Hairy root induction and secondary metabolite production in *Perovskia abrotanoides* Karel

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Abstract:

Induction of hairy roots in *Perovskia abrotanoides* Karel, a medicinally important member of the Lamiaceae family, was evaluated using different strains of *Agrobacterium rhizogenes* and various carbon sources. An efficient *Agrobacterium*-mediated transformation system was established for the first time for this plant species using the bacterial strains of ATCC15834, TR105, and R1000. Meanwhile, the effects of acetosyringone and carbon sources on both rooting and root biomass production were investigated and the potential of hairy root cultures for producing tanshinones was explored. Hairy root induction in the nodes of intact seedlings was found to be successful with all the bacterial strains tested but R1000 was found to be more effective than TR105. The highest transformation frequency belonged to the nodes infected with ATCC15834 (47.33%). Moreover, transformation frequency increased by up to 60.99% when 100 µM of acetosyringone was added. The strain ATCC15834 recorded the highest root dry and fresh weights per 50 ml of the nutrient medium (0.149 and 1.996 g, respectively) and root length (34.45 cm) in a medium containing 3% sucrose. However, R1000 induced maximum lateral root density in hairy roots (7.1 branches cm⁻¹) in a medium containing 3% glucose. PCR analysis using rolC specific primer with an amplicon of 612 bp confirmed the transgenic nature of the hairy roots. Finally, HPLC analysis revealed that the cryptotanshinone and tanshinone IIA levels in the hairy roots induced by TR105 (clone T7) were 53.17±0.26 and 14.48±0.30 µg g⁻¹ DW, respectively.


Introduction:

Medicinal plants have of long been used as important therapeutic agents. Almost one fourth of the drugs presently manufactured contain plant extracts or active compounds obtained from the plants (Yaniv, 2005). *Perovskia abrotanoides* Karel (a perennial herb from the Lamiaceae family), grown in Iran, Afghanistan, Pakistan, and Turkmenistan (Rechinger, 1982), has been used in Iranian traditional medicine for the treatment of leishmaniasis (Moallem and Niapour, 2008). In recent years, the anti-plasmodial, anti-microbial, anti-nociceptive, anti-inflammatory, and cytotoxic activities of this species have also been reported (Nassiri-Asl et al., 2002; Hosseinzadeh and Amel, 2001; Sairafianpour et al., 2001; Ashraf et al., 2014). However, while few scientific reports are available on the natural products from this species, most studies have focused on the essential oils obtained from their aerial parts (Mahboubi and Kazempour, 2009; Safaei-Ghomí and Batoollí, 2010; Ashraf et al., 2014). Moreover, phytochemical analysis has demonstrated that the roots of *P. abrotanoides* are rich sources of diterpenoid quinone compounds known as tanshinones (Sairafianpour et al., 2001; Zaker et al., 2015), which reportedly have such biological properties as anti-microbial (Lee et al., 1999), anti-inflammatory (Jang et al., 2003), antioxidant (Cao et al., 1996), anti-diabetes (Kim et al., 2007), and anti-cancer (Liu et al., 2009; Chiu et al., 2010; Pan et al., 2010) effects.

The commercially used secondary metabolites are generally higher-value, lower-volume products than the primary ones (Balandrin et al., 1985). These metabolites are generally accumulated by plants in small quantities often at certain developmental stages and under given environmental conditions. The pharmaceutical substances of *P. abrotanoides* (e.g. tanshinones) are typically produced in plants roots; thus, hairy root cultures may enhance the production and accumulation of these metabolites.

Since chemical synthesis of natural products is not readily available, an alternative is to produce them through cell-tissue culture technology (Vanisree et al., 2004). In this regard, particular emphasis is commonly laid on hairy root culture as a means of enhanced production of secondary metabolites for commercial applications (Kong et al., 2003). These cultures generally have a higher capacity than cell suspension cultures or even roots of intact plants for synthesizing secondary metabolites (Giri and Narasu, 2000; Ono and...
Tian, 2010). Moreover, hairy root cultures are not only more reliable in terms of genetic stability but are characterized by faster growth and easier maintenance in a hormone-free medium.

