Effect of Activated Carbon on Rooting of Tissue Culture Seedlings of Qiuzi Pear

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Abstract In order to improve the rooting rate of Qiuzi pear tissue culture seedlings, this study took ‘sorb’ as experimental material, based on Qiuzi pear rooting tissue culture medium (1/2MS+IBA 0.5 mg/L), and added different concentrations of activated carbon (AC): 0.25 g/L, 0.50 g/L, 0.75 g/L, 1.00 g/L. The results showed that the rooting rate, rooting number and root length of tissue culture seedlings cultured with activated carbon were significantly higher than those of the control group. The most suitable medium for rooting Qiuzi pear was 1/2MS+IBA 0.5 mg/L+AC 1.00 g/L, and the rooting rate of the treated ‘sorb’ tissue culture seedlings reached 90%, which could obviously promote the growth of Qiuzi pear tissue culture. The results provide a theoretical basis for improving the rooting rate and survival rate of tissue culture seedlings of pear plants, and provide a reference for the establishment of rapid propagation system of woody plants.

Keywords Qiuzi pear; Tissue culture technique; Activated carbon

Autumn pear (Pyrus ussuriensis Maxim.) is one of the important pear species in China, which is abundant in Northeast China, Hebei and Shandong. The seedlings of Qiuzi pear are often cold-resistant rootstocks of pears in orchards, which are excellent raw materials for producing juice drinks and also excellent greening tree species in autumn. The propagation of Qiuzi Pear is mainly carried out by seed propagation and grafting. It takes a long time for seed propagation, and the uniformity of offspring growth is inconsistent. The cultivation time of grafted seedlings is long, the technology is complex, the propagation efficiency is low, the virus is easy to spread quickly, and the degree of genetic heterozygosity is high (Xu et al., 2002). With the development of modern biotechnology, plant tissue culture technology (Xiao and Liu, 2011) can be applied in the rapid propagation of plants, variety improvement and germplasm resources preservation, which can realize the rapid acquisition of high-quality germplasm and contribute to the popularization and demonstration of excellent varieties (Miao et al., 2019). Using plant tissue culture technology, ‘sorb’ can be propagated rapidly and the seedlings can be consistent. In the traditional plant tissue culture technology, the rooting induction effect of pear is poor. In the research of Liu et al. (2016), the rooting rate of pear tissue culture seedlings is only 33%. Sun et al. (2015) found that adding activated carbon can improve the rooting quality and rooting rate of emerald pear in the study of tissue rapid propagation technology. In the rooting process of plant tissue culture, activated carbon (AC) can provide a dark environment for rooting, which is beneficial to root induction and root growth, prevent browning, increase the content of soluble protein and total sugar in culture, and adsorb plant growth regulators and substances beneficial to rooting (Sun et al., 2010), thus affecting the culture results. Therefore, by adding different concentrations of activated carbon to rooting medium, this experiment explored the effect of activated carbon on rooting efficiency of tissue culture seedlings, and provided reference and guidance for improving rooting efficiency of pear tissue culture seedlings.

1 Results and Analysis
1.1 Effects of different AC concentrations on the growth of tissue culture seedlings of ‘sorb’
On the 14th day, red young roots appeared in the roots of ‘sorb’ tissue culture seedlings. With the increase of activated carbon concentration, the number of red young roots increased obviously. On the 21st day, the roots grew...
obviously. On the 28th day (Figure 1), 35th day (Figure 2) and 41st day (Figure 3), the rooting rate and rooting situation in each culture medium were measured.

Figure 1 Rooting of ‘sorb’ seedlings with different AC concentrations on the 28th day
Note: when the concentration of activated carbon is 0.00 g/L, the rooting status and length of ‘sorb’ seedling; T1, when the concentration of activated carbon is 0.25 g/L, the rooting status and length of ‘sorb’ seedling; T2, when the concentration of activated carbon is 0.50 g/L, the rooting status and length of ‘sorb’ seedling; T3, when the concentration of activated carbon is 0.75 g/L, the rooting status and length of ‘sorb’ seedling; T4, when the concentration of activated carbon is 1.00 g/L. The rooting status and length of ‘sorb’ seedlings

Figure 2 Rooting of ‘sorb’ seedlings with different AC concentrations on the 35th day
Note: CK, activated carbon concentration is 0.00 g/L; T1, activated carbon concentration is 0.25 g/L; T2, activated carbon concentration is 0.50 g/L; T3, activated carbon concentration is 0.75 g/L; T4, activated carbon concentration is 1.00 g/L

