Pollination and CPPU Treatment Increase Endogenous IAA and Decrease Endogenous ABA in Muskmelons during Early Development

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ABSTRACT. An investigation was conducted to determine how pollination and CPPU treatment influence endogenous IAA and ABA content in netted muskmelon [Cucumis melo L. (Reticulatus Group) ‘Crest Earl’s’], and to clarify their roles in fruit set and development in relation to these endogenous plant hormones. CPPU treatment at anthesis significantly increased the fresh weight of ovaries, whether the flowers were pollinated or not, but from 6 days after anthesis (DAA) the growth rate in the nonpollinated + CPPU treatment tended to be lower than the growth rates in the pollination treatment plots. Ovaries of nonpollinated flowers not treated with CPPU failed to grow and turned brown within 4 DAA. IAA content in the placenta of fruit from pollinated flowers increased rapidly from the day of anthesis to 2 DAA and remained at relatively high levels. IAA content in the placenta of parthenocarpic fruit induced to develop by CPPU treatment was lower than that of fruit from pollinated flowers but the pattern was almost the same as that in fruit of pollinated flowers. Conversely, IAA content in the placenta of fruit from nonpollinated flowers not treated with CPPU decreased sharply after anthesis. IAA content in the mesocarp of CPPU-treated fruit, whether or not the flowers were pollinated, increased significantly from the day of anthesis to 2 DAA, then decreased to almost the same level as that of the pollination-only treatment by 10 DAA, while the IAA content of nonpollinated CPPU-treated fruit declined even further. IAA content in the mesocarp of fruit from nonpollinated flowers not treated with CPPU decreased sharply. ABA contents in both the placenta and mesocarp of muskmelon that would set decreased after anthesis while the ABA content of muskmelon that would not set increased rapidly. Results suggest that pollination and CPPU treatment increased endogenous IAA content and decreased endogenous ABA content to promote the set and growth of fruit during early development. Chemical names used: [1-(2-chloro-4-pyridyl)-3-phenylurea] (CPPU); indole-3-acetic acid (IAA); abscisic acid (ABA).

Fruit grow most rapidly during the early stages following anthesis, and whether plants will retain or shed these fruit is determined at this time. Pollination and fertilization are the main predictors of fruit growth during these early stages and it has been suggested that seed-produced auxin is involved in fruit set and development (Weaver, 1972). This view is supported by reports showing that IAA content in the seeds is higher than that in the shoot of strawberry (Fragaria x ananassa Duchesne) (Nitsch, 1950), peach [Prunus persica L. Batsch (Peach Group)] (Miller and Walsh, 1990) and tomato (Lycopersicon esculentum Mill.) (Hoher et al., 1992).

Effects of plant bioregulators on fruit set of muskmelon (Cucumis melo L.) has been investigated and auxin-derivative plant hormones, such as p-chlorophenoxyacetic acid, 2,4-dichlorophenoxyacetic acid, and naphthaleneacetic acid, as well as chemicals with cytokinin activity such as 6-benzylaminopurine and CPPU, all promote fruit set (Hayata et al., 2000). CPPU promotes fruit set even under conditions that are unfavorable for pollination and fertilization. However, few studies have investigated the relationship between fruit set of muskmelon and endogenous growth substances. Xu et al. (1989) reported that IAA content increased during the period when fruit grew rapidly in the early growth stage. Although Lingle and Dunlap (1991) reported that IAA levels were the highest at 2 d before anthesis, they also showed that IAA levels increased after anthesis to 2 d after anthesis (DAA), and that there were some increases in free IAA, esterified and amide forms of IAA during the maturing period. On the other hand, Lee et al. (1997) reported that IAA content in the ovary did not fluctuate much during early development. Although IAA appears to be an important plant hormone in the growth of muskmelons, changes in endogenous IAA activity in muskmelons during development are inconsistent among those studies. Research on ABA content during early development of muskmelons is limited, with one report that ABA decreased during development (Xu et al., 1989). Therefore, the following study was conducted to investigate how pollination and CPPU treatment at anthesis influence fruit set, growth, and endogenous IAA and ABA levels in muskmelons.