Hairy root growth and biomass productivity are affected by different factors. For instance, it is established that addition of a carbon source to the cultivation medium is necessary for the growth of hairy roots. Study has shown that the production of most biologically active compounds through hairy roots is affected by the type and concentration of the carbon source present in the nutrient media (Kochan et al., 2014). Moreover, plant growth, development, metabolism, and gene expression are reportedly affected by sugars (monosaccharide and disaccharide), as signal molecules (Praveen and Murthy, 2012; Petrova et al., 2015).

To date, there have been no published reports on hairy root induction in *P. abrotanoides* and the effects of *Agrobacterium* strain and carbon sources on its hairy root growth. In this study, hairy root cultures of *P. abrotanoides* were established using three different strains of *Agrobacterium rhizogenes* and different carbon sources to determine the best carbon source for the production of tanshinones.

**Materials and methods:**

**Plant material:** Mature seeds of *P. abrotanoides* Karel were collected in September from wild-growing plants in Kalat, Razavi Khorasan Province, Iran. The plant was identified in the herbarium of Ferdowsi University of Mashhad, Mashhad (No. 39299). The seeds were surface sterilized with 20% NaOCl (V/V) for 4 minutes and rinsed three times with sterilized water before they were aseptically germinated on the hormone-free half-strength Murashige and Skoog (MS) medium (Murashige and Skoog, 1962) with 3% sucrose (W/V) and 0.7% agar (W/V) in the culture room at 25±1 °C under darkness. The pH of the medium was adjusted to 5.7 before autoclaving at 121 °C for 15 min. After germination, the cultures were kept at 25±1 °C during 16/8 h light/dark photoperiods with an irradiance level of 45 µmol m⁻² s⁻².

**Growth of Agrobacterium rhizogenes:** The wild type of *A. rhizogenes* strains including ATCC15834, TR105, and R1000 were cultured in a solid Yeast Mannitol Broth medium (YMB) containing 0.2 g L⁻¹ MgSO₄·7H₂O, 10 g L⁻¹ mannitol, 0.5 g L⁻¹ K₂HPO₄, 0.4 g L⁻¹ yeast extract, and 0.1 g L⁻¹ NaCl adjusted to pH=7. Briefly, a single bacterial colony was inoculated in 15 mL of liquid YMB medium and grown overnight at 28 °C on a shaker incubator (100 rpm) to mid-log phase (OD₆₀₀=0.6-0.8) (Vincent, 1970). The bacterial cells were then collected by centrifugation (Hermle Z230A, Germany) at 107g for 10 minutes and resuspended at a cell density of OD₆₀₀=0.6-0.8 in a liquid inoculation medium (half-strength MS medium containing 3% sucrose).

**Induction of hairy roots in *P. abrotanoides* using *A. rhizogenes*:** *In vitro* grown (45-day-old) *P. abrotanoides* seedlings were used for transformation with the three *A. rhizogenes* strains (ATCC15834, TR105, and R1000). In an early experiment, excised leaf and internode explants from the *in vitro* seedlings and nodes from intact seedlings were inoculated using insulin syringes with and without acetylsyringone (100 µM) (Wawrosch et al., 2014). Infected leaf and internode explants together with whole seedlings inoculated at their nodes were then cultured on a half-strength MS medium with 3% sucrose and 0.7% agar for 72 hours at 25±1 °C under 16 h day⁻¹ light. After 3 days of co-cultivation, the shoots were transferred to an agar-solidified half-strength MS medium containing 500 mg L⁻¹ cefotaxime to remove residual bacteria. Subsequently, they were sub-cultured in a half-strength MS medium containing 300 mg L⁻¹ cefotaxime after 10 days (Yao et al., 2016). Controls consisted of explants and seedlings treated similarly, except that they were not co-cultivated with *A. rhizogenes*. Hairy roots emerged from the nodes of intact seedlings after 6 to 8 days. The percentage of hairy root induction was recorded after two months. In cases where no hairy roots initiated from the leaf and internode explants irrespective of the bacterial strain and the presence or absence of acetylsyringone, they were excluded from the experiment.

