Development of a highly efficient virus-free regeneration system of *Salvia miltiorrhiza* from Sichuan using apical meristem as explants

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**Abstract**

**Background:** The dry root and rhizome of *Salvia miltiorrhiza* are used to treat cardiovascular diseases, chronic pain, and thoracic obstruction over 2000 years in Asian countries. For high quality, Sichuan Zhongjiang is regarded as the genuine producing area of *S. miltiorrhiza*. Given its abnormal pollen development, *S. miltiorrhiza* from Sichuan (S.m.-SC) relies on root reproduction and zymad accumulation; part of diseased plants present typical viral disease symptoms and seed quality degeneration. This study aims to detect unknown viruses from mosaic-diseased plants and establish a highly efficient virus-free regeneration system to recover germplasm properties.

**Results:** Tobacco mosaic virus (TMV) and cucumber mosaic virus (CMV) were detected from mosaic-diseased plants. Primary apical meristem with two phyllo podium in 0.15–0.5 mm peeled from diseased plants were achieved 73.33% virus-free rate. The results showed that the medium containing MS, 0.5 mg/L 6-BA, 0.1 mg/L NAA, 0.1 mg/L GA³, 30 g/L sucrose and 7.5 g/L agar can achieve embryonic-tissue (apical meristem, petiole and leaf callus) high efficient organogenesis. For callus induction, the optimal condition was detected on the medium containing MS, 2 mg/L TDZ, 0.1 mg/L NAA by using secondary petiole of virus-free plants under 24 h dark/d condition for 21 d. The optimal system for root induction was the nutrient solution with 1/2 MS supplemented with 1 mg/L NAA. After transplant, the detection of agronomic metric and salvianolic acid B content confirmed the great germplasm properties of S.m.-SC virus-free plants.

**Conclusions:** A highly efficient virus-free regeneration system of S.m.-SC was established based on the detected viruses to recover superior seed quality. The proposed system laid support to control disease spread, recover good germplasm properties in S.m.-SC.

**Keywords:** *Salvia miltiorrhiza*, Virus-free, Regeneration, Callus-organogenesis, Hydroponic

**Introduction**

*Salvia miltiorrhiza* Bunge, also called Danshen, is a perennial herb belonging to the family Lamiaceae and has been used to treat menstrual disorders, cardiovascular diseases, chronic pain, and thoracic obstruction over 2000 years in Asia countries [1–4]. At present, the annual consumption of medicinal materials exceeds 80,000 tons.

Due to the huge market demand, the natural population of *S. miltiorrhiza* has been drastically harvested and severely damaged. *S. miltiorrhiza* has been gradually domesticated and induced to various places in China, such as Sichuan, Shandong, Hebei, etc. [1, 5]. In particular, *S. miltiorrhiza* produced from Zhongjiang of Sichuan (S.m.-SC) is considered the best-quality product for stick, bright red, and high content of salvianolic acids in the...
roots; it has been exported overseas, and Zhongjiang is deemed as genuine producing area [5, 6]. However, the pollen of S.m.-SC develops abnormally and relies on root reproduction [7]. With disease accumulation in cultivated populations of S.m.-SC, the quality and quantity of diseased plants have declined seriously [8]. Diseased plants of S.m.-SC present whitening, mottled and yellow leaves, and premature withering, and some plants present typical symptoms of viral disease. Ding et al. reported that cucumber mosaic virus (CMV) infected S.m.-HB (S. miltiorrhiza produced from Hebei province, China) with symptoms of mosaic, stunting, chlorosis, and mottle [9]. No research has reported on virus in S.m.-SC. The unknown virus of diseased S.m.-SC must be detected for disease prevention and control.

Virus elimination in vitro plant culture is the most effective approach to obtain virus-free germplasm resources of superior variety to be applied to many crops, such as Solanum tuberosum [10, 11]. Virus-free regenerated plants grow fast and have strong disease resistance ability and significantly improved production and quality [12]. Few regeneration systems have been established through callus genesis for S. miltiorrhiza [13–15]. All of these systems use the seeds or vegetable organ (leaf, root, etc.) as the starting materials but cannot remove zymad from the plant, explants, and callus, showing browning or vitrification; moreover, the salvianolic acid B content in the regeneration plant is low. None of the systems are suitable for in vitro preservation, purification, and rejuvenation of S.m.-SC plants. A highly efficient virus-free regeneration system for S.m.-SC remains to be developed.