Materials and Methods

PLANT MATERIAL AND CULTURE. Seedlings of ‘Crest Earl’s’ netted muskmelon [Cucumis melo (Reticulatus Group)] were transplanted 45 cm apart in two rows (1.2 m wide × 17 m long, two beds) in a heated greenhouse at Hiroshima Prefectural University Hiroshima, Japan, on 1 June 1999. The plants were grown under natural photoperiod and irradiance from 1 June through 26 Sept. Air temperature was maintained between 18 and 28 °C. The soil in the beds was a sandy loam with 2% organic matter. Soil pH was 6.5. Irrigation of the beds was regulated automatically by a tensionmeter (DM-8P Melon; Takemura Electric Works Ltd., Tokyo). Fertilizer was applied twice. The first application was a preplant broadcast application of 900 kg·ha⁻¹ of 14N–14P–36K; the second application was a sidedress application at anthesis of N at 326 kg·ha⁻¹. Each lateral shoot between the 13th and 15th nodes of the main shoot was cut above the 2nd node; and female flowers adhering to the first nodes of the lateral shoots were used in this experiment. Other lateral shoots were removed. The main shoots were trained upwards and topped at the 24th node.

TREATMENTS. Forty plants were assigned randomly to each of
four treatment groups and treatments were applied as follows: female flowers were hand-pollinated at anthesis without any other treatment (P); female flowers were hand-pollinated and sprayed with a solution of CPPU at 20 mg·L⁻¹ (Sigma Chemical Co., St. Louis) at anthesis (PC); female flowers covered with paper bags from the day before anthesis were not pollinated (E); and the nonpollinated female flowers were sprayed with a solution of CPPU at 20 mg·L⁻¹ at anthesis (EC). The paper bags were removed 3 DAA. Fruit in all treatments were sampled the day before anthesis and on 0, 2, 4, 6, and 10 DAA. Each harvested fruit was placed immediately in a cold room (4 °C) where it was sliced horizontally and the slices divided vertically into four sections. The placenta (containing ovule) and the mesocarp of each fruit were sampled and stored at –80 °C until analysis.

**IAA AND ABA ANALYSES.** IAA and ABA contents were measured by gas chromatography–mass spectrometry–selected ion monitoring (GC–MS–SIM) according to a modified procedure of Kojima et al. (1999). Ten grams of frozen sample and 100 mL of 80% (v/v) cold methanol containing butylated hydroxytoluene at 20 mg·L⁻¹ were homogenized for 1 min in a GLC homogenizer (Yamato Scientific Co. Ltd., Tokyo) at 30,000 rpm. The homogenate was centrifuged at 13,000 g for 10 min, and the residue extracted once again. Supernatants were pooled, ¹³C₆-IAA (Kojima, 1995) and [3',5',5',7',7',7'-2H₆]ABA [purity 99%, Shoko Co., Ltd., Tokyo, synthesized according to the method of Rivier et al. (1977)] were added as internal standards, and the mixture concentrated at 35 °C to the aqueous phase using a rotary evaporator. The aqueous phase was centrifuged at 13,000 g for 10 min, and the residue extracted once again. Supernatants were pooled, and the mixture concentrated at 35 °C to the aqueous phase using a rotary evaporator. The aqueous phase was acidified to pH 2.8 with 6 N phosphoric acid and again partitioned three times against ethyl acetate. The ethyl acetate fraction was evaporated, and the residue was dissolved in 1 mL 80% methanol and purified using the Sep-Pak C18 cartridge (Waters Corp., Milford, Mass.). The extract was fractionated with a Shimadzu-10A high-performance liquid chromatography (HPLC) system (Shimadzu Co., Ltd., Kyoto, Japan) equipped with a Shimadzu SPD-10A UV-VIS ultraviolet spectrophotometer at 254 nm. The HPLC column was a Shimpack CLC-ODS (150 × 6.0 mm i.d., Shimadzu Co., Ltd.) maintained at 35 °C with a 40% methanol solution adjusted to pH 3.5 with acetic acid at a flow rate of 1 mL·min⁻¹. The IAA and ABA fractions were dried in vacuo and methylated by adding an excess of diazomethane in diethyl ether. The methylated IAA sample was dried with N₂ and then trimethylsilylated with 20 μL of N-methyl-N-trimethylsilyl-trifluoroacetamide (MSTFA, Pierce Chemicals Co., Rockford, Ill.). The methylated ABA and the trimethylsilylated IAA were dissolved with 50 μL methanol and injected into a QP5050 GC–MS system (Shimadzu Co., Ltd.) with selected ion monitoring, respectively. A DB-1 (J & W Scientific, Folsom, Calif.) capillary column (30 m × 0.32 mm i.d. × 0.25 μm film thickness) was coupled directly to the ion-source interface, the temperature of which was 250 °C, and the column temperature was maintained at 50 °C for 2 min, then increased at 30 °C/min to 170 °C, which was maintained for 2 min, and then at 10 °C/min to 250 °C. The helium carrier gas inlet pressure was 50 kPa. IAA and ABA contents were calculated by monitoring target ion m/z 202/208 and reference ion m/z 261/267, and target ion m/z 190/194 and reference ion m/z 162/166, respectively.

