Induction and Characterization of Fruit Abscission during Early Physiological Fruit Drop in Citrus

Xi Li1, Akira Kitajima2*, Tsuyoshi Habu2**, Keiko Kataoka2**, Rihito Takisawa2 and Tetsuya Nakazaki2

1Graduate School of Agriculture, Kyoto University, Kyoto 606-8502, Japan
2Experimental Farm, Graduate School of Agriculture, Kyoto University, Kizugawa 619-0218, Japan

A new method of inducing fruit abscission by incubating detached ovaries and fruits in agar medium was developed in citrus. Ovary and fruit abscission in the satsuma mandarin, ‘Kiyomi’, hyuganatsu, and ponkan during early physiological fruit drop was characterized using this method. For primary physiological fruit drop, the abscission of detached ovaries could be divided into three patterns, including an early type in hyuganatsu, a medium type in ‘Kiyomi’, and a late type in satsuma mandarin and ponkan. The cumulative abscission ratio of four species and cultivars was over 80% at 96 h after treatment. However, for secondary physiological fruit drop, the initiation of fruit abscission was earliest in hyuganatsu, and latest in satsuma mandarin. The cumulative abscission ratio was highest in ponkan at 84% and lowest in hyuganatsu at 6%. The pattern of abscission was different for primary and secondary physiological fruit drop in the four species and cultivars. High temperature promoted the abscission of detached ovaries and fruits in satsuma mandarin and ponkan. Leaf attachment suppressed fruit abscission in ‘Kiyomi’, hyuganatsu, and ponkan at 7 weeks after anthesis in 2012 and ponkan at 5 weeks after anthesis in 2013. Ovary and fruit abscission in four species and cultivars during early physiological fruit drop was characterized by incubating detached ovaries and fruits in agar medium.

Key Words: agar medium, fruit abscission induction, physiological fruit drop.

Introduction

In citrus, physiological fruit drop is a major problem in commercial fruit production. In general, physiological fruit drop including primary physiological fruit drop and secondary physiological fruit drop affects ovary and young fruit abscission. During primary fruit drop, ovary abscission occurs between peduncle and pedicel. During secondary fruit drop, young fruit drop occurs between fruit and peduncle. Many studies have previously been conducted to elucidate the mechanism underlying physiological fruit drop.

Current evidence indicates that carbohydrate availability when growing young fruits is strongly associated with fruit abscission during physiological fruit drop in citrus (Iglesias et al., 2003). This suggestion has also been advanced for other tree species, such as the persimmon (Kitajima et al., 1990, 1992), peach (Morandi et al., 2008), and grape (Aziz, 2003). Defoliation reduces the number of leaves (Mehouachi et al., 1995) and shading induces a reduction in photosynthetic activity (Byers et al., 1990), resulting in increased fruit abscission. Girdling treatment reduces fruit abscission by blocking photoassimilate transport from leaves to roots, thus increasing carbohydrate availability for the growing fruits (Yamanishi, 1995). From these results it was concluded that fruit set is highly dependent on carbohydrate availability (Iglesias et al., 2003). All these treatments have been applied to trees. For fruits on trees, the amount of photoassimilate products translocated into sink organs regulates fruit abscission. Fruit abscission is the result of

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* Corresponding author (E-mail: kitajima@kais.kyoto-u.ac.jp).
** Present address; Faculty of Agriculture, Ehime University, Matsuyama 790-8566, Japan.

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competition for carbohydrate between fruits or between growing shoots and fruit. Environmental conditions, such as high temperature, rain, and insufficient light, also increase fruit abscission during physiological fruit drop (Kajiura, 1942; Sato et al., 2010). Regarding the influence of many variables on trees, it is difficult to elucidate the mechanism of fruit abscission during physiological fruit drop.

