Microclonal propagation of *Dasiphora fruticosa* (Rosaceae)

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**ABSTRACT**

Microclonal propagation of *Dasiphora fruticosa* Duham. (wild form (Df0) and cultivars ‘Lovely Pink’ (Df1), ‘Bella Bianca’ (Df2), ‘Bella Sol’ (Df3)) was investigated in details. Microplants were cultured on hormone-free solid ½ Murashige and Skoog medium. Well-formed plants with developed root system were established after 2 months of culturing. The maximum values of shoot height, number of internodes, and root mass were observed in the Df1 microplants, with a greater number of roots in the Df0 microplants. The use of half-strength medium for both growth initiation and microplant propagation allows saving reagents and reducing the cost of *D. fruticosa* microclonal multiplication *in vitro*. The use of ready-mixed soil with pH 5.6–6.7, and cultural vessels for growth of microplants under controlled conditions such as temperature, light and humidity made it possible to successfully adapt more than 90% of test-tube microplants to the soil conditions.

**Keywords:** *Dasiphora fruticosa*, *in vitro*, medicinal plant, micropropagation, cultivation, adaptation to soil

**REЗЮМЕ**

Гафицкая И.В., Орловская И.Ю., Наконечная О.В., С.В. Нестерова. Микроклональное размножение лапчатника кустарникового (*Dasiphora fruticosa*, Rosaceae). Подобраны условия для микроклонального размножения лапчатника кустарникового (*Dasiphora fruticosa* (L.) Rydb.) – лекарственного и декоративного растения. В работе использованы *D. fruticosa* из природной популяции – Df0 и 3 декоративных сорта: «Lovely Pink» – Df1, «Bella Bianca» – Df2, «Bella Sol» – Df3. При выращивании микрорастений на безгормональной питательной среде Мурасиге и Скуга с половинной концентрацией (*) минеральных удобрений через 2 месяца были получены хорошо сформированные растения с развитой корневой системой. Максимальные значения высоты побегов, числа междоузлий и массы корней выявлены у растений Df1, большее число корней отмечено на растениях Df0. Единый состав питательной среды для инициации и пролиферации позволяет экономить реактивы и делает более дешевым микроклональное размножение *D. fruticosa* и сортов. В контролируемых лабораторных условиях (температура и влажность воздуха, освещенность) более 90% пробирочных растений успешно адаптируются к условиям почвогрунта в культуральных сосудах объёмом 0,5 л при использовании готовой почвенной смеси (pH 5.6–6.7).

**Ключевые слова:** *Dasiphora fruticosa*, *in vitro*, лекарственное растение, микроклонирование, сорт, культивирование, адаптация к почвогрунту

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antimicrobial, immunomodulatory, and antioxidant properties (Tomczyk et al. 2008; Tomczyk & Latte 2009, Zeng et al. 2019). D. fruticosa decoctions and infusions are used for the treatment of gastrointestinal diseases in Tibetan, Indian, and Mongolian medicine (Stalnaya 2015, Zeng et al. 2019). Decoctions also normalize blood pressure, strengthen the walls of blood vessels, prevent their rupture and, thereby, heart attacks and strokes (Stalnaya 2015); the extract is used in the treatment of disbiosis in infants, as a tonic and sedative, as well as a diuretic (Khramova et al. 2003, Khramova, 2009). Dried extract can be applied in manufacture of confectionery, chewing gum, and others (Stalnaya 2015).

Dasiphora fruticosa is a melliferous and pergans plant. Mature plants inhibit soil erosion (Korzn & Valvilo 2015). Kuril tea is an ornamental shrub and, therefore, is used in decoration of alpine hills, rocky gardens, hedges in city parks (Fig. 1A).

Studies on vegetative propagation of the Kuril tea were conducted previously, including propagation for industrial production (Korzn & Valvilo 2015, Urusov & Lobanova 2018). Some species of Rosaceae family such as Potentilla alba and intergeneric hybrids (Fragaria vesca × Potentilla fruticosa, Fragaria × Potentilla palustris) were propagated microclonally in vitro (Silva & Jones 1996, Sutan et al. 2010, Tikhomirova et al. 2016 et etc.). Meanwhile, microclonal propagation method for valuable specimens of ursine plants offers great opportunities for mass plant propagation, due to a high breeding rate (up to 10–10 specimens per year per 1 plant), possibility of all year round cultivation, little space for sterile microplant cultivation, elimination of plants from viruses, and long-term storage of microplants in tubes at low temperatures, allowing to establish a bank of genotypes of valuable species and forms (Kataeva & Butenko 1983).

