Genotype, Temperature, and Fall-applied Ethephon Affect Plum Flower Bud Development and Ovule Longevity

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Abstract. Flower bud growth and ovule longevity of plum (Prunus domestics L.) cultivars Italian and Brooks and the effects of fall-applied ethephon and of temperature were studied. Fresh and dry weights of terminal flower buds were measured at 1-week intervals from 50 days to 1 day before bloom in 1988. Buds were also analyzed for N, P, K, Ca, and B. After bloom, ovule longevity was determined using a fluorescence method after staining with aniline blue. Ovule longevity was determined in 1990 using shoots excised at full bloom from untreated and ethephon-treated trees of both cultivars and held in growth chambers for 18 days at 5, 10, 15, or 20°C. ‘Brooks’ flower buds showed a higher accumulation of fresh and dry weight than ‘Italian’, and ethephon reduced bud weights in both cultivars. Ethephon did not affect mineral content of flower buds of ‘Brooks’, but ‘Italian’ flower buds contained a higher concentration of Ca and a lower concentration of P when treated with ethephon. Boron content was higher in the ethephon-treated buds of ‘Italian’ trees on some sampling dates. Ovule longevity was higher for ‘Brooks’ than for ‘Italian’ in both years. Ethephon treatment delayed ovule senescence in ‘Italian’ flowers, but had little or no effect on ‘Brooks’ flowers. Increasing temperatures induced faster ovule senescence in both cultivars. Chemical name used. 2-chloroethylphosphonic acid (ethephon).

Insufficient fruit set in plum trees may be due to a genetic predisposition for abnormal embryo sac development (Thompson and Liu, 1973), low temperature during and after bloom time (Jaumien, 1968), or B deficiency in the flowers that results in poor pollen tube growth (Hanson and Breen, 1985a). Poor fruit set may result from a shortened effective pollination period (EPP), the difference between ovule longevity and the time required for the pollen tube to reach the egg sac (Stösser and Anvari, 1982; Williams, 1966, 1970). A short EPP can result from slow pollen tube growth rates and/or short ovule longevity.

Fall ethephon applications delay flowering in several fruit tree species (Coston et al., 1985; Dennis, 1976; Gianfagna et al., 1986; Proebsting and Mills, 1973; Webster, 1986), but such treatments have often not resulted in increased fruit set and yield. The benefit of bloom delay is often decreased by side effects, such as gummosis, floral abscission, or failure of floral buds to open (Coston et al., 1985; Dennis, 1976; Proebsting and Mills, 1973). Crisosto et al. (1990) reported bloom delay but decreased yield in peaches [Prunus persica (L.) Batsch.]. Bloom was also delayed when ethephon was applied to ‘Italian’ prune (Crisosto et al., 1990). However, a significant increase in yield was observed in 1 year of trials. Durner et al. (1990) reported that in a 2-year trial, ethephon treatment enhanced yield of ‘Cresthaven’ peach in both years, but that of ‘Jerseydawn’ in only 1 year. Regardless of its effect on yield, the ethephon treatment caused smaller pistils to develop and heavier prethinning crop loads, which led to smaller fruit and a later harvest date.

Temperatures up to 3 weeks after full bloom had a pronounced effect on the rate of ‘Italian’ prune embryo sac development and abortion (Thompson and Liu, 1973). Keulemans (1984) found the same relationship, but also that genotypes varied in sensitivity to cool postbloom temperatures. Shorter ovule longevity in warm than cool seasons adversely affects fertilization in sweet cherry [Prunus avium (L.) L.] (Eaton, 1959) and apple (Malus domestica Borkh.) (Dorsey, 1929; Williams, 1965). Cool weather may prolong ovule longevity, but retards pollen tube growth sufficiently so that fertilization is not possible before ovular breakdown (Williams, 1970).

The objectives of this study were to determine the effect of genotype and fall ethephon application on flower bud growth, mineral content of buds, and ovule longevity of flowers from the plum cultivars. ‘Brooks’ and ‘Italian’ were selected because of their consistent and erratic annual production, respectively. The effect of temperature on ovule viability of the two cultivars, with or without a fall ethephon application, was determined.