As a vital hormone regulating plant abscission, ethylene has been investigated for decades. Many studies suggest that ethylene is closely associated with fruit abscission in tree species (Gómez-Cadenas et al., 2000). The plant growth regulators benzylaminopurine (BAP) and aminooxyacetic acid (AOA) have been applied to investigate possible regulation of ethylene biosynthesis and its role in fruit abscission (Dal Cin et al., 2005; Tudela and Primo-Millo, 1992). However, whether the mechanism of fruit abscission induced by ethylene treatment is similar to that in trees during physiological fruit drop or not is not well understood.

To investigate the mechanism of fruit abscission, a new method for induction of fruit abscission could be developed by blocking carbohydrate translocation to the fruit. In the present study, the utility of this method was tested. Using this method, an investigation was undertaken to 1) characterize ovary and fruit abscission in several species and cultivars during primary and secondary physiological fruit drop and 2) determine the effect of temperature and sucrose addition on ovary and fruit abscission.

**Materials and Methods**

**Plant materials**

‘Miyagawa-wase’ satsuma mandarin (*Citrus unshiu* Marcow.), ‘Soren tangelo’ (C. sp.), ‘Kiyomi’ (*C. unshiu* × *C. sinensis* (L.) Osbeck), ‘Futsukei’ hyuganatsu (*C. tamurana* hort. ex Tanaka), and ‘Teishoukei’ ponkan (*C. reticulata* Blanco) were used in these experiments.

**Induction of abscission in detached and attached fruit**

Satsuma mandarin and ‘Soren tangelo’ were used in an experiment performed 6 weeks after anthesis (WAA) in 2011. The trees were grown on the experimental farm of Kyoto University, Osaka, Japan. The experiment was performed with 100 open-pollinated fruits from leafy inflorescence on each cultivar. Of these, 50 fruits were detached from the tree as the “attached fruit” treatment and the remaining 50 were left on the tree as the “attached fruit” treatment. Peduncles of detached fruits were cut 2 cm from the sepal and were incubated in 1% agar medium at 25°C.

In 2011, during anthesis, detached ovaries of satsuma mandarin, hyuganatsu, and ponkan were used. The ovaries were incubated in 1% agar medium with 0% sucrose at 15°C and 25°C or in 1% agar with 1% and 3% sucrose under 25°C. Numbers of abscised ovaries were recorded every 6 h after the induction of ovary abscission. The percentage of cumulative ovary abscission was calculated. Fifty detached ovaries were used for each treatment.

**Abscission of detached ovaries under different conditions during primary physiological fruit drop**

Satsuma mandarin, ‘Kiyomi’, hyuganatsu, and ponkan were used at 4 WAA in 2012. Trees were grown on the experimental farm of Kinki University. Fifty open-pollinated fruits on leafy shoots were detached as described above, and incubated in 1% agar medium under 25°C.

In 2011, during anthesis, detached ovaries of satsuma mandarin, hyuganatsu, and ponkan were used. The ovaries were incubated in 1% agar medium with 0% sucrose at 15°C and 25°C or in 1% agar with 1% and 3% sucrose under 25°C. Numbers of abscised ovaries were recorded every 6 h after the induction of ovary abscission. The percentage of cumulative ovary abscission was calculated. Fifty detached ovaries were used for each treatment.

**Abscission of detached fruit under different conditions during secondary physiological fruit drop**

Satsuma mandarin, ‘Kiyomi’, hyuganatsu, and ponkan were used at 4 WAA in 2012. Trees were grown on the experimental farm of Kinki University. Fifty open-pollinated fruits on leafy shoots were detached as described above, and incubated in 1% agar medium under 25°C.

Satsuma mandarin, hyuganatsu, and ponkan were used at 5 WAA in 2012. Trees were grown on the experimental farm of Kinki University. One hundred and fifty open-pollinated fruits on leafy shoot were detached as described above and incubated in 1% agar medium under 20, 25, and 30°C.