The purpose of the study was the introduction into culture in vitro and propagation of valuable medicinal and ornamental plant D. fruticosa and its cultivars. It was necessary to find out the conditions for successful growth and adaptation of the obtained plantlets to a soil. The study was of scientific interest and high practical value, as it represented the first stage of the shrubby Kuril tea plantations establishment in Primorsky Region. In prospect, the obtained plants could be used as a source of biologically active metabolites in pharmacetics.

**MATERIAL AND METHODS**

The work on micropropagation of D. fruticosa plants in vitro was performed in 2015–2018 (Vladivostok). The plant sample of D. fruticosa (Df0) was found in collection in the Botanical Garden-Institute FEB RAS, Vladivostok. Samples of cultivars ‘Lovely Pink’ (Df1), ’Bella Bianca’ (Df2), ’Bella Sol’ (Df3) were provided from the private collection (Ussuriysk).

The shoot cuttings of 0.5 to 1.0 cm long with 1–2 accessory buds were used as explants. Shoot cuttings were sterilized according to a standard method (Kataeva & Butenko 1983), using 0.1 % commercial Diocide solution for 2–4 min, and placed upright in 15-mm diameter test tubes on a half-strength MS medium (Murasige and Scoog) with half-concentration of micro- and macroelements. The medium was autoclaved for 20 minutes at 0.8 atmospheres. The tubes with explants were further placed under luminous light (4000 lux) at 24°C, 60–70 % relative humidity, and 16h/8h photoperiod.

Plantlets were cut, and shoot samples with one leaf were transplanted to fresh nutrient medium after 2 months. The main qualitative and quantitative characteristics of plant development during propagation in vitro are shoot length and number of internodes, and micropropagation coefficient. Characteristics of root system are important for transplanting to the ground. Morphometric parameters (are shoot length, number and length of roots) were measured weekly. We determined micropropagation coefficient as the mean number of microcuttings obtained from one regenerant plant for 1 passage (Shumikhin 2005).

We used plastic cultural containers with 0.5 l volume, and soil mixture containing nitrogen (0.2 %), phosphate (0.1 %), potassium (0.1 %), water (no more than 60 %), with pH 5.6–6.7. The soil mixture was manufactured by ECO-AGTI Company (Korolev, Russia). We cultivated plants transmitted to soil in controlled conditions of growth chamber, described above. The data were processed using Microsoft Office Excel and STATISTICA version 13.3. The table show the arithmetic mean value of growth parameters with standard error (n = 30).

**RESULTS AND DISCUSSION**

**Explant sterilization.** We used 0.1 % commercial Dio- cid solution for 4 min to provide qualitative surface sterilization of explants. As a result, more than 90% of viable explants (Fig. 1B) were obtained from each plant specimen, indicating that sterilizing agent and sterilization time for the Kuril tea were selected optimally.

**Microclonal propagation.** We obtained well developed microclones after 2 months of explants cultivation (Fig. 1C) on half-strength MS medium with half concentrations of macro- and microelements. The explants developed into microplants with 9–14 internodes (Table). This method allowed us to obtain the necessary number of microplants (about 1000 microplants) for a short period (4 months). The use of the medium with half salt concentrations allowed us to avoid the preparation of several different cultural media for initiation and proliferation, and helped to make the propagation process easier with less material and labor spendings.

**Plant development during microcloning.** The microplants obtained in the experiment had different morphometric characteristics (Table). Df0 explants grew actively during the third and fifth week of culturing (Fig. 2). After two months they had the smallest stem height and number of internodes, comparing with other experimental groups (Table 1). During the first week of cultivation we observed root formation in 15 % of Df0 explants. After three weeks of growth 65 % of explants had two or more roots. At the end of the experiment (2 month of culturing) we detected the maximum mean root number per Df0 plant comparing with other groups (Table 1), which probably determined high growth rates in Df0 clones during first five weeks of growth, comparing with three other experimental groups.
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We registered active growth of Df1 explants from fifth to eighth week of culturing (Fig. 2). Twenty percent of clones formed roots during the first week; and 60 % of clones formed abundant root system five weeks later. At the end of the experiment shoot length, number of internodes and root mass of the Df1 clones were the highest comparing with other groups (Fig. 2, Table). Maximal internodes number determined the maximal propagation coefficient (12), meaning greater number of explants for further micropropagation.

Active growth of Df2 explants was observed after six weeks of cultivation (Fig. 2). Thirty-five percent of explants formed roots two weeks after start of the experiment. We registered roots on all microplants at the end of culturing; root system was formed with the main root and roots of the second and third order.

