Flowering Induction in Camellia chrysantha, a Golden Camellia Species, with Paclobutrazol and Urea

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Additional index words. early flowering, exogenous phytohormone, golden camellia flower, reproduction

Abstract. The flowers of Camellia chrysantha, commonly named as golden camellia, are treasured for their unique yellow color and are popularly used for tea. Compared with common camellia flowers that are either red, purple, pink, or white, golden camellia flowers are rare and are in high market demand. Our study was aimed to induce flowering in juvenile C. chrysantha grafted plants with urea and paclobutrazol (PBZ), a growth retardant. Generally, it takes 6–8 years for C. chrysantha seedlings and 5–6 years for grafted plants to set flower buds. With a 4 × 4 factorial design, four dosages of urea (1, 3, 5, or 8 g/plant) and four concentrations of PBZ (50, 150, 350, and 750 ppm) were tested on 4-year-old C. chrysantha grafted plants. Significant interaction between urea and PBZ was observed, and nine of the 16 combinations produced significantly more flower buds than the control, although not all flower buds could open because of abscission. High concentrations of PBZ and high dosages of urea were generally associated with severe defoliation and slow growth of basal stem diameter. When taking bud abscission into account, combinations of 150 ppm PBZ with 1 g urea and 350 ppm PBZ with 3 g urea resulted in significant flowering in juvenile C. chrysantha grafted plants without negative effects on vegetative growth and flower bud size and severe defoliation. This is the first report on flowering induction in a golden camellia species using juvenile plants. Our results suggest that application of optimized PBZ and urea doses can be a potential means for manipulation of early flowering in golden camellia species.

Commonly referred to as cha-hua in Chinese, camellia flowers have been cultivated and treasured in the Orient for thousands of years and are grown throughout the world in botanical gardens and home greenhouses (Ming, 2000; Zhu et al., 2007). Their flowers bloom from winter to spring, in a range of colors, forms, fragrances, and sizes. Until fairly recently, camellia flower coloration was restricted to red, purple, pink, and white. Yellow is a long-sought-after color. Traditional breeding approaches were tried for nearly half a century, yet were unsuccessful in producing a yellow flower (Park, 2000; Shinichi et al., 2004). Therefore, when a camellia with deeply golden-yellow petals was first discovered in Guangxi, China, by Hu (1965), it generated great excitement in the horticultural sector of the world. Hu (1965) first named the species as Theopsis chrysantha. This taxon was subsequently transferred to the genus Camellia by Tuyama (1975) as Camellia chrysantha (Hu) Tuyama. Later taxonomic studies proved that C. chrysantha was the same as Camellia nitidissima Chi, which was introduced much earlier in the 1930s, yet received no attention from the public or horticulturists (Deng et al., 2000). Since then, both names have been used in publications.

Camellia chrysantha, along with other similar yellow-flowering camellia species that were discovered later, are grouped into section Chrysanthha Chang (Theaceae), which comprises more than 24 species and five variants and are commonly known as golden or yellow camellias (Zhuang, 2008). Because of their large unique golden flowers (5–10 cm in width) and high ornamental value, these yellow camellias are honored as ‘‘the Queen of Camellia’’ (Liang, 1993). Unlike other camellia species that are globally cultivated, this subgroup occurs in a narrow region of southwest China and North Vietnam, ranging between 20°32′–23°53′N and 104°–108°56′E and at altitudes of 50–650 m (Gao, 2005; Zhang and Ren, 1998). In recent years, there has been a dramatic decline in natural population size for these species because of a combination of factors such as increasing anthropogenic pressure, deforestation, and destructive collection of seedlings (Xing, 2005). As a result, yellow camellias have also unfortunately earned the nickname ‘‘the Giant Panda of the Plant Kingdom’’ because they have been listed as a first-grade endangered species in China (Xu, 1995). In addition to its importance in floriculture, golden camellias are priced for their beneficial components in leaves and flowers. According to Song et al. (2011), golden camellias contain more diverse phenolic compounds compared with commonly consumed tea leaves (Camellia sinensis). Ellagitannins, proanthocyanidins, taxifolin deoxyhexose, apigenin derivatives, kaempferol derivative, quercetin derivatives, glucosylisorhamnetin, (epi)catechin-(epi)afzelechinpolymerisations, and phyllospine were found in golden camellias, whereas negligible in common tea leaves. Many of these compounds have been shown to possess antioxidant and anti-inflammatory activities that may be beneficial to health (Lin et al., 2013; Wang et al., 2015). Unlike common tea leaves, no caffeine was detected in the six species of golden camellias included in the study of Song et al. (2011). Currently, golden camellias have been introduced to Japan, Australia, and North America as a genetic resource for commercial cultivation (Li et al., 2013; Nybom and Bartish, 2000; Zhuang, 2008). Because flowers are usually harvested before seeds are set, golden camellia seeds are not readily available. As a result, grafting and cutting are two main propagation methods for golden camellia.