Materials and Methods

Ethephon was applied at 0 or 500 mg-liter to runoff with a hand gun sprayer to ‘Italian’ and ‘Brooks’ prune trees at the 60% leaf drop stage on 9 Nov. 1988 and 26 Oct. 1989. Regulaid [0.1% (v/v)] was added to assist spreading. The orchard used in this study was planted in 1975 and located at the Lewis-Brown Horticulture Research Farm, Oregon State Univ., Corvallis. Ten single-tree replicates were used for each treatment.

Growth measurements. Three replicates of 100 terminal flower...
buds were excised from the terminal region of 2 to 3-cm-long spurs at \( \approx 1 \) week intervals, beginning 17 Feb. 1989 for the four treatments. Fresh and dry weights were obtained for each sampling date.

**Tissue mineral analysis.** Nutritional analysis of the buds was performed for each sampling date. A modified Kjeldahl method (Schuman et al., 1973) was used for total N analysis. Inductively coupled plasma spectrometry (Isaac and Johnson, 1985) measured P, K, Ca, and B.

**Ovule longevity determination**

**Field experiment.** Flower buds were emasculated before bloom to prevent self-pollination and covered with cheesecloth bags to prevent pollination by bees. Ten flowers, excised from the apical position of a spur (\( \approx 2 \) cm) for each treatment, were sampled every 2 days from full bloom until 20 days after full bloom (DAFB). The number of open flowers per limb were counted periodically and full bloom was considered as the time when at least 80% of the flowers were open. The full-bloom dates for the first year of study were 11, 12, 12, and 13 Apr. 1989 for ‘Brooks’, ‘Brooks’ + ethephon, ‘Italian’, and ‘Italian’ + ethephon trees, respectively. In the second season, the recorded full-bloom dates for the same treatments were 30 Mar. and 3, 1, and 10 Apr. 1990. Five flowers of each treatment were sampled every 3 days from full bloom until 15 DAFB. Samples were fixed in 5 formaldehyde : 5 propionic acid : 90 95% ethanol (FAP) (by volume) until observations were made.

Flowers. were removed from the FAP solution, washed in distilled water for 3.0 rain, soaked in 1% sodium bicarbonate for 1 h, and rinsed three times in distilled water. Pistil and ovary samples were softened by autoclaving in 1% sodium sulfite for 2 min. Pistils were excised, ovaries were split longitudinally, and the two ovules were removed with the aid of fine forceps under a light microscope. Ovules were then mounted on slides by direct squash in 0.5% (w/v) aqueous aniline blue (C.I. 42755; Baker Chemical Co., Phillipsburg, N.J.) in 0.15 M potassium phosphate buffer. Slides were observed under a Carl Zeiss (Oberkochen, Germany) universal fluorescence microscope equipped for epi-illumination with near-ultraviolet excitation (Crisosto, 1987; Pimienta et al., 1983).

Ovule longevity was tested by differential intensity of ovule fluorescence after staining with aniline blue. Strong fluorescence of the ovule at the chalazal end indicates nonviable ovules (Polito and Pimienta, 1982; Stosser and Anvari, 1982). Each flower was rated for the number of viable (nonfluorescing) ovules of temperature and ethephon treatment on ovule longevity, three flowers per twig were sampled for the 10 flowers per treatment on each sampling date. A modified P, K, Ca, and B.

**Growth chamber experiment.** In 1990, to determine the effect of temperature and ethephon treatment on ovule longevity, three replicates of 20 twigs each, up to 15 cm long, were cut from flowering trees of the two genotypes and placed in shallow trays in moist florist’s foam (Oasis Floral Products, Smithers-Oasis, France) (Jefferies et al., 1982). Three flowers per twig were emasculated to prevent self-pollination. The pots were then transferred to controlled environment chambers. Conditions were 5, 10, 15, and 20°C, with a 16-h light/8-h dark period. A Murashige and Skoog basal salt mixture (M5524; Sigma Chemical, St. Louis) solution was added to the trays to fulfill nutritional requirements. Distilled water was subsequently added periodically to avoid moisture loss. Samples of 10 flowers were taken every 2 days for at least 18 DAFB, depending on the temperature. The flowers were fixed in FAP solution and prepared as above.