Overall, the current study was aim to detect the virus of diseased plant and development a high efficient virus-free regeneration system for S.m.-SC to recover good germplasm properties and improve its yield and quality.

Results

Virus detection and symptom observation of diseased plants

Table 1 and Fig. 2B indicate that the media components significantly influenced the survival of primary meristem. Exogenous cytokinin (6-BA) has a crucial effect on the growth of apical meristem. When 6-BA was used in the media, the survival rate significantly increased compared with that in the control, resulting in the lowest survival rate of 31.82% in B0 media to...
the highest survival rate of 75.47% in B3 media (supplied with 0.5 mg/L 6-BA), while the bud was emaciated and vitrified. The concentrations of sugar and agar were adjusted in B5–B9 medium at the same exogenous cytokinin level. The result shows that the survival rate of primary meristem significantly decreased with increasing agar concentration in B5, B7, and B8 medium. The buds were emaciated or vitrified (Fig. 2B: d, f, g). At the same time, increasing the concentration of sucrose was beneficial to the survival of primary meristem by
comparing the 30 g/L group (B3, B6 and B9 medium) and the 20 g/L group (B5, B7, and B8 medium).

The highest survival rate was 77.36% in B6 media (Table 2), and vigorous directly regenerated shoot was obtained after culture (Fig. 2B: e). The virus carrying states of 45 directly regenerated shoots from 15 primary apical meristems trimmed from five virus-infected plants were tested, and the virus-free rate was 73.33% (Figs.2).

Therefore, the B6 media containing MS + 0.1 mg/L NAA + 0.1 mg/L GA3 + 0.5 mg/L 6-BA + 30 g/L sucrose + 7.5 g/L agar was selected for the initial culture of primary apical meristem.

### Callus induction from directly regenerated tissue

Secondary apical meristem, leaf and petiole were took up from virus-free directly regenerated buds obtained by detoxification culture to induce callus. The results show that the yellow-greenish callus was induced from the secondary apical meristem (SM) without phyllo podium (Fig. 3A). Most of secondary leaf (SL) and secondary petiole (SP) manifest browning and have poorly induced callus under the PM growth condition (Fig. 3B, C).

The inducing effect of the six media for three cytokines on callus formation under two light conditions (16 h light/8 h dark and 24 h dark/day) was investigated using secondary leaf and petiole to optimize the induction condition for callus formation. L0 (without exogenous cytokines) was used as control.

Significant difference (P < 0.05) in the main effect of light and exogenous hormone kinds was observed in the formation of secondary callus (Figs. 3 and 4). Overall, secondary leaf has higher callus-genesis potential, with the induction rate of 73.32%. Light significantly represses the formation of secondary callus. When the light condition changed, the callus induction rate (33.33%–45.90%) of secondary leaf and petiole under 16 h light/8 h dark

| Group | Medium components | Survival Rate (%) | Growing status |
|-------|-------------------|-------------------|----------------|
| B0    | 0  0  0  8.5  30  | 34.09±4.69 d      | –              |
| B1    | 0.1 0.1 0  8.5  30 | 40.48±4.41 d      | –              |
| B2    | 0.1 0.1 0.25 8.5  30| 67.35±4.24 b      | Vitrified bud  |
| B3    | 0.1 0.1 0.50 8.5  30| 75.47±2.53 a      | Partial health buds |
| B4    | 0.1 0.1 1.00 8.5  30| 58.00±2.81 c      | Vitrified buds, callus formed |
| B5    | 0.1 0.1 0.50 7.5  20| 75.47±2.89 a      | Emaciated, vitrified buds |
| B6    | 0.1 0.1 0.50 7.5  30| 77.36±2.61 a      | Vigorous bud    |
| B7    | 0.1 0.1 0.50 8.5  20| 62.50±3.37 b      | Emaciated bud   |
| B8    | 0.1 0.1 0.50 9.5  20| 53.06±5.65 c      | Emaciated bud   |
| B9    | 0.1 0.1 0.50 9.5  30| 69.44±4.34 b      | Vitrified bud   |

Table 2  Effect of medium components in combination with MS medium on the survival of apical meristem

Number of apical meristem inoculated per group was twelve; three replication of each group were laid. Data were collected after 4 weeks. Different letters indicate significant differences among mediums for each group as determined by Duncan’s multiple range test followed by one-way ANOVA at the p < 0.05 significant level.