**Establishment of hairy root cultures:** Induced hairy root lines produced with *A. rhizogenes* ATCC15834, TR105 and R1000 strains, were aseptically dissected in 2 cm lengths from the inoculated seedlings and cultured in 200 mL flasks containing 50 mL of a liquid half-strength MS medium with 200 mg L⁻¹ cefotaxime and 3% sucrose. All the cultures were maintained at 25±1 °C in the dark on a rotary shaker at 90 rpm. Isolated clones were sub-cultured every 4 days into a fresh liquid medium but with gradually reduced cefotaxime concentrations (from 200 to 100 and 50 µM) (Yao et al., 2016). After complete removal of cefotaxime, the cultures were transferred into a liquid half-strength MS medium with 3% sucrose. Hairy roots were maintained by sub-culturing into a fresh medium every 14 days. The root lines A9, R5, and T7 already transformed with ATCC15834, R1000, and TR105 strains, respectively, were used for further experiments.

**PCR Analysis of hairy roots:** Genomic DNA was extracted from 200 mg of both fresh hairy and untransformed roots using Sharp’s method (Sharp et al., 1988). Polymerase chain reaction (PCR) was performed to detect the rolC gene using the following primer sets: forward-5’-CTGCTGACATAAACTGTC-3’ and reverse-5’-TGCTTCGAGTTAGGGTACA-3’. The reaction mixture consisted of 0.9 µl of 1 Unit Taq polymerase, 0.6 µl of 0.2 mM dNTP, 1 µl of 0.5 mM of each primer, 2 µl of 100 ng template DNA, 1.5 µl of 2 mM MgCl₂ and 2.5 µl of 10X reaction buffer; the mixture was replenished up to 25 µl with sterilizing distilled water. The following procedure was used in the PCR reaction: initial denaturation at 95 °C for 4 min followed by 95 °C for 1 min, annealing of the primer at 55 °C for 30 seconds and amplification at 72 °C for 1
min; the procedure was repeated for 35 cycles with a final extension at 72 °C for 7 min (Wang et al., 1993). Amplification products were analyzed by electrophoresis on 1.2% agarose gel, stained with ethidium bromide, visualized, and documented using a Trans illuminator equipped with a gel documentation system (Major Science, USA). Plasmid DNA from the A. rhizogenes strain ATCC15834 was used as the positive control and DNA from the untransformed root served as the negative control.

Effect of carbon source on hairy root growth: Hairy roots induced by three strains of A. rhizogenes (clones A9, R5, and T7 transformed with ATCC15834, R1000 and TR105 strains, respectively) were cultured in a half-strength MS medium containing 3% sucrose, glucose, or sorbitol to investigate the effects of carbon sources on hairy root growth. All the cultures were kept at 25±1 °C on a rotary shaker (90 rpm) in the dark. The hairy roots were harvested after 50 days and fresh and dry weights, root length, and lateral root density (i.e., number of lateral branches per 1 centimeter root) were recorded.

Analysis of tanshinones in hairy roots: One hairy root clone of P. abrotanoides induced by A. rhizogenes strain TR105 which grew vigorously with lateral branches in the medium containing 3% sucrose was selected to determine its tanshinones content. Briefly, dried hairy roots were ground into powder and extracted with methanol (0.5 g/30 mL) at room temperature in a sonic water-bath for one hour. The extract was evaporated to dryness and dissolved in 1 mL methanol, centrifuged at 5000 rpm for 5 minutes, and subjected to HPLC analysis. HPLC was performed on a C18 column (250×4 mm, 5 µm particle size) using gradient elutions of acetonitrile/water (20:80 V/V to 95:5 V/V) as the mobile phase at a flow rate of 1 mL min⁻¹. UV detection was conducted at 254 nm. Cryptotanshinone and tanshinone IIA contents were detected and quantified against authentic standards obtained from Sigma.

Statistical analysis: A completely randomized design was used for all the experiments. Significance was determined by analysis of variance (ANOVA) using SPSS software and means were compared using the Duncan’s Multiple Range Test at the 5% significance level. Data were reported as means ±standard error.