**Statistical analysis.** Both experiments were conducted as a completely randomized design. Data for the tissue mineral analysis and growth measurements were analyzed using two models. The data for fresh weight (FW), dry weight (DW), and N and P content were fitted to a nonlinear analysis of covariance, exponential growth model:

\[
Y_{ij} = \alpha_i \exp(\beta_i \times d_{ij}) + \tau_i + \epsilon_{ij}; \\
i = 1-4, \ j = 1-8. \]  

The ij’t’ response is denoted \( Y_{ij} \), the \( j \)’th regression is denoted \( d_{ij} \) and day 1 = 0 corresponds to 17 Feb. The index \( i = 1, 2, 3, 4 \) corresponds to the treatments ‘Italian’ (1), ‘Italian’ + ethephon (1 + E), ‘Brooks’ (B), and ‘Brooks’ + ethephon (B + E), respectively. The random error terms are assumed to be independent and identically distributed with a mean of zero. The 12 parameters for \( \alpha_i, \beta_i, \) and \( \tau_i \) were estimated by nonlinear least squares. The parameters, \( \alpha, \beta, \) and \( \tau \) do not have a simple geometrical and biological interpretation. Consequently, instead of testing whether each individual parameter differed among treatments, we decided to test jointly whether all three parameters differed among treatments. This procedure corresponds to testing whether the regression functions coincide. The hypotheses of coincident responses among treatments were tested with likelihood ratio, approximate F tests (Seber and Wild, 1989).

The responses of P and Ca were fitted to the linear analysis of covariance model:

\[
Y_{ij} = \alpha_i + \beta_i \times d_{ij} + \epsilon_{ij}; \ i = 1-4, \ j = 1-8. \]

The notation is the same as in the exponential growth model. The eight parameters for \( \alpha, \beta, \) and \( \epsilon \) were estimated by least squares. We again tested whether the regression lines were coincident.

For B, actual B content, rather than B concentration, for a sample of 100 buds was obtained. Ovule longevity data for the field experiment, as well as B content of buds, were analyzed by using a general linear model procedure in SAS (SAS Institute, 1987). Differences among treatments were tested by using orthogonal contrasts.

Data from the growth chamber experiments were combined for a response surface regression analysis. Separate regressions were performed for each cultivar/ethephon treatment/temperature combination for a total of four analyses. The regression model used was a complete quadratic response surface model of the form:

\[
Y = \beta_0 + \beta_1 X_1 + \beta_2 X_2 + \beta_3 X_3 + \beta_4 X_4 + \beta_5 X_1 X_2 + \beta_6 X_1 X_3 + \beta_7 X_1 X_4 + \beta_8 X_2 X_3 + \beta_9 X_2 X_4 + \beta_{10} X_3 X_4 + \beta_{11} X_1 X_2 X_4 + \beta_{12} X_1 X_2 X_3 + \beta_{13} X_2 X_3 X_4 + \beta_{14} X_1 X_2 X_3 X_4, \]

where \( Y \) = number of viable ovules, \( X_1 = \) temperature, and \( X_2 = \) DAFB. Components were deleted from the model if their parameter estimate (\( \beta \)) was not significant (\( \alpha = 0.05 \)). If a component increased the mean square error (\( \delta^2 \)) or if its deletion did not decrease the \( R^2 \) of the model, it was also deleted. Results are presented as graphs of the predicted values from the models.

**Results**

**Growth rate and mineral content of buds.** Ethephon-treated flower buds of both cultivars had different growth rates when compared to their controls. Tests of hypothesis and probability levels for the equations fitted to each particular treatment (Table 1) show the significance of these differences. The model for FW and DW has an \( R^2 \) of 0.989 and 0.986, respectively. In both cases, the ethephon treatment resulted in a lower rate of bud growth (Fig. 1). ‘Brooks’ flower buds showed a higher accumulation of FW and DW than ‘Italian’ buds.

The model for N has an \( R^2 \) of 0.982 (Table 1); however, there were no differences in N concentration between the treatments (Fig. 1). For K, the model has an \( R^2 \) of 0.988 (Table 1).