Golden camellia flower, bud, leaf, and seed cake are popularly used as tea in China. However, the price is high, particularly for flower tea. According to Su (2010) and a report in China Daily published in Sept. 2014 (http://guangxi.chinadaily.com.cn/hechi/2014-09/05/content_18551816.htm), the price of golden camellia flower tea is about 30,000 Yuan ($4600) per kilogram, whereas bud tea is about 20,000 Yuan (~$3050), seed cake tea 1050 Yuan ($165), and leaf tea about 300 (~$460) Yuan. Hence, there is a large market for golden camellia flowers. Whereas mature golden camellias can produce as many as 300 flowers per tree, it generally takes 6–8 years for seedlings and 5–6 years for grafted plants to start setting flower buds (Chai et al., 2009; Jiang and Zhao, 1997). This long juvenile phase not only hinders conventional breeding, but also has an adverse impact on economic incomes for breeders. Our study was aimed to induce flowering in juvenile C. chrysantha. We assessed the effects of different combinations of nitrogen fertilizer, urea [CO(NH2)2], and growth regulator, PBZ ([2RS,3RS]-1-(4-chlorophenyl)-4,4-dimethyl-2-[1H-1,2,4-triazol-1-yl]pentan-3-ol, PBZ),...
on flowering induction. The objective was to identify optimal combinations that could promote flowering in young *C. chrysanthana* plants while minimizing the undesirable effects.

Paclorobutrazol is a triazole-type cytochrome P-450 inhibitor used extensively in horticulture as a plant growth retardant and fungicide. Its dual biological functions are because of the facts that PBZ possesses effects. One isoform targets the ecdysis of insects and fungal sterols, whereas the other inhibits gibberellin (GA) biosynthesis by blocking the oxidative conversion of ent-kaurene to kaurenoic acid in plants, leading to decrease in endogenous GA levels and abscisic acid catabolism (Fletcher and Hofstra, 1988; Rademaker, 2000). PBZ has been found to be predominantly effective in inducing and manipulating flowering or fruiting in plants such as mango (Blakie et al., 2004; Kulkarni, 1888; Yadav et al., 2005), Eucalyptus (Griffin et al., 1993), *Consolida orientalis* (oriental knight’s-spur) (Mansuroglu et al., 2009), plum (Oliveira and Browning, 1993), red camellia hybrids (*Camellia × Williamsii*) (Wilkinson and Richards, 1988), and grapes (Christov et al., 1995). However, as an antagonist of GA, which promotes cell division and/or cell elongation in plants and is relatively abundant in juvenile plant tissues, PBZ reduces stem length (e.g., Mabngwwe et al., 2016; Pal et al., 2016). Adverse effects on fruit shape, leaf size and color, and timing of anthesis have also been reported (Stinchcombe et al., 1984; Williams, 1984; Zheng et al., 2012).

Nitrogen (N) is an essential nutrient for healthy plant growth and development, and urea is the predominant form of N fertilizer used worldwide. In addition to promoting vegetative growth and green coloration of foliage, nitrogen fertilizers can improve crop yield and quality. Elimination of N in cotton production could lead to an estimated yield reduction of 37% (Stewart, 2002). According to George and Nissen (1992), nitrogen alone increased the peach fruit set by 48% and tree fruit yields by about 40%, and when nitrogen and PBZ were applied together, the yield efficiency was estimated to increase by about 60%. Application of both nitrogen fertilizer and PBZ also substantially increased the occurrence of precociously flowering Eucalyptus trees over that of either treatment applied alone (Williams et al., 2003). Here, we report for the first time that application of optimized PBZ and urea doses can be a potential strategy to achieve early flowering in golden camellia species.

**Materials and Methods**

*Plant materials and study site.* Partially lignified *C. chrysanthana* branches (0.5- to 1-year-old) were grafted in Aug. 2012 on 1.5-year-old *Camellia osmantha* rootstocks. As a fast-growing oil camellia species, *C. osmantha* can tolerate drought, flooding, and high temperatures (Wang et al., 2014). Grafting compatibility of *C. chrysanthana* on *C. osmantha* is high, with a survival rate of at least 90% (Ma et al., 2013). The grafted plants were grown in nonwoven fabric garden bags (30 cm in height, 25 cm in diameter) containing 85% yellow podzolic soil, 10% coconut husk, and 5% fertilizer (v:v:v). Major components of the fertilizer included 