Results:

Effects of different A. rhizogenes strains on hairy root induction: No hairy roots were induced from infected leaves or internodes regardless of the bacterial strain used and the presence or absence of acetosyringone. The fragments of leaves and internodes turned brown and gradually became necrotic and died. Results of A. rhizogenes inoculation on the nodes of intact P. abrotanoides seedlings after 60 days of in vitro culturing showed that all the tested strains were able to induce hairy roots even in the absence of acetosyringone. At the site of inoculation (nodes of in vitro P. abrotanoides seedlings), transformed hairy roots were successfully induced. The appearance and growth of hairy roots were monitored over a period of two months. The first ATCC15834-induced hairy roots emerged from the inoculated nodes after 6 days; in the case of R1000 and TR105-infected nodes, however, hairy roots appeared 8 days after inoculation (Figure 1). No hairy roots were induced in the control groups which had only been inoculated in a half-strength MS medium.

Hairy root formation percentages in the seedlings of P. abrotanoides transformed with ATCC15834, R1000, and TR105 strains of A. rhizogenes bacteria are reported in Table 1. It was found that bacterial strains had significant effects on hairy root formation (p≤0.05). The three strains used varied in their ability to infect P. abrotanoides seedlings. In the absence of acetosyringone, the highest percentage of root induction was achieved with ATCC15834 (47.33%) followed by R1000 (38.16%) and TR105 (28.99%) (Table 1). Finally, the two strains ATCC15834 and TR105 recorded significantly (p<0.05) different capabilities for inducing hairy roots.

Confirmation of transformation: The transgenic nature of hairy roots was confirmed by PCR using rolC specific primers. A 612 bp rolC expected size fragment was present only in hairy roots but absent in normal roots (Figure 2).

Effect of acetosyringone on hairy root induction: According to Table 1, in the presence of acetosyringon, different strains of A. rhizogenes displayed varying degrees of ability to induce hairy roots on the nodes of P. abrotanoides seedlings. The presence of 100 µM acetosyringone increased hairy root induction on plants inoculated with the strain ATCC15834 from 47.33% to 60.99%. The use of acetosyringone with R1000 and TR105 strains also increased transformation efficiencies (from 38.16 to 49.49 in R1000 and from 28.99 to 36.33 in TR105) although the increments were not significant when compared with acetosyringone-free treatments (Table 1).

Effect of carbon source on hairy root growth: In the present study, application of sucrose, rather than glucose or sorbitol (p≤0.05), in the culture medium resulted in increased dry weight, fresh weight, and length of transformed (hairy) roots of P. abrotanoides induced by all the strains of A. rhizogenes. It is evident from Figures 3, 4, 5, and 6 that not only were the highest fresh and dry weights (1.996 and 0.149 g, respectively) and maximum root length (34.45 cm) observed in the ATCC15834-transformed hairy roots grown in the half-strength MS medium supplemented with 3% sucrose but that glucose had a weaker effect on the growth (root length) and biomass (fresh and dry weights) of hairy roots than did sucrose (p≤0.05). The transformed roots grown in the medium containing 3% sorbitol had significantly lower (p≤0.05) weights, lengths, and lateral branchings in most cases. The lowest dry and fresh weights of hairy roots (0.222 and
Figure 1. Hairy roots induced from the nodes of *Perovskia abrotanoides* seedlings after inoculation by different *Agrobacterium rhizogenes* strains. A: 12 days after inoculation with ATCC15834 strain; B: 2 weeks after inoculation with ATCC15834 strain (using acetalosyringone); C: 9 days after inoculation with R1000 strain (using acetosyringone); D: 12 days after inoculation with TR105 strain (using acetosyringone). Black arrows indicate the hairy roots developed in the sites of inoculation.

Table 1. The effects of different strains of *Agrobacterium rhizogenes* and acetosyringone application on the percentage of hairy root formation in *Perovskia abrotanoides* seedlings, 60 days after inoculation.

<table>
<thead>
<tr>
<th>A. rhizogenes strain</th>
<th>Percentage of hairy root induction</th>
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<tr>
<td></td>
<td>Without acetosyringone (100 µM)</td>
</tr>
<tr>
<td>ATCC15834</td>
<td>47.33 ± 5.06bc</td>
</tr>
<tr>
<td>R1000</td>
<td>38.16 ± 4.13bcd</td>
</tr>
<tr>
<td>TR105</td>
<td>28.99 ± 2.52d</td>
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All data were represented as the means (n=20) ± standard deviation. Different script letters characterize significant differences in each column (or row) according to Duncan’s Multiple Range Test at p≤0.05.