Figure 3 Rooting of ‘sorb’ seedlings with different AC concentrations on the 41st day
Note: CK, activated carbon concentration is 0.00 g/L; T1, activated carbon concentration is 0.25 g/L; T2, activated carbon concentration is 0.50 g/L; T3, activated carbon concentration is 0.75 g/L; T4, activated carbon concentration is 1.00 g/L

1.2 Effects of different AC concentrations on rooting rate and rooting number of ‘sorb’ seedlings in tissue culture
As shown in Figure 4, on the 41st day of culture, the rooting rates of T2 and T3 were both 50%, T4 was 90%, CK was the lowest, which was 40%. The rooting rates of rooting seedlings cultured with activated carbon were higher than those of the control group. The rooting rate of T1 was 80%. The rooting rate of T4 was significantly higher than that of CK, T2 and T3.

In different treatments, there were significant differences in the rooting number of pear tissue culture seedlings, which were 2.2 for T1, 2.2 for T2, 3.6 for T3, 6.2 for T4 and 0.8 for CK. As shown in Figure 5, with the increase of activated carbon concentration, the number of rooting strips increased. T4 was significantly higher than CK, T1 and T2.
1.3 Effects of different AC concentrations on root morphological indexes of tissue culture seedlings of ‘sorb’

Measure the average root length of ‘sorb’ seedlings under different treatments on the 41st day (Figure 6). It can be seen from the figure that the average root length of T4 is significantly different. Compared with the other four groups of results, the total root length is far greater than other treatments.

As shown in Figure 7, with the increase of AC concentration, the root surface area of ‘sorb’ seedlings increased. The root surface area of T4 was significantly higher than that of other treatments.

According to Figure 8, with the increase of AC concentration, the volume of Qiuze pear seedlings increased.
Figure 6 Effects of different AC concentrations on total length of ‘sorb’ seedlings
Note: CK, activated carbon concentration is 0.00 g/L; T1, activated carbon concentration is 0.25 g/L; T2, activated carbon concentration is 0.50 g/L; T3, activated carbon concentration is 0.75 g/L; T4, activated carbon concentration is 1.00 g/L

Figure 7 Effects of different AC concentrations on the surface area of ‘sorb’ seedlings
Note: CK, activated carbon concentration is 0.00 g/L; T1, activated carbon concentration is 0.25 g/L; T2, activated carbon concentration is 0.50 g/L; T3, activated carbon concentration is 0.75 g/L; T4, activated carbon concentration is 1.00 g/L

Figure 8 Effects of different AC concentrations on seedling volume of ‘sorb’
Note: CK, activated carbon concentration is 0.00 g/L; T1, activated carbon concentration is 0.25 g/L; T2, activated carbon concentration is 0.50 g/L; T3, activated carbon concentration is 0.75 g/L; T4, activated carbon concentration is 1.00 g/L
2 Discussion
During the growth and development of plants, hormones in the body are in a dynamic process, and plant growth regulators can regulate the growth of tissue culture seedlings. Different varieties have different responses to the types and concentrations of growth regulators, and only by choosing appropriate hormone ratio can better proliferation effect be achieved (Xu et al., 2008). Qiu et al. (2015) thought that the most suitable rooting medium for North American pear was MS+TDZ 0.5 mg/L+ABT1 # 1.5 mg/L+Sucose 20 g/L+Agar 4.5 g/L+AC 2.0 g/L, and the rooting rate was 52.03%. Zhang et al. (2009) thought that the most suitable rooting medium for Huangguan pear was 1/2MS+IBA 1.0 mg/L+Sucrose 20 g/L, and the rooting rate was 33%. Wen and Zhang (2007) thought that the most suitable rooting medium was 1/2 ms+NAA 1.0 mg/L+IBA 1.0 mg/L+AC 3.0 mg/L, and the rooting rate was 95%. Sun et al. (2001) thought that the medium for strong seedling of Fengshui pear was 1/2 MS+BA 0.5 mg/L+GA 0.5 mg/L+3% sucrase. The experiment found that 1/2MS+IBA 0.5 mg/L+AC 1.0 g/L was suitable for rooting culture of ‘sorb’, and the rooting rate was 90%. It is difficult to take root in tissue culture of Pyrus. One of the important factors affecting rooting of tissue culture seedlings is the type and concentration of auxin. IBA, IAA and NAA are commonly used growth regulators to promote rooting. IBA was used for rooting and tissue culture seedlings of seedless Rosa roxburghii (Jiang et al., 2017) and Douli (Li et al., 2012), but the suitable concentration was different. 0.2 mg/L IBA is suitable for seedless Rosa roxburghii and 1.5 mg/L IBA is suitable for bean pear. In this experiment, the rooting of ‘sorb’ seedlings in tissue culture was studied. The results showed that the most suitable rooting medium for ‘sorb’ seedlings was 1/2 ms+IBA 0.50 mg/L+AC 1.00 g/L.