![Fig. 3](image-url) Callus growth of different secondary materials (cutted from directly regenerated buds) on B6 media under the light condition with 16 h light/8 h dark. A Callus from secondary apical meristem, B culture effect of secondary leaf in B6, and C culture effect of secondary petiole in B6. Three replication of each group were laid. Data were collected after 3 weeks.
condition significantly increased to 57.4%–73.32% under 24 h dark/d. The induction varied rate of leaf callus was 40% compared 16 h light/8 h dark with 24 h dark, and that of petiole was 12% only, indicating that leaf was more sensitive to light condition than petiole (Fig. 3A). Under the light condition of 16 h light/8 h dark, most explants appeared as drying and browning after 30 days, and poorly grown brown callus was also induced. The browning callus from petiole and leaf developed hyperhydric shoots, or growth was stopped after 40 days of induction. The leaf and petiole product, namely, yellow–greenish callus was obtained from the incision under 24 h dark/day, and the growth was normal, and browning was not observed (Fig. 3B).

The influence of different kinds and concentrations of cytokines for callus induction from petiole and leaf explants was measured under the light condition of 24 h dark/d (Fig. 4A, B). With increasing hormone concentration, the callus induction rate significantly increased. On media L3 and L4 (supplied with 1 or 2 mg/L KT), explants were yellowish-brown, scarce callus formation (2.78%–8.70%) was observed in the leaf or browning callus was formed at the petiole incision. By contrast, on the media supplemented with 6-BA or TDZ, 52.94%–84.21% of the explants produced callus. The highest induction rate (84.21%) was obtained on the media supplemented with 2 mg/L TDZ for secondary petiole callus induction, and the callus was compact and greenish-yellow.

The optimized light and medium condition for secondary callus induction is that cutting secondary petiole into 10 mm length and inducing in the media containing MS + 2 mg/L TDZ + 0.1 mg/L NAA + 30 g/L sucrose + 7.5 g/L agar. The culture was kept for 3 weeks under 24 h dark/d light condition.

**Induction and growth of adventitious buds**

Callus reached the differentiation phase of organ after almost one month of vegetative growth phase. Callus from secondary apical meristem, secondary leaf, and secondary petiole fragment was inoculated in screened media (Table 2, B6) to induce bud organogenesis.

The formation of adventitious buds was an event observed in the callus of different origins, did not all occur healthy buds after 40 days of culture (Fig. 5). The regeneration ability of secondary apical meristem was examined. Hyperhydric buds were produced from the
pale yellow-green callus (Fig. 5A: SM). The callus from apical meristem had better plant regeneration ability, as verified in terms of regeneration rate and growth coefficient. The callus of secondary apical meristem with an average of 15.33 buds/explant and 78.86% regeneration rate, has a greater regeneration potential for buds organogenesis, compared with secondary leaves and petiole (Fig. 5B, C). This finding indicated that tender tissue was sensitive to medium component and negative to healthy bud proliferation. Regarding the development of secondary leaf callus, adventitious buds were produced from spherical embryoid growing from the surface bulge of massy callus (Fig. 5A: SL). However, vigorous adventitious buds were produced on the callus of petiole incision. The structure had a deeply yellow-green dense callus, and the inner is lignified (Fig. 5A: SP). The numbers of adventitious buds per callus of secondary leaf and secondary petiole were 8.33 and 8.67, and the regeneration rates were 58.61% and 66.03%, respectively (Fig. 5B, C).

In short, apical meristem is efficient in micro-propagation because of its high regeneration frequency. Embryonic callus has compact structure in deep yellow-greenish color in in vitro culture for S.m.-SC. Thus, the convenience and regeneration ability of explant were comprehensively considered. Secondary petiole callus was selected as the material for high-efficient adventitious bud induction.

**Root induction and plant acclimatization**

Root regeneration is a crucial step for successful regeneration of plants. The quality of the adventitious roots directly affects subsequent acclimatization and transplant survival rate. When the regeneration buds were dissected into single bud and grew to 3–5 cm high in the media, adventitious roots were induced in the solid medium and improved hydroponics. Two styles for nine combinations were designed for root regeneration of S.m.-SC. The plant growth effect is presented in Table 3 and Fig. 6.