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Table 1. Tests of hypothesis and probability levels for the exponential growth model fitted to the responses for FW, DW, and concentration of N and K, and the simple linear model fitted to the responses for P and Ca concentration for nontreated and ethephon-treated ‘Italian’ and ‘Brooks’ flower buds.

| Null hypothesis                      | FW F P value | FW P value | DW F P value | DW P value | N F P value | N P value | K F P value | K P value | P F P value | P F P value | Ca F P value | Ca P value |
|--------------------------------------|--------------|------------|--------------|------------|-------------|-----------|-------------|-----------|-------------|-------------|--------------|-------------|
| B, B + E, I, and I + E coincide     | 21.60 0.000  | 15.79 0.000 | 1.21 0.341   | 4.64 0.002  | 2.65 0.041  | 3.34 0.015 |
| B and B + E coincide                | 10.42 0.000  | 7.77 0.001  | --- ---      | --- ---     | --- ---     | --- ---     |
| B and I coincide                    | 25.42 0.000  | 22.49 0.000 | --- ---      | --- ---     | --- ---     | --- ---     |
| B and I + E coincide                | 60.33 0.000  | 41.04 0.000 | 4.14 0.020   | 2.42 0.110  | 9.01 0.001  |
| B + E and I coincide                | 3.60 0.031   | 4.65 0.013  | --- ---      | --- ---     | --- ---     | --- ---     |
| B + E and I + P coincide            | 20.89 0.000  | 14.85 0.000 | 9.92 0.000   | 5.02 0.015  | 3.61 0.043  |
| I and I + E coincide                | 9.09 0.001   | 4.01 0.022  | 1.16 0.348   | 6.70 0.005  | 4.58 0.021  |

Based on data obtained from three replicates of 100 terminal buds each.

'B = 'Brooks'; B + E = 'Brooks' + ethephon; I = 'Italian'; I + E = 'Italian' + ethephon.

For both genotypes, there were no significant differences between the ethephon treatment and the controls (Fig. 1). However, in comparisons between ethephon-treated ‘Italian’ and ‘Brooks’ flower buds, a significant increase in K concentration in the ‘Brooks’ flower buds was observed.

The tests of hypothesis and probability levels for P and Ca appear in Table 1. There was no effect of ethephon application for P responses ($R^2 = 0.884$) of ‘Brooks’ flower buds. Ethephon-treated ‘Italian’ buds had lower P concentrations than the control (Fig. 1). The Ca response ($R^2 = 0.965$) was similar to that for P. Ethephon treatment had no effect on ‘Brooks’. Calcium concentration was significantly higher in ‘Italian’ buds treated with ethephon than in nontreated buds (Fig. 1).

There was no difference in B content per bud between the ethephon treatment and the controls during the first three dates of sampling (Table 2). From 10 Mar., ‘Italian’ + ethephon had significantly higher B content than the control treatment. The B content of treated and nontreated ‘Brooks’ flower buds were
Table 2. The effect of fall-applied ethephon on B content (in milligrams) of ‘Brooks’ and ‘Italian’ prune flower buds sampled from 50 days before bloom until just before full bloom.

| Sample date | 17 Feb. | 24 Feb. | 3 Mar. | 10 Mar. | 18 Mar. | 25 Mar. | 31 Mar. | 7 Apr. |
|-------------|---------|---------|--------|---------|---------|---------|---------|--------|
| P           | 11.13   | 12.17   | 14.23  | 12.60   | 15.20   | 14.87   | 20.10   | 45.66  |
| I + E       | 12.43   | 13.83   | 15.13  | 17.53   | 20.40   | 21.60   | 27.07   | 48.60  |
| B           | 10.20   | 10.37   | 12.00  | 16.47   | 15.40   | 15.10   | 31.13   | 54.43  |
| B + E       | 8.76    | 9.87    | 11.03  | 13.13   | 16.50   | 16.10   | 30.77   | 64.00  |

Significance

Contrasts

| I vs. I + E | NS | NS | NS | ** | * | ** | ** | ** |
| B vs. B + E | NS | NS | NS | ** | * | *** | * | * |
| I + (I + E) | NS | NS | NS | * | NS | NS | NS | * |
| vs. B + (B + E) | NS | NS | NS | NS | NS | NS | NS | ** |

‘Average B content of three samples (100 buds each) from 10 trees.

B = ‘Brooks’; B + E = ‘Brooks’ + ethephon; I = ‘Italian’; I + E = ‘Italian’ + ethephon.

NS, *, **, ***Nonsignificant or significant at $P = 0.05$, 0.01, or 0.001, respectively.