Satsuma mandarin, hyuganatsu, ponkan, and ‘Kiyomi’ fruits at 7 WAA in 2012 and hyuganatsu and ponkan fruits at 5 WAA in 2013 were used. Trees were grown on the experimental farm of Kinki University. For each species and cultivar, 100 open-pollinated fruits were detached from the peduncle with leaves. Fifty fruits of each treatment with two leaves (leaf attachment) and the fruits with no leaf (leaf detachment) were
incubated in 1% agar medium at 25°C with a 24 h light period under about 10000 lux fluorescent lighting.

Abscised young fruits were counted every 6 h after induction of fruit abscission and the percentage of cumulative fruit abscission was calculated.

**Results**

**Induction of detached and attached fruit abscission**

Figure 1 shows the abscission patterns of detached and attached satsuma mandarin and ‘Soren tangelo’ fruits during 186 h after induction treatment for fruit abscission at 6 WAA in 2011.

For attached fruits, the abscission of ‘Soren tangelo’ began at 36 h after induction (HAI) with a 2.9% cumulative abscission ratio. The cumulative abscission ratio then increased and reached 72.9% at 186 HAI. For attached satsuma mandarin, abscission began at 60 HAI with 0.7% and the cumulative abscission ratio at 186 HAI reached 21.3%. Initiation of attached fruit abscission in ‘Soren tangelo’ was earlier and the final abscission ratio was higher than that in satsuma mandarin. The fruit abscission pattern of attached ‘Soren tangelo’ was different from that of satsuma mandarin. For detached fruits, abscission of ‘Soren tangelo’ began at 60 HAI and the cumulative abscission ratio increased from 2.9% to 95.7% during 186 HAI. For satsuma mandarin, the cumulative abscission ratio was only 6.7% at 186 HAI. The fruit abscission pattern of detached ‘Soren tangelo’ was different from that of satsuma mandarin. The abscission pattern of detached fruits was similar to that of attached fruits for ‘Soren tangelo’ and satsuma mandarin.

Figure 2 shows the abscission patterns of attached and detached fruits in satsuma mandarin, ‘Kiyomi’, and hyuganatsu during 156 HAI at 6 WAA in 2013.

For detached fruits, the abscission of hyuganatsu began at 72 HAI and the cumulative abscission ratio increased from 10% to 56% during 156 HAI. The abscission ratio was higher than that of satsuma mandarin and ‘Kiyomi’. For satsuma mandarin and ‘Kiyomi’, fruit abscission began at 108 HAI and 96 HAI, respectively, and the cumulative abscission ratios in the two species at 156 HAI were 28% and 16%, respectively. The fruit abscission pattern was similar for satsuma mandarin and ‘Kiyomi’, and different for hyuganatsu. The abscission pattern of detached fruits was similar to that of attached fruits for all species and cultivars.
Abscission of detached ovary under different conditions during primary physiological fruit drop

Figure 3 shows the cumulative abscission of detached ovaries in satsuma mandarin, ‘Kiyomi’, hyuganatsu, and ponkan during 96 HAI at anthesis in 2012. The ovary abscission pattern could be divided into the following three types: (1) an early type in hyuganatsu, with abscission beginning at 12 HAI; (2) a medium type in ‘Kiyomi’, with abscission beginning slightly later than that in hyuganatsu; (3) a late type in satsuma mandarin and ponkan. The initiation of ovary abscission in both species was at 48 HAI. The cumulative abscission ratio of the four species and cultivars at 96 HAI was a little over 80%.

Figure 4 shows the influence of temperature on detached ovary abscission in satsuma mandarin, hyuganatsu, and ponkan at anthesis in 2011. Detached ovaries were incubated in 1% agar with 0% sucrose at 15°C and 25°C. Higher temperature promoted the cumulative abscission ratio during 72 HAI. In hyuganatsu, the cumulative abscission ratio at 72 HAI was 23.3% at 15°C, but reached 100% at 25°C. In satsuma mandarin and ponkan, the cumulative abscission ratios at 72 HAI reached 95.7% and 85%, respectively, at 25°C, whereas ovary abscission did not occur below 15°C.