The rooting effect was compared among M0–M3 groups. The result showed that the regenerated plants grew slowly, malnutrition occurred, and the lowest rooting rate (38.46%) was found in the control group. Under the same auxin concentration (1 mg/L), applying a single auxin in the medium had better root induction effect than mixed use. The M3 group supplied with 1 mg/L NAA had greater rooting rate (80.00%) than M1 and M2. In the H0 group without added nutrients and auxin, the plants grew slowly and the rooting rate was lower. After adding the basal medium, the plants grew vigorously, and the rooting rate was significantly increased to more than 70%. The H4 group had the best growth effect, and the rooting rate reached 87.88%. Based on comparison of the solid medium and improved hydroponics, the roots induced by hydroponics were thinner and longer (Table 3).

After hard-seeding in nutrient soil for 21 days, the virus-free regenerated plants were transplanted into soil. The growth effect and survival rate after the transplant of the regenerated plants were compared between M3 and H4 groups. The leaves of the M3 plants were easy to be dehydrated and wilt after transplanting (Fig. 6A, B), and the survival rate was only 65.29% after 30 days. However, the leaves in the H4 group plants had healthy growth, the leaves were stretched and emerald green, the root system was slender (Fig. 6C, D), and the survival rate reached
88.64% after transplanting. The leaves of whole regenerated plants were collected after rooting, and no positive reaction for TMV or CMV was detected in the Dasselisa test.

The result showed that inducing root formation in the improved hydroponics is a suitable option to replace root induction in sterile solid media. Nutrient solution with 1/2 MS supplemented with 1 mg/L NAA and free-sugar in hydroponics can facilitate efficient plant regeneration and survival of S.m.-SC.

Evaluation of yield metric and active ingredient

The whole virus-free plants were transferred to soil after acclimatization. Common plants were planted under the same condition as control. In total, 10 virus-free and common plants were collected after 10 months of culture for yield metric and main active ingredient analysis of roots.

The yield metric and active ingredient of the root of the virus-free plant significantly improved compared with those of the common plants (P < 0.05, Table 4). The average root fresh weight of virus-free plants was 398.50 ± 87.88 g/plant, and the yield was increased by 62.96%. The average main root length

| Plants | Root Weight (g/plant) | Main root Number | Main root Length (mm) | Main root Diameter (mm) | Salvianolic acid B (mg/g) |
|--------|----------------------|------------------|-----------------------|-------------------------|-------------------------|
| Virus-free | 398.50 ± 87.88 a | 6.60 ± 1.82 a | 250.00 ± 36.51 a | 14.06 ± 1.84 a | 87.87 ± 14.74 a |
| Common | 244.54 ± 133.95 b | 6.46 ± 1.96 a | 200.81 ± 58.85 b | 11.20 ± 2.73 b | 67.29 ± 16.66 b |

* Means with different letters are significantly different at P < 0.05 according to Student's t-test. Three replication of each plants were laid. Yield data were obtained by measuring fresh roots after harvest, and salvianolic acid B content was measured after drying.
increase (P < 0.05) by 30.58% in the dried roots in the virus-free plants and that of the common plants had 67.29 ± 14.74 mg/g salvianolic acid B, and that of the common plants had 67.29 ± 16.66 mg/g. The values represented a significant increase (P < 0.05) by 30.58% in the dried roots in the virus-free plants compared with those in the common plants.

Salvianolic acid B is one of the main biologically active secondary metabolites in the dry roots and rhizomes of S. miltiorrhiza [16]. The HPLC analysis showed that the dry root of the virus-free plants had 87.87 ± 14.74 mg/g salvianolic acid B, and that of the common plants had 67.29 ± 16.66 mg/g. The values represented a significant increase (P < 0.05) by 30.58% in the dried roots in the virus-free plants compared with those in the common plants.

Virus-free plants grow robustly, the main roots are thick and long, the yield is high, and salvianolic acid B content is significantly increased (P < 0.05).