Similar to Df2 microplants, Df3 microplants displayed active growth from sixth to eighth week of cultivation (Fig. 2). The explants formed no roots during the first 2 weeks of growth. After three weeks of culturing, 55 % of explants had roots. At the end of the experiment, 95 % of microplants formed, in average, 2 roots per plant. The root mass was the least, comparing with other groups of microplants, because the roots were thin (Table 1). The greatest propagation coefficient in *vitro* (12) was achieved for Df1 cultivar (‘Lovely Pink’). The lowest coefficient, 8, was provided for Df0 sample. The coefficient for Df2 (‘Bella Bianca’) and Df3 (‘Bella Sol’) was equal to 11.

The data on the cultivation features of the plant species from the Rosaceae family can be found in the literature. Thus, Ren and colleagues (2003) reported that MS culture medium supplemented with 2 mg/L 6-BAP (6-benzylaminopurine) and 0.2 mg/L NAA (α-naphthaleneacetic acid) suited the best for cultivation of tender stems of *Potentilla glabra* G. Lodd. The rooting of the established stems was the best on ½ MS+NAA 0.1 mg/L cultivation medium (Ren 2003). He and colleagues (2006) studied micropropagation of *P. potaninii* Th.Wolf and used hypocotyl and cotyledon explants. They achieved the highest frequency of adventitious shoot regeneration using MS medium supplemented with 5.0 mg/L benzyladenine (BA) and 1.0 mg/L NAA). The reproduction effect of the *P. fruticosa* ‘Gold Drop’ explants was the best when the concentration of 6-BAP was 2.0 mg/L. The medium MS+2.0 mg/L 6-BAP +0.6 mg/L NAA +0.2 mg/L K (kinetin) was defined as the optimum for the *P. fruticosa* ‘Gold Drop’ explants proliferation. And the optimal rooting medium for the same species was MS+0.1 mg/L NAA + 1.0 mg/L KT (Ma et al. 2009). The optimum concentration of phytohormones for the *P. alba* L. *in vitro* micropropagation was 1.0 μM 6-BAP + 0.5 μM IBA (indolebutyric acid)+0.05 μM GA (gibberellic acid) (Tikho-
mireva et al. 2016). Production of *P. fulgens* Wall. ex Hook. plantlets with better rate of shoot multiplication and elongation obtained on MS medium supplemented with 1 mg/L Kalone or combined with 1 mg/L NAA (Sambyal et al. 2006).

It is interesting to note that the *in vitro* propagation coefficient for *P. alba* (average fold multiplication) was 9.3 (Tikhomirova et al. 2016) and 11.6 for *P. fulgens* (Sambyal et al. 2006). *P. alba* and *P. fulgens* are herbaceous plants. Our study with *D. fruticosa* also revealed high propagation coefficient for the representative of the Rosaceae family with different growth form – a shrub. Herbaceous plants are often characterized with a high propagation coefficient, but for *P. alba* grown on a hormone-free medium it was equal to 1 (Tikhomirova et al. 2016). It is likely, that different species of Rosaceae family have different breeding potential. In some cases, the use of hormones was necessary to increase the propagation coefficient. The use of hormones for *D. fruticosa* was unnecessary, as it displayed high propagation coefficient on a hormone-free medium, and the resulting plants were not overloaded with hormones.

**Planting D. fruticosa into soil.** Adaptation of test tube microplants to soil conditions is a complex process, therefore, conditions with gradual air and substrate moisture regulation in cultural vessels were conducted when transferring the microplants into soil. Substrate quality is an important factor of microplant adaptation. The use of a ready for use soil mixture for plant planting allowed us to make the transfer into soil gentle and effective. In result, microplant survival in the soil was more than 90%. Plants grew actively after the transfer, and for 350 plants allowed us to obtain more than 90% of sterile well-growing microplants, when introducing the Kuril tea, *D. fruticosa*, into culture. The explants started to grow actively on nutrient medium from the third (D1) to the sixth (D2, D3, D4) weeks of culturing. Despite some distinction between microplants of different cultivars (shoot length, internodes number, root number and fresh weight), all of them transformed into well-developed plants with abundant root system and strong shoots within 2 months of cultivation on hormone-free half-strength MS medium. The unified composition of medium for both growth initiation and proliferation, applied in this work, allows decreasing material costs and reagents consumption for microclonal propagation of *D. fruticosa* and cultivars. Application of a ready-made soil mixture for plant growing in controlled conditions (temperature, illuminance, and air humidity) led to a successful adaptation of more than 90% of test tube microplants to soil conditions.

**ACKNOWLEDGMENTS**

The authors are grateful to Mikhail Viktorovich Mikhailin for provided *D. fruticosa* cultivars (‘Lovely Pink’, ‘Bella Bianca’, ‘Bella Sol’) for microclonal propagation.

**LITERATURE CITED**


