle et al., 1998). According to Brandle and Telmer. (2007), rebaudioside synthesis is inhibited with the preferred synthesis of steviolbioside. In the present study, steviolbioside was detected qualitatively in the leaves of shoots grown on selected media (WPM supplemented with 2.27 µM TDZ, 4.54 µM TDZ, 2.22 µM BA + 2.69 µM NAA, 2.22 µM BA + 5.37 µM NAA, 2.32 µM Kn + 5.71 µM IAA or 2.32 µM Kn + 2.69 µM NAA) as well as rubusoside and no rebaudioside A was determined. This may be because the SGs pathway was directed rubusoside synthesis and thus rebaudioside A synthesis was slowed down. In addition to stevioside, steviolbioside, and rubusoside, dulcoside A were also detected. The determination of dulcoside A in the selected media can be explained by the presence of stevioside. Conclusion A simple and low-cost in vitro propagation procedure was developed for *S. rebaudiana*. We were able to develop a one-step in vitro propagation protocol. On the PGR-free WPM, shoots and roots were formed simultaneously and well-developed plants were obtained without the use of an additional rooting medium or the use of PGRs. Furthermore, the highest stevioside content was determined in leaves of plants regenerated on this medium. All plants obtained from PGR-free control medium were successfully acclimatized in a glasshouse and then plants were transferred to the field. This in vitro propagation procedure has been adapted by a commercial plant tissue culture laboratory. literature Cited Aickgoz, N., E. Ilker, and F. Gokcen. 2004. Analysis methods of biological researches. Seed Technology Center, Publication 2, Izmir, Turkey. Ahmed, M.B., M. Salahin, R. Karim, M.A. Razvy, M.M. Hannan, R. Sultana, M. Hossain, and R. Islam. 2007. An efficient method for in vitro cloning propagation of a newly introduced sweetener plant (*Stevia rebaudiana*) Bertoni in Bangladesh. American-Eurasian J. Sci. Res. 2:121–125. Aman, N., F. Hadi, S.A. Khalil, R. Zamin, and N. Ahmad. 2013. Efficient regeneration for enhanced stevioside glycosides production in *Stevia rebaudiana* (Bertoni). C. R. Biol. 336:486–492. Bondarev, N., O. 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