Figure 5 shows the influence of 1% and 3% sucrose added to 1% agar medium on detached ovary abscission in satsuma mandarin, hyuganatsu, and ponkan at anthesis in 2011. In hyuganatsu and ponkan, abscission began at 18 HAI and 54 HAI and the cumulative abscission ratio at 72 HAI was over 90% under 1% and 3% sucrose treatment, similar to that under 0% sucrose treatment. Sucrose addition did not affect detached ovary abscission of hyuganatsu or ponkan during 72 HAI. However, in satsuma mandarin, 1% sucrose addition suppressed ovary abscission compared to 0% and 3% sucrose addition treatment.

Abscission of detached fruit under different conditions during secondary physiological fruit drop

Figure 6 shows the cumulative abscission ratio of detached fruits in satsuma mandarin, ‘Kiyomi’, hyuganatsu, and ponkan during 96 HAI at 25°C during secondary physiological fruit drop at 4 WAA in 2012.
The fruit abscission of ponkan was initiated at 48 HAI and the cumulative abscission ratio then reached 84% at 96 HAI. The fruit abscission of ‘Kiyomi’ began at 48 HAI and the cumulative abscission ratio was 48% at 96 HAI. The abscission of fruit of hyuganatsu and satsuma mandarin began at 18 HAI and 90 HAI, respectively, and the cumulative abscission ratio at 96 HAI was approximately 20% in both species. The cumulative abscission ratio was highest in ponkan and lowest in hyuganatsu.

Figure 7 shows the effect of different temperatures on detached fruit abscission in satsuma mandarin, hyuganatsu, and ponkan at 5 WAA in 2012. In ponkan, under the treatments at 25 and 30°C, abscission began at 36 and 48 h, respectively, and the cumulative abscission ratios at 96 HAI were 76% and 84%, respectively. Abscission under the 20°C treatment began at 60 HAI, and the final abscission ratio at 96 HAI was 40%. In hyuganatsu, fruit abscission was low under each temperature treatment. The cumulative abscission ratio under 30°C treatment at 96 HAI was only 20%. In satsuma mandarin, fruit abscission was induced under the 30°C treatment, and the cumulative abscission ratio at 96 HAI was 76%, whereas that under the 20°C and 25°C treatments was only 12% at 96 HAI.

Figure 8 shows the influence of detaching fruits with leaves on fruit abscission in satsuma mandarin, ‘Kiyomi’, hyuganatsu, and ponkan at 7 WAA in 2012, and Figure 9 shows this influence in hyuganatsu and ponkan at 5 WAA in 2013.

At 7 WAA in 2012, in satsuma mandarin, the cumulative abscission ratios of leaf attachment and detachment were 0% and 4%, respectively, at 96 HAI. In ‘Kiyomi’, the initiation of fruit abscission in leaf attachment and detachment was at 72 and 66 HAI, respectively, and the cumulative abscission ratios at leaf attachment and detachment treatment were 32% and 74%, respectively, at 96 HAI. Leaf attachment of hyuganatsu and ponkan delayed initiation of abscission, and the cumulative abscission ratios were 32% and 28%, respectively, at 96 HAI.

At 5 WAA in 2013, in ponkan, leaf attachment delayed initiation of fruit abscission, and the final cumulative abscission ratio at 96 HAI was 24.6%, in contrast to leaf detachment, which increased the cumulative abscission ratio to 64% at 96 HAI. However, in hyuganatsu, irrespective of leaf attachment, the abscission pattern was similar and the cumulative abscission ratio was below 20% during 96 HAI.

