Effect of Cooling of Medium on Fruit Set in High-bench Strawberry Culture

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Abstract. We demonstrated the effect of cooling of the medium on the fruit set of strawberries (Fragaria × ananassa Duch.) grown on high benches for forcing culture. The cooling by water evaporation promoted by a fan enabled to cool the medium by an average of several degrees compared with no cooling. When runner plants were transplanted in late summer, cooling accelerated flower bud emergence almost 10 days on the primary axillary branch compared with plants grown in uncooled medium. Also, with cooling, fruit was harvested from the inflorescence of the primary axillary branch almost 10 days earlier. We expect that this technique will allow early transplanting around the end of summer and will shorten the time between fruit set on the terminal inflorescence and that on the inflorescence of the primary axillary branch.

Materials and Methods

High-bench culture with medium cooling. The high-bench culture system was constructed of plastic film bags on a framework (Ikeda et al., 2006). The film bag containing the culture medium was composed of two layers: an inner layer of silver-colored plastic film and an outer layer of cotton cloth. During the day, the cotton layer was kept wet by dripping water from the framework. The heat of the culture medium evaporated the water from the cotton surface of the bags, and the culture medium was cooled by the latent heat of water evaporation (Ikeda et al., 2006). Underneath the culture system, a polyethylene duct connected to a fan (Ka-120 Nepon, Tokyo) allowed more water to evaporate from the cotton.

The culture medium contained carbonized chaff and peatmoss (Majestic, Agawam, Mass.) in a ratio of 1:1. Water for irrigation of the culture medium was supplied by a line differentiated from that used for medium cooling. Initially, 15 g slow-release fertilizer (Rongotani, 13.0N–10.8P–4.8K; JA, Tokyo) and 10 g of lime (JA) per runner plant were contained in the culture medium. Irrigation water was supplied to the culture medium twice a day during the experiment. In the 2003 experiment, the excess irrigated water was drained off, but in the 2005 experiment, we modified the system to reuse it. The pH of the drainage in the 2005 experiment was kept by addition of potassium hydroxide. We supplied liquid fertilizer (Otsuka Chemical Co. Ltd., Osaka, Japan) as necessary. The high-bench culture system was constructed in an arched plastic-film greenhouse.

We designed two regimes: dripping water with fan cooling and control (no water and no fan). Water was supplied to the cotton cloth during the day (from 6 AM to 6 PM) between 4 Sept. and 3 Oct. 2003 and between 2 Sept. and 30 Sept. 2005. The fan operated over the same period. The temperature of the medium was determined by thermocouples placed 5 cm below the surface (medium depth was 10 cm) and recorded by a data logger (Solac III MP-090; Eko Instrument Co. Ltd., Tokyo).

Plant materials. Pot-grown strawberry (Fragaria × ananassa Duch. cv. Sachinoka) runner plants with four leaves were used. In the 2003 experiment, they were stored in a growth cabinet in total darkness at 15 °C for 15 d (total darkness treatment) and were then transplanted into the culture system on 4 Sept. We examined 16 runner plants for each treatment. In the 2005 experiment, the plants were stored in a growth cabinet in the dark for 16 h at 15 °C and then kept in a shaded greenhouse for 8 h during the day (8 AM to 4 PM) for 20 d (short-day treatment) and transplanted into the system on 2 Sept. We examined 58 runner plants for each treatment. Honeybees were used for pollination. The numbers of flowers were fixed to 11 for the terminal inflorescence and nine for the inflorescence of the primary axillary branch in the 2003 experiment. In the 2005 experiment, the numbers were fixed to nine and seven, respectively.

Results

Figure 1 shows the temperature in the greenhouse and in the culture medium in the (A) 2003 and (B) 2005 experiments. All data are the means of temperatures taken hourly from 7 AM to 6 PM each day. The temperature of the control medium was always higher than that of the cooled medium. The greenhouse was covered by thin shadecloth in the 2005 experiment for 2 weeks after transplant, the temperature of the control medium was lower than that in the 2003 experiment.

Flower bud emergence on the primary axillary branches under the cooling regime was always faster than in the controls in the 2003 experiment (Fig. 2A). The emergence rate was 38% in the controls and 63% under cooling regime on 14 Nov. Flower bud emergence reached 100% on 17 Nov. with cooling but did not reach this value until 25 Nov. in the controls. In the 2005 experiment, on 1 Dec., the percentage flower bud emergence under the cooled regime was 62%, but in the controls, it was only 33% (Fig. 2B). Under the cooling regime 80% emergence was reached on 7 Dec., but in the controls, it was not reached until 15 Dec.