**DATA ANALYSIS.** Means and se values were calculated based on the data obtained from 10 replications for fresh weight (FW) and

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**Fig. 1.** Effects of pollination and CPPU treatment on fresh weight of ‘Crest Earl’s’ muskmelon fruit during the early growth stage. Each symbol represents the mean (n = 10) ± se.

**Fig. 2.** Effect of pollination and CPPU treatment on the IAA content of (A) placenta and (B) mesocarp of ‘Crest Earl’s’ muskmelon fruit during the early growth stage. Each symbol represents the mean (n = 3) ± se.
three replications for IAA and ABA analyses by each sampling time.

Results and Discussion

CPPU treatment at anthesis significantly increased fruit FW whether flowers were pollinated or not, and at 2 DAA the FW increased 2-fold compared with treatment P. Thereafter, fruit growth rate in treatment P increased, and FW became the same as that in treatment PC at 10 DAA, while FW in treatment EC showed the reverse pattern (Fig. 1). These results are consistent with our previous finding (Hayata et al., 2000) and confirm the promoting effect of CPPU on growth of young muskmelons. Nonpollinated fruit failed to grow; instead, FW decreased by 20% by 2 DAA, the ovary turned yellow and then brown by 4 DAA.

IAA content in the placental tissues of the muskmelons was 85.5 ng·g⁻¹ FW on the day before anthesis, and increased thereafter, whether the flower was pollinated or not (Fig. 2A). In both treatments P and PC, IAA content in the placenta increased rapidly from the day of anthesis to 2 DAA and was maintained at high levels even though it declined slightly thereafter; IAA content in the placenta of parthenocarpic fruit induced by CPPU was lower than the IAA contents of pollinated fruit, but showed almost the same changes as in the pollinated fruit. IAA content in the placenta of treatment E decreased dramatically, and became undetectable by 2 DAA.

IAA content in the mesocarp of CPPU-treated fruit increased more than the fruit in the P treatment group from the day of anthesis until 2 DAA, regardless of pollination (Fig. 2B). Thereafter, however, IAA content decreased; IAA levels in the PC treatment group decreased to almost the same as the levels in the P group, and IAA levels in the EC treatment group became lower than those in both the PC and P groups (34 ng·g⁻¹ FW). IAA content in the mesocarp of the E treatment group decreased as rapidly as that in the placenta, and became undetectable by 2 DAA; that is, changes in IAA content in the mesocarp of nonpollinated fruit were similar to those in the placenta. In addition, since there was a concentration gradient in IAA from the placenta to the mesocarp, IAA levels are presumed to move along a gradient. Our previous observation supports this speculation: a concentration gradient in IAA moved along from the normal seeds to the placenta and toward the mesocarp (Li et al., 2002).