Discussion

The relationship between carbohydrate supplementation and hormonal regulation of citrus fruit abscission during physiological fruit drop has been previously studied (Iglesias et al., 2003, 2006; Roitsch, 1999). Many tree treatments, such as defoliation, shading, and girdling, have been applied to investigate the mechanism of fruit abscission. Defoliation (Mehouachi et al., 1995) and shading (Byers et al., 1990) decreased photosynthetic activity, whereas girdling blocked photosimulate transport from leaves to roots, thus increasing carbohydrate availability to the growing fruits (Yamanishi, 1995). Sucrose supplementation increased citrus fruit set (Iglesias et al., 2003). These results indicate that fruit set is highly dependent on carbohydrate availability. However, hormones also influence abscission. Exogenous ethylene or plant growth regulators have been applied to investigate fruit abscission in
various fruit trees (Pozo et al., 2004; Untiedt and Blanke, 2001). Carbon shortage induced fruit abscission via ethylene, which was considered as the final hormonal product to induce fruit drop during physiological fruit drop (Gómez-Cadenas et al., 2000). Although these treatments have been commonly applied for investigation of fruit abscission, it is not known whether the mechanism of natural fruit abscission is similar to that in ethylene-induced fruit abscission. Moreover, the occurrence of natural fruit abscission is not very clear under the fruit abscission induction treatments described above because of unstable environmental conditions, such as high temperature, rain, and insufficient light. In our study, a new method, namely incubation of detached fruits in agar medium to exclude variable influences on the tree during physiological fruit drop, was applied. In this experiment, to block carbohydrate translocation to the fruit, peduncle ringing was conducted on attached fruits and incubation in agar medium was conducted on detached fruits. The abscission pattern of detached fruit incubation in agar medium was similar to that of fruits on the tree in satsuma mandarin and ‘Soren tangelo’ at 6 WAA in 2011 (Fig. 1) and in satsuma mandarin, ‘Kiyomi’, and hyuganatsu at 6 WAA in 2013 (Fig. 2). These results indicate that a similar mechanism regulates abscission for both detached fruits in agar medium and attached fruits on the tree. Detached leaf explant incubation has also been performed to investigate the leaf abscission process in citrus (Agustí et al., 2008). We conclude that incubation of detached fruits in agar medium is a useful method for inducing fruit abscission.

In the primary physiological fruit drop, the final 96 HAI cumulative abscission ratio of detached ovaries in four species and cultivars uniformly exceeded 80% (Fig. 3), although the time of abscission initiation was different for each species and cultivar. However, in the secondary physiological fruit drop, the final 96 HAI cumulative abscission ratio of satsuma mandarin, ‘Kiyomi’, and hyuganatsu was under 50%, whereas that of ponkan reached 84% (Fig. 6). Detached ovaries reached a higher abscission ratio during primary physiological fruit drop, whereas detached fruits did not reach abscission ratios as high as detached ovaries during secondary physiological fruit drop. This result indicates that different mechanisms trigger ovary and fruit abscission in the primary and secondary physiological fruit drop stages. In citrus, physiological fruit drop was activated through two abscission zones located between the peduncle and pedicel during primary fruit abscission and between the peduncle and fruit during secondary fruit abscission (Iglesias et al., 2007). 1-aminocyclopropane-1-carboxylic acid (ACC) treatment was unable to reactivate the abscission zone between
peduncle and pedicle or peduncle and fruit after the period of physiological fruit drop (Iglesias et al., 2006), indicating that the mechanisms regulating abscission in the two physiological fruit drops are different. In citrus, the involvement of separation layers in young fruit abscission has rarely been investigated. Further studies are needed to clarify the differences in formation of two abscission zones in citrus.

Plant hormones are among the internal signals promoting abscission. A high level of endogenous gibberellins (GAs) resulted in high fruit set and elevated parthenocarpic ability in satsuma mandarin (Talón et al., 1992). Satsuma mandarin showed the lowest cumulative abscission ratio during 96 HAI among four species and cultivars in secondary physiological fruit drop (Fig. 6). This observation probably results from reduction by GAs in satsuma mandarin fruit abscission. In citrus, carbohydrate shortage resulted in young fruit abscission by triggering increased levels of ACC (Gómez-Cadenas et al., 2000). The fruit abscission mechanism may depend on the ability of the abscission zone to sense ethylene (Bonghi et al., 2000). Further research on the expression of ethylene biosynthesis and ethylene receptor genes may identify their relationships with fruit abscission in species and cultivars.