Figure 2. PCR analysis of a 612 bp fragment of a *rolC* gene in *Agrobacterium rhizogenes*. Lane 1: molecular weight marker (1000 bp ladder); Lane 2: amplified band from the DNA of strain ATCC15834 (positive control); Lane 3: amplified band from the DNA of hairy roots induced by TR105 strain (clone T7); Lane 4: amplified band from the DNA of hairy roots induced by R1000 strain (clone R5); Lane 5: amplified band from the DNA of hairy roots induced by ATCC15834 strain (clone A9); Lane 6: the band is absent in non-transformed root (negative control).
Hairy root induction and secondary metabolite production

Figure 3. Growth of hairy roots induced by the ATCC15834 strain of *Agrobacterium rhizogenes* in ½ MS medium after 50 days culture under the dark condition. A: in the medium containing 3% (W/V) glucose; B: in the medium containing 3% (W/V) sucrose.

Figure 4. The effect of carbon sources on the fresh weight of hairy roots in *Perovskia abrotanoides*, induced by different strains of *Agrobacterium rhizogenes* after 50 days of culture. Data are means (n=3) ± standard deviation. Bars with different letters are significantly different from one another (Duncan’s Multiple Range Test at \( P \leq 0.05 \)).

Figure 5. The effect of carbon sources on the dry weight of hairy roots in *Perovskia abrotanoides*, induced by different strains of *Agrobacterium rhizogenes* after 50 days of culture. Data are means (n=3) ± standard deviation. Bars with different letters are significantly different from one another (Duncan’s Multiple Range Test at \( P \leq 0.05 \)).

0.025 g, respectively) were obtained when roots induced by TR105 were cultured in the medium supplemented with 3% sorbitol (Figures 4, 5, and 6). The highest lateral root density (7.1 branches cm \(^{-1} \)) belonged to the roots induced by R1000 and grown in the medium containing 3% glucose. The least lateral root branching (1 branch cm \(^{-1} \)) was recorded for TR105-transformed roots grown in a medium containing 3% sorbitol (Figure 7).

**HPLC Analysis of tanshinones in hairy roots:** HPLC analysis revealed cryptotanshinone and tanshinone IIA contents of 53.17±0.26 and 14.48±0.30 µg g \(^{-1} \) DW, respectively, in the TR105-induced hairy roots (clone T7). These data cannot be compared with those from the control seedlings (not inoculated with *A. rhizogenes*) since they did not produce any adventitious roots in the half-strength MS medium.

**Discussion:**
It has been well established that hairy root formation in different plant species greatly depends on the type of explant. Wound response has been found to be the most important factor affecting successful transformation. Different explants reportedly vary in their wound response (i.e., in the number of competent cells produced for transformation) (Potrykus, 1991). This is because explant cells differ in their DNA synthesis and
Figure 6. The effect of carbon sources on the length of hairy roots in *Perovskia abrotanoides*, induced by different strains of *Agrobacterium rhizogenes* after 50 days of culture. Data are means (n=3) ± standard deviation. Bars with different letters are significantly different from one another (Duncan’s Multiple Range Test at p≤0.05).

Figure 7. The effects of carbon sources on the density of lateral hairy roots in *Perovskia abrotanoides*, induced by different strains of *Agrobacterium rhizogenes* after 50 days of culture. Data are means (n=3) ± standard deviation. Bars with different letters are significantly different from one another (Duncan’s Multiple Range Test at p≤0.05).

Cell division due to differences in the physiological maturity of the cells (Karmarkar *et al*., 2001). The specificity of various strains of *A. rhizogenes* to induce hairy roots in different explants and tissues is related to their age, cell division ability, and hormonal status (Karmarkar and Keshavachandran, 2001).

Our results revealed that neither the leaves nor the internodes of *P. abrotanoides* were capable of forming hairy roots. Consistent with this finding, no hairy roots reportedly formed in the leaves or internodes of *Hollostemma ada-kodien* K. Schum (Karmarkar and Keshavachandran, 2001) and *Salvia officinalis* L. (Grzegorczyk *et al*., 2006) transformed with *A. rhizogenes*. The observation by Karmarkar *et al*., (2001) that, contrary to leaves and internodes, seedling hypocotyls and shoot buds induced hairy roots might have been due to their ability to produce greater numbers of wound-adjacent cells competent for regeneration and transformation. To this may be added the fact that wounds induce the production of certain inhibitory compounds and that not all the *A. rhizogenes* strains are capable of degrading these inhibitory compounds (Karmarkar *et al*., 2001). It has been accepted that leaves or plant aerial parts contain compounds that might affect the transformation process in a positive and/or negative way (Bivadi *et al*., 2014).