Light conditions and culture methods also have effects on adventitious root induction of tissue culture seedlings. Tang et al. (2006) found that early dark culture promoted the rooting induction of ‘Zaosu’ and ‘Shenbuzhi’, but had poor rooting induction effect on ‘Bali’ and ‘Komisi’. In this experiment, active carbon was added to provide a dark environment and placed under light conditions, which improved the rooting rate and rooting number of ‘sorb’.

Yang et al. (2008) induced Qiuzi pear to take root, the medium was 1/2MS+IBA 0.5 mg/L, the rooting rate was 45%, and the number of rooting strips was 3. When Wang et al. (2018) induced Qiuzi pear to take root, the medium was 1/2MS+IBA 0.5 mg/L, the rooting rate was 45.5%, and the number of rooting strips was 2.24. In this experiment, based on the rooting media used in Yang Fang and Wang Defen, activated carbon was added to promote the rooting of ‘sorb’ seedlings. The effect of the medium added with activated carbon was higher than that without activated carbon in rooting rate, rooting quantity, average root length, surface area and volume. When the concentration of activated carbon was 1 g/L (1/2MS+IBA 0.5 mg/L+AC 1.0 g/L), the rooting rate of ‘sorb’ seedlings was as high as 90%, and the number of rooting strips was 6.2. In this study, the efficiency and quality of tissue culture of ‘sorb’ seedlings were significantly improved by adding activated carbon, which provided reference for the tissue culture technology of fruit trees.

3 Test Materials and Methods
3.1 Test materials
The tissue culture seedlings of ‘sorb’ cultivated in Key Laboratory of Fruit Tree, College of Horticulture, Anhui Agricultural University were used as test materials.

3.2 Culture conditions
The activated carbon used in this experiment is black powder. When the culture medium is prepared, if it is added in advance, it will agglomerate and adhere to the pot wall. Therefore, it is advisable to add activated carbon after the culture medium melts and before sterilization. It is necessary to constantly shake the culture medium before solidification, so that it can be evenly distributed during solidification. Rooting medium was 1/2MS+IBA(0.5 mg/L)+AC (activated carbon concentration gradients were 0.25 g/L, 0.50 g/L, 0.75 g/L and 1.00 g/L, respectively). The medium was used after sterilization (sterilized in a sterilization pot at 121℃ for 20 minutes). The culture temperature is 25℃, the illumination is 2000 lx, and the illumination time is 24 h/d.
3.3 Test design
The experiment was conducted in November, 2020, with a single factor randomized block design. Take the tissue culture seedlings of ‘sorb’ in the same growth condition in the same period, and carry out increment culture on the tissue culture seedlings of ‘sorb’, and inoculate 2 plants in each bottle, totally inoculating 50 bottles. Taking tissue culture seedlings of ‘sorb’ as materials, more than one third of petiole was cut, and the petiole was beveled at 45 with a scalpel on an ultra-clean workbench. The cut seedlings of ‘sorb’ were put into rooting medium containing different concentrations of activated carbon for culture.

3.4 Index determination
Electronic vernier caliper and instruments regent scanner calibrated for image analysis with regent instruments software were used for determination and analysis. Rooting rate(%)=(number of rooted stem segments/number of inoculated stem segments)×100%
Rooting number(number)=total root number of stem segment/stem segment number of rooting
Average root length(cm)=total root length/total number of rooted plants

3.5 Data analysis
SPSS software (IBM SPSS statistics 26) was used for data statistical analysis, Duncan method was used for variance analysis and multiple comparison (p<0.05).

Authors’ contributions
Luan Xiaolong is the executor of this experimental study, finishing data and writing the first draft of the paper; Shi Hao participated in experimental design and analysis of experimental results, Xu Bo and Zhang Qiannan participated in some experimental studies, and Li Liu was the project leader, guiding the design of experimental studies and the revision of papers. All authors read and approved to the final manuscript.

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