- **C. chrysantha**: 44% urea treatment per plant (date: 16 Apr. 2016) (g)

| Urea treatment per plant (date: 16 Apr. 2016) (g) | 50 ppm | 150 ppm | 350 ppm | 750 ppm |
|------------------------------------------------|--------|---------|---------|--------|
| 1                                               | T1     | T2      | T3      | T4     |
| 3                                               | T5     | T6      | T7      | T8     |
| 5                                               | T9     | T10     | T11     | T12    |
| 8                                               | T13    | T14     | T15     | T16    |

PBZ treatment (date: 26 Apr. 2016, 29 May 2016, 24 June 2016)

| PBZ treatment (date: 26 Apr. 2016, 29 May 2016, 24 June 2016) | 50 ppm | 150 ppm | 350 ppm | 750 ppm |
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| 1                                                           | T1     | T2      | T3      | T4     |
| 3                                                           | T5     | T6      | T7      | T8     |
| 5                                                           | T9     | T10     | T11     | T12    |
| 8                                                           | T13    | T14     | T15     | T16    |

**Results**

*Effects of urea and PBZ on flower bud induction.* All plants used in our study produced flower buds for the first time in 2017 except for two plants in the T1 treatment (1 g urea + 50 ppm PBZ), two plants in T5 (3 g urea + 50 ppm PBZ), and one plant in T12 (5 g urea + 750 ppm PBZ). The three control plants produced a total of eight flower buds in Sept. 2016. Statistical analysis showed that
seven of the 16 treatment types, T1, T5, T6 (3 g urea + 150 ppm PBZ), T12, T13 (8 g urea + 50 ppm PBZ), T14 (8 g urea + 150 ppm PBZ), and T16 (8 g urea + 750 ppm PBZ), resulted in a similar flower bud number to the control. In contrast, treatment types of T2 (1 g urea + 150 ppm PBZ), T3 (1 g urea + 350 ppm PBZ), T4 (1 g urea + 750 ppm PBZ), T7 (3 g urea + 350 ppm PBZ), T8 (3 g urea + 750 ppm PBZ), T9 (5 g urea + 50 ppm PBZ), T10 (5 g urea + 150 ppm PBZ), T11 (5 g urea + 350 ppm PBZ), and T15 (8 g urea + 350 ppm PBZ) significantly increased the flower bud number. Thus, these treatment types were effective in flower induction (Fig. 2, dark color bars).

Likely because of the competition for nutrition, an average of 40.6% of the flower buds observed in Sept. 2016, were lost by mid-January when flower buds started to break. There was a moderate positive correlation between initial flower bud number and dropping rate, with a Pearson R value of 0.54 and a P value of 0.03. However, after this abscission, plants receiving the T2, T3, T7, and T11 treatments still had significantly more flowers and flower buds than the control group (11, 11, 16, and 14, respectively, vs. two per tree) (Fig. 2, light color bars). There was no significant difference among these four treatment groups, and their average number of combined flowers and flower buds was 13 per plant. In comparison, the control group had only two flowers per plant. Whereas T3, T7, and T11 differed in urea amount, 1, 3, or 5 g/tree, respectively, they all included three times’ application of 350 ppm PBZ. This indicates that in combination with 350 ppm PBZ, applying more than 1 g of urea per tree is not necessary because it did not result in extra flowers. In fact, when the applied urea reached 8 g/tree.

Fig. 1. Vegetative and reproductive buds and flower of golden camellia. (A) Buds in late Sept. 2016 without abnormal defoliation. (B, C) Buds in late Sept. 2016 with severe abnormal defoliation. (D, E) Buds in mid-Jan. 2017. (F) Flower in late Jan. 2017. r = flower buds; v = vegetative bud.

Fig. 2. Average number of flower and flower bud per *Camellia chrysantha* tree in Sept. 2016 and Jan. 2017. T1–T16: treatments with different combinations of paclobutrazol and urea. * indicates significant difference from control at P < 0.05.
Treatments with combined urea and PBZ were lower than 0.35, P number and abscission (absolute of defoliation severity with the flower bud correlation analysis found a weak relationship the defoliation effect of 50 ppm PBZ but in comparison, applying 8 g of urea worsened plants treated with 750 ppm suffered the most. had the lowest degree of defoliation, whereas in Table 2, plants treated with 50 ppm PBZ suffered severe defoliation (>50%). As shown medium category (30% to 50%), and 24 light defoliation (<30%), six were in the after the second and third PBZ treatments Defoliation occurred in all treated plants when they started to break, with a range of 10.9–17.7 mm and 12.5–18.5 mm, respectively (Fig. 1F). There was a strong correlation between flower bud diameter and length (R² = 0.85, P < 0.00001).