Discussion

**Virus-free and regeneration of apical meristem**

The selection of suitable explant is the base of success for in vitro culture. Plant meristem has remarkable regenerative and antiviral capacity, enabling them to either recover damage tissues or established de novo organ [17, 18]. Wu et al. reported that the stem cell regulator WUSCHEL was expressed in response to viral infection and inhibited its accumulation to protect the central and peripheral areas in Arabidopsis meristem [19]. However, apical meristem isolation is very time consuming, and injury or bruising during lengthy isolation can usually kill the meristem [20]. Cytokinin enhances the mitotic activity of the stem cell to reduce the damage by disinfection and splicing of explants. The survival rate of primary apical meristem of S.m.-SC was increased by improving the concentration of cytokinin. Isah et al. demonstrated that cytokinin promoted the growth of Gymnema sylvestre meristem, which turned green earlier, and increased the survival rate significantly [21]. During in vitro culture, endogenous and exogenous plant hormones can trigger cellular reprogramming, and osmotic pressure and hardness significantly affect explant survival and regeneration [22]. In the present study, the apical meristem on high-hardness medium presented with vitrification symptom, and the survival rate was improved by increasing the sugar concentration. Hence, in the construction of in vitro culture environment, the key factors of growth hormones should be given attention, and the adjustment of other factors, such as osmotic pressure and hardness, should not be ignored.

**Effect of PGR and light during tissue culture**

The ratio of auxin and cytokinin directs the differentiation progress of explants in in vitro culture [23]. A high concentration of cytokinin promotes callus formation. The formation of embryogenic tissue is the key to regeneration. Wounding stress induces the local accumulation of cytokinin at the wound site and modifies hormone biosynthesis, thereby activating the transcription and translation processes and leading to changes in cell cycle phase and callus formation [22]. In early stage of red maple in vitro culture, callus induction and embryo were more dependent on PGR, especially TDZ [24]. In this article, TDZ showed better embryonic callus induction ability of S.m.-SC compared with 6-BA and KT, consistent with the result reported by Chen [13]. Although hormone effects are stage specific, the medium containing TDZ only was not effective for bud regeneration or induced vitrification and emaciation of apple buds; high shoot regeneration was achieved in the media supplied with 6-BA only [25]. Here, vigorous buds were induced from deeply yellow–green dense callus in B5 media supplied with 0.5 mg/L 6-BA, 0.1 mg/L NAA, and 0.1 mg/L GA3.

Light is one of the key factors that affect plant regeneration and growth. Callus genesis and shoot formation are suppressed by light in vitro culture of apple [25, 26]. Here, browning explants and poor callus genesis were observed during 16 h light/8 h dark condition after 30 days of culture of S.m.-SC, which may be caused by the oxidation of abundant photosensitive substances (phenolic acids, flavonoids, and tanshinones) in the explants [27, 28]. Change in light condition is essential during redifferentiation and bud growth stage, and dark pretreatment of the explant for 3 weeks under 16 h light/8 h dark condition enhanced the formation of apple shoots [25]. After dark culture for a month, pale shoots with elongated stem and lack of chlorophyll in the apex were detected after in vitro regeneration of eggplant [29]. The seedlings of S.m.-SC show slender stems, yellow–green leaves, and vitrification under long-term dark conditions.

**Rooting and acclimatization of regenerated buds**

In regenerated plants under in vitro condition, the surface of leaf lacks cuticle, the stoma is highly opened, and transpiration is extremely strong. Changes in environment humidity and organ damage (such as leaf or root breakage) during transplantation can easily cause water loss and withering and even death [24, 30, 31]. Similarly, the regenerated plants in solid medium experienced severe water loss and wilting after transplantation, resulting in a low survival rate of only 65.29%. Meanwhile, the regenerated plants in hydroponics grew vigorously, and the survival rate reached 88.64%. The great root induction effect may be due to the fact that humid environment and good air circulation can make the regenerated plants to better adopt to the environment and growth faster [31]. Kyle et al. developed an improved axenic
hydroponic propagation system to produce large quantities of the roots of *Taraxacum kokssaghyz*; the plants grew faster in the system than in the solid medium [32].

**Change in the content of the main active ingredient of plants after virus-free**

Salvianolic acids are the main active secondary metabolites in the dry roots and rhizomes of Danshen. Phenolics participate in plant stress resistance as constitutive expression products of genes, while ketones are considered secondary metabolites of inducible expression. For example, higher constitutive levels of total phenols were found in resistant synthetic hexaploid wheat compared with those in moderately resistant and susceptible genotypes [33]. During the development process, salvianolic acids exist in both the above-ground and underground parts of *S. miltiorrhiza* [34, 35], indicating that phenolic acids may be synthesized in both parts. After detoxification, the virus and other diseases in the plant are reduced, the growth is vigorous, the leaves are dark green, and the photosynthesis is enhanced, thereby promoting the synthesis of salvianolic acids and significantly increasing (P < 0.05) the yield of *S.m.-SC* compared with those in the common plants (Additional file 1).