Transformation efficiency has been shown to be greatly influenced by bacterial strain (Jain and Sing, 2015). In our investigation, all the three strains of *A. rhizogenes* were able to induce hairy roots in the nodes of intact *P. abrotanoides* seedlings albeit they differed significantly in the percentage of hairy roots they induced. ATCC15834 was the most efficient *Agrobacterium* strain in inducing hairy roots. Similar results have indicated the higher susceptibility of certain explants from various plant species to the strain ATCC15834. Some of these include *Helicteres isora* (Kumar *et al*., 2014), *Hypericum perforatum* (Bivadi *et al*., 2014), *Plumbago indica* (Gangopadhyay *et al*., 2010), *Solenostemon scutellaroides* (Saleh and Thuc, 2009), *Ipomoea batatas* (Chandran and Potty, 2008) and *S. officinalis* (Grzegorczyk *et al*., 2006). Virulence of *A. rhizogenes* varies depending on its strain, which thereby influences its transformation efficiency (Brijwal and Tamta, 2015). Moreover, the hairy roots obtained by different strains of *A. rhizogenes* vary with respect to their morphology, growth rate, and potential for the production of secondary metabolites, which may be attributed to the different plasmids the *Agrobacteria* harbor (Giri and Narasu, 2000; Kim *et al*., 2002; Rhodes *et al*., 1989). It seems that agropine strains such as ATCC15834, R1000, TR105, LBA9402, and A4, whose
plasmids contain both TR and TL regions, are more effective than mannopine strains in transforming a wide range of plant species (Otani et al., 1993). The TR fragment of T-DNA delivers genes for auxin synthesis that triggers cellular division; hence, the ability of these strains to transform a wide range of species.

Wounding is a precondition for the genetic transformation process through Agrobacteria. It may not only help in the production of signal phenolic molecules such as acetosyringone but also enhance the availability of assumed cell-wall binding factors to the bacterium (Gelvin, 2000). Acetosyringone, as a phenolic inducer molecule, has been successfully applied to elevate transformation in Agrobacterium-mediated genetic transformation of plants due to the activation of bacterial virulence genes which is necessary for the delivery of T-DNA to plant tissues (Brijwal and Tamta, 2015; Kumar et al., 2006). This was confirmed by our results as acetosyringone treatment was found to lead to a significant enhancement in transformation frequency (nearly 1.3 fold) for all the bacterial strains tested. Similar observations have been documented in Solanum melongena (Jain and Singh, 2015), Cucumis melo (Mohiuddin et al., 2012), S. scutellarioides (Saleh and Thuc, 2009), S. officinalis (Grzegorczyk et al., 2006), wheat (Rashid et al., 2011), rice (Tripathi et al., 2010), and cotton (Afolabi-Balogun et al., 2014).

Carbohydrates, both as sources of energy and carbon and osmotic regulators, play a crucial role in in vitro cultures. In addition, sugars can act as signaling molecules and/or regulators of gene expression regulating plant growth and development (Rolland et al., 2006; Smeekense et al., 2010). Recent work has shown that exogenous sucrose can compensate for the regulators needed for root meristem maintenance (Wahl et al., 2010). Expression of a homeobox gene, which maintains localized auxin maxima in the root apical meristem, is induced by a non-metabolizing sucrose analogue (Gonzali et al., 2005).

Many studies have shown that different sugars significantly affect hairy root growth (Jain and Singh, 2015; Petrova et al., 2015). The results of the present study showed that carbon source had significant effects on root dry and fresh weights, root length, and lateral root density. In general, sucrose was identified as the best carbon source for P. abrotanoides hairy root growth while sorbitol was the weakest. Maximum growth of hairy roots was achieved in a 1/2 MS medium containing 3% sucrose. In agreement with our findings, Sivanesan and Jeong (2009) found that fructose or glucose (3%, w/v) used instead of sucrose in the culture medium led to a significantly slower growth rate of Plumbago zeylanica L. hairy roots.