Gustafson (1939) reported that auxin was a key substance for fruit set, based on his observations that IAA content in the ovary of parthenocarpic grape (Vitis labrusca L.), orange (Citrus aurantium L.), and lemon (Citrus limon Burm.f.) was higher than IAA content in nonparthenocarpic ovaries at anthesis. Takeno and Ise (1992) reported that IAA content in the ovary of parthenocarpic cucumber (Cucumis sativus L.) was three times higher than in cucumber that dropped fruit several days after anthesis. Since we observed that pollination increased IAA content in the placenta and ovary of netted muskmelon, IAA appears to be involved in fruit set. This is supported by the observation that application of CPPU, a fruit set promoting treatment for muskmelon, increased IAA content. Thus, it is appears that parthenocarpy induced by CPPU and the promotive effect of CPPU on fruit set are closely associated with biosynthesis of endogenous IAA, and it is clear that the promotive effect of CPPU on biosynthesis of IAA is temporary, occurring only at the time of treatment. There have been some reports about IAA levels in the ovary of melon at anthesis. Lingle and Dunlap (1991) showed that although the highest IAA level was reached before anthesis, IAA levels increased again after anthesis to 2 DAA, and Xu et al. (1989) reported that IAA levels increased after pollination. These increasing patterns coincide with our results. Furthermore, CPPU treatment increased the growth of the ovary at anthesis when cell division was still occurring (Masuda and Hayashi, 1959); that is, CPPU promoted cell division using its own active cytokinin (Takahashi et al., 1978). It is also possible that CPPU promoted biosynthesis of IAA, which is involved in cell enlargement, resulting in enhancement of growth.

Baldi et al. (1989) reported that under alkaline extraction conditions, IAA ester was hydrolyzed to free IAA. However, they showed that significant hydrolysis occurred only after a 2-h exposure to pH 9.0 or above. Since our samples were exposed to mild alkaline conditions for <1 h during extraction, hydrolysis of IAA-ester would have been minimal. Additionally, it was apparent that there were differences in IAA levels between the treatments.

Endogenous ABA content in the mesocarp and placenta of muskmelons was =180 to 190 ng·g⁻¹ FW at anthesis, and ABA levels of all treatment groups except E decreased gradually by 6 DAA, and thereafter maintained almost the same level (Fig. 3A). This declining pattern of ABA in fruit agrees with findings of Xu et al. (1989). On the other hand, ABA content in treatment E increased rapidly from the day of anthesis, and at 2 DAA it was three times higher than that of the day before anthesis: 578 ng·g⁻¹ FW in the placenta (Fig. 3A), and 620 ng·g⁻¹ FW in the mesocarp (Fig. 3B). In addition, previous studies showed the same declining pattern of ABA content in satsuma mandarin [Citrus unshiu (Mak) Maric.] (Kojima et al., 1996), persimmon (Diospyros kaki
Thunb.) (Kojima et al., 1999), strawberry (Archbold and Dennis, 1984), and pear (Pyrus communis L.) (Gil et al., 1972). It may be that fruit set in various fruits, including muskmelon, requires a decrease of ABA content.

Results herein demonstrate that pollination and CPPU treatment increase endogenous IAA content of muskmelons and decrease endogenous ABA content at fruit set. The presence of high IAA content and low ABA content may be closely related to fruit set and growth during the early growth stages. Moreover, it appears that CPPU not only operates directly as a cytokinin but also induces parthenocarpy and promotes growth in the early growth stage periods by increasing endogenous IAA in muskmelons.

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