Ovule longevity

Field experiment. In 1989, senescence clearly progressed differently in the two cultivars in the field. During the first 8 days of study, nontreated and ethephon-treated ‘Italian’ flowers had fewer viable ovules than flowers of treated or nontreated ‘Brooks’ (Table 3). After this date, this difference became nonsignificant until the last two sampling dates. The rate of ovule death of ‘Brooks’ flowers was generally the same for the ethephon treatment and the control. There was no difference in any of the sampling dates, except 8 DAFB (Table 3). For ‘Italian’, flowers started with fewer viable ovules until 8 DAFB, when there was no difference between the ethephon-treated and control flowers. After that date, ethephon-treated flowers showed significantly more viable ovules than nontreated ones.

Table 3. Number of viable ovules in ‘Brooks’ and ‘Italian’ prune flowers treated with or without ethephon from full bloom until 20 DAFB under field conditions in 1989.

| Treatment | 0 | 4 | 6 | 8 | 11 | 13 | 17 | 20 |
|-----------|---|---|---|---|----|----|----|----|
| P         | 1.8| 1.5| 1.4| 1.0| 0.8| 0.8| 1.0| 0.6| 0.4|
| I + E     | 1.2| 1.2| 1.1| 1.1| 1.1| 1.0| 0.9| 1.0| 0.8|
| I         | 2.0| 1.8| 1.8| 1.7| 1.2| 1.1| 1.2| 1.1| 1.0|
| B + E     | 1.8| 1.8| 1.8| 1.5| 1.2| 1.2| 1.1| 1.1| 1.2|

Significance

Contrasts

| I vs. I + E | NS | NS | NS | ** | * | NS | NS | ** |
| I + (I + E) | NS | NS | NS | * | NS | NS | NS | NS |
| vs. B + (B + E) | NS | NS | NS | NS | NS | NS | NS | ** |

‘Average of 10 replicate flowers from 10 trees.

B = ‘Brooks’; B + E = ‘Brooks’ + ethephon; I = ‘Italian’; I + E = ‘Italian’ + ethephon.

NS, *, **, ***Nonsignificant or significant at $P = 0.05$, 0.01, or 0.001, respectively.

only different at two sampling dates, 10 Mar. and 7 Apr. When comparing the overall B content for each cultivar, it was higher in ‘Brooks’ only at the last two sampling dates.

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Days after full bloom

Fig. 2. Frequency distribution of the number of flowers with two viable ovules from full bloom until 20 DAFB excised from nontreated and ethephon-treated ‘Italian’ and ‘Brooks’ prune trees.

Growth chamber experiment. The predicted number of viable ovules in ‘Italian’ flower buds at 0 DAFB was fewer than two, indicating that ovular senescence had already occurred before bloom. This was not so for ‘Brooks’, where two viable ovules were present at full bloom in all flowers (Table 4, Fig. 3).

Across all temperature regimes, ‘Italian’ had fewer viable ovules than ‘Brooks’. At 5°C, the rate of ovule death was similar for both cultivars. The difference between the predicted average number of ovules across days is explained by the initial lower ovule viability of ‘Italian’. As temperature increased, ovule senescence was faster in ‘Italian’ than in ‘Brooks’. At 15°C, only one ovule per flower remained viable by 8 DAFB for ‘Italian’. For ‘Brooks’, higher temperatures also resulted in a more rapid decrease in ovule longevity over time, but at a lower rate than for ‘Italian’. Total ovule senescence for some ‘Italian’ flowers had already occurred by 2 DAFB at 20°C. For ‘Brooks’ flowers at the same temperature, senescence of both ovules began 4 days later (6 DAFB).

For ‘Brooks’, flowers from treated and nontreated spurs had the same number of viable ovules at anthesis (Fig. 3). The surface response for the ethephon-treated buds was different than that of the nontreated control buds. At low temperatures, the rate of ovule senescence was similar in both cases and a clear difference only started to appear between 15 and 20°C. At 20°C, ovule longevity was slightly enhanced by ethephon treatment. By 18 DAFB, the ethephon-treated flowers had more viable ovules than the control.

Full-applied ethephon reduced the rate of ovule death in both cultivars (Fig. 3). For ‘Italian’, ethephon application translated into fewer viable ovules at anthesis. However, those ovules remained viable for a longer period. Ovule longevity was enhanced, but this effect also depended on temperature. This interaction was particularly important at 20°C. Without ethephon, some flowers were already lacking viable ovules at 2 DAFB. However, when treated with ethephon, at least one viable ovule was present until 10 DAFB. Also, at 15°C, more ovules per flower remained viable in the ethephon-treated flowers. At 5°C, there appeared to be little difference between the ethephon treatment and the control.