Sucrose is the most widely used carbon source in plant tissue cultures. This is due to its efficient absorption through the cell membrane (Petrova et al., 2015). This is also confirmed by the findings of Sivakumar et al. (2005) who reported that sucrose effectively enhanced the growth of ginseng hairy roots. The transgenic hairy roots of Arnica montana (Petrova et al., 2015), S. melongena (Jain and Singh, 2015), Solanum trilobatum (Shilpha et al., 2015), Momordica charantia L. (Thiruvengadam et al., 2014), Withania somnifera (L.), and Dunal (Praveen and Morthy, 2012) have also achieved their best growth and highest biomass in an MS medium containing 3% sucrose.

Hairy root cultures aimed at producing secondary metabolites have been employed with different plant species including Ophiiorrhiza pumila (Kitajima et al., 2002; Asano et al., 2013), Panax ginseng (Yu et al., 2005), Ruta graveolens (Kuzovkina et al., 2004), Scutellaria lateriflora (Wilczanska-Barska et al., 2012), and Lithospermum canescens (Syklowska-Baranek et al., 2012). Successful production of tanshinones in hairy root cultures of Salvia miltiorrhiza Bunge (Hu and Alfermann, 1993; Wang and Wu, 2010) and enhanced production of these secondary metabolites by elicitation have also been reported (Ge and Wu, 2005; Yan et al., 2005; Wu and Shi, 2008; Wang and Wu, 2010; Gupta et al., 2011).

Hu and Alfermann (1993) first reported on hairy root culture of S. miltiorrhiza induced by infecting explants from sterile plants with the ATCC15834 and TR105 strains of A. rhizogenes. The authors cultured the hairy roots in a liquid MS medium without ammonium nitrate and detected seven types of tanshinones (containing tanshinone IIA and cryptotanshinone) in the hairy roots thus produced. They also noted a 22-fold increase in the dry weight of the hairy roots in 20 days and a total root tanshinone content of 43 mg g⁻¹ DW (20 mg g⁻¹ DW as cryptotanshinone). In a later study by Chen et al. (2001), hairy roots of S. miltiorrhiza were induced from plantlets using A. rhizogenes ATCC15834 with only trace amounts of cryptotanshinone (0.001% DW) and other tanshinones (tanshinone IIA, tanshinone IIB, and tanshinone I) produced in the MS medium lacking ammonium nitrate. Upon elicitation with yeast elicitor, not only was the intracellular content of cryptotanshinone observed to increase from 0.001% to as much as 0.096% of the dry weight but the growth of hairy roots also improved. Our HPLC analysis results indicated measurable presence of cryptotanshinone and tanshinone IIA in the hairy root clone T7 of P. abrotanoides after less than two months of cultivating in a liquid medium. The amount of cryptotanshinone (53.17±0.26 µg g⁻¹ DW or 0.005% DW) was 5 times higher than that recorded by Chen et al. (2001) for S. miltiorrhiza. However, the plant species, bacterial strains, and culture conditions in our experiment with no elicitors used were not exactly the same as those in Chen et al.’s. Hairy root culture of P. abrotanoides coupled with techniques for increasing secondary metabolite production seems to provide evidence for the useful application of biotechnology to enhance tanshinones production.

**Conclusion:**
Transgenic hairy roots of Perovskia abrotanoides were
successfully induced and established using three different strains of *A. rhizogenes* (ATCC15834, R1000, and TR105). Whole in vitro seedling nodes were found to be explants most favorable to hairy root induction. Also, the half-strength MS medium containing 3% sucrose was identified appropriate for the growth and biomass accumulation of hairy roots. The hairy roots thus produced were found capable of growing in the nutrient medium in the absence of growth regulators. Moreover, the hairy roots of *P. abrotanoides* were found to possess the potential for producing such medicinally important components as tanshinones. The findings of the present study can be used for further investigation of the metabolite profiles and production of pharmaceutically valuable compounds such as tanshinones in the hairy roots of the species.

**Conflict of interest:**
The authors declare that they have no conflict of interests.

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**Reference:**