Citrus physiological fruit drop has been proposed to depend on temperature. Inoue and Chien (1987) reported that satsuma mandarin grown in sunlit phytotron rooms showed severe fruit abscission under high temperature. Sato et al. (2010) also reported that physiological fruit drop in citrus was promoted by a 2°C increase in air temperature after full bloom. These results suggest a close relationship between temperature and fruit abscission. In our experiment, the cumulative abscission ratio of detached ovaries in satsuma mandarin, hyuganatsu, and ponkan was extremely high at 25°C compared with that at 15°C (Fig. 4), indicating a result similar to previous observations, that is, there is a clear relationship between ovary abscission and temperature in each species and cultivar. In contrast, detached fruit abscission at 30°C in hyuganatsu was only slightly induced, whereas that in satsuma mandarin and ponkan was promoted (Fig. 7). However, 25°C and 15°C treatment did not promote fruit abscission in satsuma mandarin and hyuganatsu. These results suggest that the response of the abscission mechanism to temperature is different in ovaries and fruits or among species and cultivars. It appears that the ovary is much more sensitive to temperature compared with fruit and that each species and cultivar has a different mechanism for responding to the same temperature during fruit abscission.

We investigated the effect of sucrose supplementation on induction of abscission of detached ovaries and fruits. Addition of 1% and 3% sucrose in agar medium showed no effect on suppression of ovary abscission in satsuma mandarin, hyuganatsu, or ponkan, except for 1% sucrose addition for satsuma mandarin (Fig. 5). It appears that ovary abscission may be not affected by carbohydrate supplementation during flowering. Therefore, more investigations are needed to clarify factors affecting induction of ovary abscission during primary physiological fruit drop.

Partial defoliation promoted young fruit abscission and carbohydrate availability was important for fruit abscission (Iglesias et al., 2003). Fruit abscission was suppressed by leaf attachment in hyuganatsu and ‘Kiyomi’ at 7 WAA, and ponkan at 5 WAA (Figs. 8 and 9), which probably resulted from carbohydrate supplementation from leaves. However, the effects of leaf attachment on suppression of fruit abscission were different in cultivars and times. The signal of fruit abscission induction that related to reduction of photosynthesis transport or changes in plant hormones from leaves may be different in species and cultivars at 7 WAA and 5 WAA. We considered that the difference in the fruit abscission pattern of leaf attachment and detachment in hyuganatsu and ponkan at 7 WAA and 5 WAA could be useful for gene expression analysis to identify fruit abscission induction signals.

Environmental conditions such as high temperature, rain, and insufficient light increased fruit abscission during physiological fruit drop (Kajiura, 1942; Sato...
et al., 2010). However, besides environment conditions, fruit abscission is also influenced by genetic factors. For example, endogenous levels of auxins and GAs are higher in the ovaries of parthenocarpic than in normal tomato lines (Gillaspy et al., 1993). Satsuma mandarin has a high degree of natural parthenocarpy and a high fruit set (Talón et al., 1992). Characterization of fruit abscission in each species and cultivar was difficult for attached fruits in the field. In the present study, the abscission pattern of ovaries and fruits in four species and cultivars during physiological fruit drop was determined using incubation of detached ovaries and fruits in agar medium. Therefore, we propose that detached fruit incubation allows characterization of fruit abscission potential in each species and cultivar during physiological fruit drop.

In this study, abscission of ovaries and fruits in four species and cultivars was characterized by incubation of detached ovaries and fruits in agar medium, and the pattern of abscission differed between primary and secondary physiological fruit drop. Temperature was closely associated with abscission of detached ovaries and fruits. Leaf attachment suppressed abscission of detached fruits.

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Literature Cited