**Conclusion**

CMV and TMV were first determined from *S. miltiorrhiza* from Sichuan diseased plants. An efficient virus-free regeneration system for *S.m.-SC* was established using apical meristem as explants for the first time (Fig. 7). In summary, the optimal bud induction media contained MS, 0.5 mg/L 6-BA, 0.1 mg/L NAA, 0.1 mg/L GA3, 30 g/L sucrose, and 7.5 g/L agar. For callus induction, the optimal condition was L6 media containing MS, 2 mg/L TDZ, 0.1 mg/L NAA, 30 g/L sucrose and 7.5 g/L agar and using 10 mm secondary petiole of virus-free plants under 16 h light/d condition for 21 days. For rooting induction, the optimal system was using nutrient solution with 1/2 MS supplemented with 1 mg/L NAA. After transplant, virus-free plants grew robustly, the main roots were thick and long, the root yield and salvianolic acid B content in the dry root were significantly increased to 398.50 g/plant and 87.87 mg/g by 62.96% and 30.58% compared with those in common plants.

**Materials and method**

**Plant material**

The diseased plants of *S. miltiorrhiza* were collected from a planting base in Zhongjiang, Sichuan, China. The whole living plants were kept in the Teaching and Research Base of Sichuan Agriculture University (Ya’an, Sichuan, China). The buds sheared from the diseased plants were selected as the experiment material.

**Virus detection of plants**

Leaves collected from 17 *S. miltiorrhiza* diseased plants (marked as D1–D17, Table 1) and regenerated plants were serologically subjected to detection of viruses in Double antibody sandwich elisa (Das-ELISA), Tobacco mosaic virus (TMV), Cucumber mosaic virus (CMV), Tomato mosaic virus (To MV), and Tomato spotted wilt virus (TSWV) were detected. For ELISA tests, the reagents, buffers, and controls supplied by AGDIA (American) were used in accordance to manufacturer’s instructions. The detection of the regenerated buds is the same as that of the diseased mother plant.

**Explant sterilization and screening of bud induction medium**

Parts of the plants were selected to screen the bud induction medium of *S. miltiorrhiza* by using apical meristem after virus detection of the diseased plant. Buds with no fully spread leaves sheared from the diseased mother plants were surface cleaned by detergent for 5 min and rinsed for 30 min under running water before sterilization at gnotobasis. The explants were immersed in 75% alcohol for 45 s and washed twice with sterilized distilled water. The explants were cleaned by soaking into 0.1% mercury chloride solution for 7 min and 30 s and washed six times with sterilized distilled water under aseptic condition.

Primary apical meristem (PM, Fig. 2A) was exfoliated from the sterilized plant material under digital microscope (yookdd DM3) and transferred into the bud induction media (BIM, B0–B9) containing 1 × Murashige and Skoog (MS) base medium (Chembase, China) with 0.1 mg/L 1-naphthaleneacetic (NAA) (Chembase, China) and 0.1 mg/L Gibberellin A3 (GA3) (Chembase, China) and supplied with different concentrations of 6-benzylaminopurine (6-BA, 0, 0.25, 0.50, and 1.00 mg/L) (Chembase, China), agar (7.5, 8.5 and 9.5 g/L), and sucrose (20 and 30 g/L); B0 containing 1 × MS basal salt without any phytohormone was used as control (Table 2). One apical meristem was inoculated in a glass culture bottle (65 mm × 115 mm with breathable plastic cover). About 50 mL of the medium was placed per bottle for meristem culture. Twelve meristems were inoculated in each medium, and the experiment was repeated three times. The cultivating bottles were placed in a thermostatic chamber at 22 ± 1 °C with a photoperiod of 16 h light/8 h dark under white fluorescent light (average illumination intensity 75 μmol m−2 s−1). The survival rate of primary apical meristem was measured after 4 weeks.