After the fruits of the terminal inflorescence had been harvested from November to the beginning of January, harvesting of the fruits of the primary axillary lateral branch of the controls in the 2003 experiment started on 23 Jan., but under the cooling regime, it began on 16 Jan. (Fig. 3A). The harvest of the two regimes was almost 7 d apart with the same quantity of fruits being harvested (Fig. 3A; e.g., to reach for two fruits, on 30 Jan. for the cooling regime but on 6 Feb. for the control). In 2005, the harvest showed a maximum 17-d difference (Fig. 3B; e.g., to reach for two fruits, on 25 Feb. for the cooling regime but on 14 Mar. for the control).
Discussion

The use of high-bench strawberry culture is increasing in Japan because it saves labor and can reduce the likelihood of problems with soil-borne diseases. Our cooling system for high-bench culture reduced the temperature of the culture medium (Fig. 1). Although the temperature of the medium never dropped below the wet bulb air temperature of the greenhouse, theoretically (Takaichi et al., 2001), the decrease in temperature was enough to accelerate flower-bud emergence and fruit set on the primary axillary branch. Because induction of flowering on the primary axillary branch presumably started during the cooling operation, flower-bud emergence on this branch was maybe faster than in the controls (Fig. 2). The flower-bud emergence and harvesting times of fruits of the terminal inflorescence did not differ between the regimes in the 2003 and 2005 experiments (data not shown), presumably because flower bud induction treatment on the terminal inflorescence had already started during cold storage (i.e., flower bud induction treatment). Because there were no duration of flower bud emergence on the terminal inflorescence (data not shown), it might have been possible to transplant the runner plants into the cooled system earlier (e.g., in late August). If so, this could give a greater marketing advantage to growers. Also, if cooling were to be used even earlier in spring again, it might be possible to cultivate more axial branches for June-bearing cultivars.

Some researchers in Japan have attempted before to remove the summer heat from culture media in high-bench culture systems (Komori and Kanke, 2002; Takaichi et al., 2001). Takaichi et al. (2001) were the first to introduce a system of cooling media by latent heat. We modified their system to make it simpler and more effective in evaporating water (Ikeda et al., 2006), and here we have demonstrated the practical use of the system for forcing culture of strawberries.

Ganmore-Neumann and Kafkafi (1983) found that a high root temperature (32°C) decreased dry weight and leaf area. Utagawa et al. (1989, 1991) observed that a relatively high temperature (23°C) in the culture solution decreased root weight, nutrient absorbance, and photosynthesis rate in ever-bearing cultivars in a nutrient film technique system. Also, Lieten (1997) observed that higher root temperatures negatively affected vegetative development in his experiment ranging from 12 to 24°C.

With this system, because it was impossible to decrease the temperature of the medium below 23°C when air temperature was high (Fig. 1), the roots of the runner plants could have been damaged by the high temperatures. However, under the cooling regime, the medium temperature rose above 23°C for only a few hours a day, even in late summer (data not shown). Thus, the cooling system allowed the runner plants to maintain their growth. Another point is that the use of refrigeration for cooling media might not be
practical because of the costs of electricity and equipment. Our system requires electricity only for pumping for irrigation and for operating the fan (apart from that needed for greenhouse climate control); it is simple and easy to handle (Ikeda et al., 2006).

In everbearing cultivars, fruits size and yield decrease above 23 °C, although fruits number increases up to 27 °C (Wagstaffe and Battey, 2004). Kumakura and Shishido (1995) also found 20 to 25 °C was optimum for the growth of four everbearing strawberry cultivars. However, June-bearing strawberry cultivars had a lower optimum temperature of ≈15 °C (Le Mière et al., 1998). Also, Lieten (1997) reported that higher root temperatures had a negative effect on fruit size in his experiment ranging from 12 to 24 °C. In our study, fruit number was fixed for both regime, and the average weight of fruits from primary axillary branch was 17.2 g under the cooling regime and 16.9 g for controls in the 2003 experiment; there was no significant difference in fruit weight between the regimes. Because the season at the time of fruit development on the primary axillary branch was already cold, fruit weight might not have been affected in the same way as flower-bud initiation.

In conclusion, cooling of the high-bench culture medium allowed the runner plants to be transplanted in late summer and accelerated flower-bud initiation on the primary axillary branch in forcing culture of strawberries.

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