Discussion

Flower buds treated with ethephon grew more slowly than the controls during the winter. This difference was maintained during the period of flower bud swell in the spring. There was a considerable difference in terms of FW and DW. According to Gianfagna (1989), pistil length was only 80%, and the length of the flower buds was 90% of nontreated ‘Cresthaven’ peach buds within 1 month of the ethephon application. In our case, buds from the ethephon-treated trees were 20% smaller than nontreated buds. This difference might be explained by reduced cell division in the ethephon-treated buds (Gianfagna, 1989).

Apelbaum and Burg (1972) have shown that ethylene inhibits cell division in the plumular hook of pea. Inhibition of the growth of dormant buds by ethephon suggests that ethylene released in the tissues may inhibit cell division during the later phases of flower differentiation in the autumn (Crisosto et al.,

Table 4. Regression equations, coefficients of determination ($R^2$), and model significance probabilities ($P > F$) of the number of viable ovules for nontreated and ethephon-treated ‘Italian’ and ‘Brooks’ prune flower buds held at 5, 10, 15, and 20°C.

| Treatment | Equation | $R^2$ | $P > F$ |
|-----------|----------|-------|---------|
| I         | $Y = 1.2872 + 0.1233X_1 + 0.0831X_2 - 0.0053X_2^2 - 0.0029X_1^2 - 0.0143X_1X_2 + 0.00002315X_1X_2$ | 0.51 | 0.0001 |
| I + E     | $Y = 1.3450 + 0.0471X_1 - 0.0024X_2^2 - 0.0034X_2X_2$ | 0.31 | 0.0001 |
| B         | $Y = 1.5679 + 0.0924X_1 + 0.0635X_2 - 0.0039X_2^2 - 0.0015X_2^2 - 0.0105X_1X_2 + 0.00001546X_1X_2$ | 0.41 | 0.0001 |
| B + E     | $Y = 1.9345 - 0.0038X_1 - 0.0011X_2 - 0.0034X_2X_2$ | 0.25 | 0.0001 |

*B = ‘Brooks’; B + E = ‘Brooks’ + ethephon; I = ‘Italian’; I + E = ‘Italian’ + ethephon; $X_i =$ temperature (°C); $X_2 =$ DAFB.

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Delayed differentiation of flower primordia could result in a delay in flower development the following spring. As the process of flower bud development is a complex phenomenon under both genetic and environmental control, it may be directly and indirectly related to a number of organic and inorganic nutritional factors. However, the literature does not provide much evidence for categorizing the essential elements in this regard. In addition, there is no report in the literature regarding the effect of ethephon application on mineral element content of flower buds. Our results show that N and K concentrations in flower bud tissue were not changed by ethephon application. ‘Jonathan’ apple trees deficient in P show delayed bud burst and retarded development of fewer vegetative and floral meristems (Taylor and Goubran, 1975). Our results agree with this report to the extent that the ‘Italian’ flower buds had a lower P concentration with the ethephon application and bloomed later.

The higher concentration of Ca as a result of the ethephon application could be associated with delayed growth rather than increased movement of Ca$^{2+}$ into the buds. The vascular bundles of the ethephon-treated prune buds developed more slowly than those of nontreated buds (Sun et al., 1990). Calcium has relatively low mobility in the plant and is translocated mainly acropetally via xylem sap with the transpiration stream (Mengel and Kirby, 1982). As was the case for ethephon-treated buds, where growth was delayed and transpiration rates were most likely low, it is improbable that additional Ca was moved into buds. What we observed was most likely a dilution effect in the case of the nontreated control as growth started early and before the spring transport of Ca into the buds.