**Callus induction from directly regenerated tissue**

The secondary meristem, leaf, and petiole from the sterile, directly regenerated plants were used for callus
induction. The leaf and petiole were cut into fragments of 10 mm and placed in callus induction media (CIM) containing 1 × MS base salt with 8.5 g/L agar, 30 g/L sucrose, and 0.1 mg/L NAA and supplied with different concentrations of 6-benzylaminapurine (6-BA, 1, 2 mg/L), kinetin (KT, 1, 2 mg/L), and thidiazuron (TDZ, 1, 2 mg/L), named as L1–L6 groups, respectively; L0 without cytokinin was used as control. About 18 leaf and petiole fragments were inoculated in each plastic cultivating pot (80 mm × 96 mm with breathable plastic cover) containing 50 mL of CIM. The cultivating bottle was placed in the same thermostatic chamber under two photo-periods of 16 h light/8 h dark and 24 h dark/d. Five pots were used for each treatment, and all of treatments was
repeated three times. The induction rate of callus and the growth of explants were detected after 3 weeks.

**Induction and growth of adventitious buds**
The callus from the secondary meristem, leaf, and petiole was inoculated in the screened meristem medium after 1 month of callus induction. Eight calluses (10 mm in diameter) from each material per container were placed in each culture pots for induction of adventitious buds. Five pots were used for each callus for induction of adventitious buds, and all of materials was repeated three times. The adventitious buds were divided into single bud to strengthen the shoots in the same medium after 30 days of induction. The inducting ratio of the adventitious buds was also determined.

**Root induction and plant acclimatization**
After the plants were growth into 3 cm high, the rooting and plant acclimatization effect were evaluated through sterile solid media and improved hydroponics. The rooting media contained 1/2 MS base salt with 8.5 g/L agar and 10 g/L sucrose supplied with different kinds and concentrations of auxin (H0–H4, Table 3). Plastic seeding boxes (365 mm × 230 mm × 110 mm, including lid and seeding plate and base plate) were used in hydroponics. A 1:1 mixture of perlite and vermiculite was used as the fixed material in the improved hydroponics. The level of culture liquid was maintained in 3 cm high, and the culture liquid contained 1/2 MS base salt with 8.5 g/L agar and 10 g/L sucrose supplied with different kinds and concentrations auxin (H1–H4, Table 3) but without agar and sucrose. H0 was supplied with distilled water and used as control. Num-

**Evaluation of yield metric and main active ingredient**
The virus-free plants were transplanted into soil in Zhongjiang, Sichuan, China after hardening–seeding in March, 2020 at the germination time of the cultivated plants. A land that has a flat terrain, smooth drainage, sunny terrain, no soil pollution, and never planted with S. m-SC was selected. The plants were alternately planted in ridges and watered daily with tap water for 1 month after transplantation.

The root and rhizome tissues were harvested separately from the virus-free plants and common plants after 10 months of culture for yield metric and main active evaluation. Root weight, main root number, main root length, and main root diameter per plant were measured. The main active ingredient, salvianolic acid B, was assayed after drying by referring to Chinese pharmacopoeia (2020 edition). Main active ingredients were determined by HPLC (1260 Infinity II, Agilent Technologies, American).

**Data statistics and analysis**
Data were analyzed using SPSS version 23.0 statistics and Duncan’s test. One-way ANOVA was used for multiple comparisons.

**Supplementary Information**
The online version contains supplementary material available at https://doi.org/10.1186/s13007-022-00872-4.

**Additional file 1.** Figs. 1 Virus detection results of diseased plants (D1–D17) of S.m.-SC by Das-ELISA test. Figs. 2 Virus detection results of regenerated plants of S.m.-SC by Das-ELISA test. Figs. 3 Caslus induction rate and growth status of SP and SL under two kinds light condition of 16 h light/ 8 h dark and 24 h dark/d.

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**Authors’ contributions**
LZ conceived this study, SCY, JF and QYL performed the experiment, SN, LW and LXY provide advice, SCY analyzed the data and wrote the manuscript, YYJ, RWY and LZ revised the manuscript. All authors read and approved the final manuscript.

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**Availability of data and materials**
All data generated or analysis during this study is included in published article. The materials are available from the corresponding author on reasonable request.

**Declarations**

**Ethics approval and consent to participate**
Not applicable.

**Consent for publication**
Not applicable.

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

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