Boron levels also were higher in ethephon-treated buds of ‘Italian’ than in the control. Dilution cannot account for the lower levels of B in the nontreated controls, as B was expressed as total content per 100 buds. Hanson (1984) stated that re-mobilization of B from ‘Italian’ prune branches was more limited than that of other elements. In intact trees, B accumulated in buds more slowly before bud swelling. However, B accumulates rapidly as buds accumulate dry matter from swelling to bloom. Hanson and Breen (1985b) explained these results by suggesting that xylem in the axis of flower buds became functional only when buds began to swell. However, the putative xylem-supplied B accounted for only 26% of that B entering buds. In the same report, they reasoned that since buds showed limited dry matter accumulation at this time, they were weak carbon sinks and suggested that symplastic flow, or xylem transport, to buds would also be limited. Ethephon may have altered the intrinsic physiology of the buds and allowed enhanced B movement to buds in the spring.

Longer ovule viability of ‘Brooks’ flowers, compared to ‘Italian’, was observed under both field and growth chamber conditions. Ethephon application did not change this difference markedly. ‘Brooks’ flowers seem to fit Williams’ (1965) description of a “strong” flower; these flowers were larger and bloomed earlier. Ten percent of ‘Brooks’ flowers were “strong” enough to maintain two viable ovules until 20 DAFB. At this time, senescence of both ovules was observed in only 10% of the flowers; the remaining 90% still had one viable ovule. ‘Italian’ flowers can be classified as “weak” blossoms according to the criteria outlined by Williams (1965). Some ovule senescence was already apparent at the time of bloom. The ovules

Fig. 3. Predicted number of viable ovules derived from equations in Table 4 for nontreated and fall ethephon-treated (500 mg·liter$^{-1}$) flowers excised from ‘Italian’ and ‘Brooks’ prune trees and held for 18 DAFB at 5, 10, 15, and 20C.
of ‘Italian’ flowers also senesced at a faster rate than those of ‘Brooks’. Ethephon-treated flowers at bloom had fewer viable ovules, but they showed a delayed senescence and, in practical terms, might have improved the chance of fertilization.

Efforts to prolong ovule longevity have been successful in apple by improving cultural conditions (Dorsey, 1929; Williams, 1963, 1965). Williams (1965) found that summer N, application provided optimal nutritional conditions at the time of flower initiation, which greatly increased the proportion of strong flowers. Kliewer (1977) stated that grape (Vitis vinifera L.) ovule fertility may be markedly affected by a supply of organic nutrients to the ovules. Perhaps, the lower fruit set observed in the ‘Italian’ genotype can be explained by a limited supply of nutrients to the flower buds (Thompson and Liu, 1973; Hanson and Breen, 1985a). If one of the ovules is senescent, as when ethephon-treated, the other ovule may compete better for a “limited” supply of nutrients to the flower buds. In addition, increased amounts of B and Ca in the ethephon-treated buds may enhance ovule longevity.

The results clearly illustrated that ovule senescence was influenced by temperature. Ovule senescence was very rapid at the higher temperatures we used. However, genotype and ethephon treatment interacted with temperature. Fruit set in grape was much better at 25°C than at 35 or 40°C (Ewart and Kliewer, 1977; Kliewer, 1977). They indicated that ovules degenerated earlier as temperature increased, relating this to an insufficient supply of organic nutrients to the ovules. Tomato (Lycopersicon esculentum Mill.) ovules were affected by high temperatures more drastically than was pollen (El-Ahmadiand Stevens, 1979). Ovule viability was reduced in all cases by high temperatures, but cultivars differed significantly in this respect. Levy at al. (1978) found that tomato male gamete viability was affected more at high temperature than that of female gametes. However, the degree of the temperature effect between the male or female gamete was found to be genotype-dependent. In our experiments, ovule longevity of ‘Brooks’ seemed to be less dependent on temperature than that of ‘Italian’. This independence may be associated with the capacity of ‘Brooks’ to consistently set a crop. The effect of ethephon, in preventing ovule senescence in ‘Brooks’ flowers was less marked and occurred only at high temperatures. Perhaps, as a result of the tendency of ‘Brooks’ to produce strong ovules, these ovules have a higher temperature stress threshold than those of ‘Italian’.

Flower buds from ‘Italian’ trees were smaller and had a lower concentration and content of Ca and B, respectively, when compared to ‘Brooks’ flower buds. Ovule longevity was shorter in ‘Italian’ flowers. Since the ‘Italian’ genotype apparently has a “weaker” flower genotype when compared to ‘Brooks’, cultural practices that ameliorate this weaker condition have the potential to enhance ovule longevity. As ‘Brooks’ flowers already have the tendency to produce strong flowers, increasing ovule longevity may not be possible.

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