Effects of urea and PBZ on defoliation. Defoliation occurred in all treated plants after the second and third PBZ treatments were applied. By Jan. 2017, 18 plants had light defoliation (<30%), six were in the medium category (30% to 50%), and 24 suffered severe defoliation (>50%). As shown in Table 2, plants treated with 50 ppm PBZ had the lowest degree of defoliation, whereas plants treated with 750 ppm suffered the most. In comparison, applying 8 g of urea worsened the defoliation effect of 50 ppm PBZ but lessened the effect of 750 ppm PBZ. Pearson correlation analysis found a weak relationship of defoliation severity with the flower bud number and abscission (absolute R values were lower than 0.35, P value < 0.05).

Effects of urea and PBZ on growth. Treatments with combined urea and PBZ did not affect the growth of stem basal diameter because there was no significant difference between the treated plants and control (P > 0.05). In contrast, plants treated with T4, or T11 to T16 were significantly shorter than the control group, suggesting that these treatment types affect height growth. All these treatments include either high urea or PBZ (Fig. 4).

When the plants were grouped based on defoliation severity (light <30%, medium 30% to 50%, and severe >50%), no statistical difference in stem basal diameter was detected among them. In terms of height, plants that suffered severe defoliation were significantly shorter than the ones having light defoliation (Fig. 5). There was also a weak positive relationship between plant height and diameter (R² = 0.40, P value = 0.0004).

Relative chlorophyll content and C/N ratio. There was no significant difference in relative chlorophyll content among the treatments and control. A weak negative correlation was found between relative chlorophyll content and plant basal diameter (R² = –0.31, P value = 0.028). C and N contents were measured for three time periods: before treatments were applied (Apr. 2016), when flower buds developed rapidly (Sept. 2016), and when flower buds started to break in Jan. 2017. Because of budget constraints, leaf samples of three plants within the same treatment types were pooled for C and N analysis. As a result, there is no statistical power in this set of data. However, positive correlation was found between the plant height and C/N ratio measured in September (R² = 0.45, P value = 0.001) and January (R² = 0.35, P value = 0.02) and between plant basal diameter and C/N ratio measured in April (R² = 0.37, P value = 0.008) and September (R² = 0.61, P value < 0.001). This indicates that plant growth is affected by C/N balance.

Discussion

As a landscaping and a high-end indoor potted plant, golden camellia is highly prized for its flowers’ large size and unique color. In addition to their aesthetic appeal, golden camellia flowers are valued for tea because they contain chemical compounds that may support good health (Lin et al., 2013; Wang et al., 2015). Compared with red, purple, pink, or white camellia flowers, golden camellia flowers are rare, despite their high market demand. Among the yellow-flowered camellia species, C. chrysantha is the first discovered and best known. In this study, we treated 4-year-old grafted C. chrysantha plants with urea and PBZ and successfully increased flowering intensity. This is the first report on flower induction in a golden species with PBZ. To our knowledge, the only previous report that used PBZ for Camellia growth and flowering was on cuttings of two hybrids of a crossing of Camellia saluenensis with Camellia japonica (Wilkinson and Richards, 1988). These two red-flowered species belong to sections different from the golden camellias. In addition, the effect of PBZ on flowering differed between the two cultivars included in the study, whereas the plant height was reduced by 30% (Wilkinson and Richards, 1988). In another study, Luo et al. (2013) successfully increased the flower number in mature C. chrysantha seedlings (15–20 years old, basal diameter 3.0–3.9 cm) with ethephon. However, the optimal concentration for flower induction, 2.50 mg·L⁻¹, caused smaller flowers and leaf area. Sprouting of new shoots was also severely affected with this concentration. According to a report on guava (Psidium guajava L.) plants, PBZ was found more effective in increasing the fruit number, fruit yield, yield efficiency, and fruting density than ethephon (Brar and Bal,
were grafted on 1.5-year-old C. osmantha rootstocks in Aug. 2012 using 0.5- to 1-year-old shoots. By 2016 when our experiments started, none of these plants had flowered. Therefore, we decided to include PBZ in our experiment.

The C. chrysantha plants used in our study were grafted on 1.5-year-old C. osmantha rootstocks in Aug. 2012 using 0.5- to 1-year-old shoots. By 2016 when our experiments started, none of these plants had flowered.

Table 2. Defoliation of *Camellia chrysantha* plants treated with paclobutrazol (PBZ) and urea, with dark green representing <5% defoliation, dark red >50%, and yellow between 5% and 50%.

| Urea amount (g/plant) | PBZ (ppm per treatment) | 50 | 150 | 350 | 750 |
|-----------------------|-------------------------|----|-----|-----|-----|
| 1                     | T1                      | T12| T23 | T24 | T25 |
| 2                     | T3                      | T15| T16 | T17 | T18 |
| 3                     | T7                      | T10| T11 | T12 | T13 |
| 4                     | T11                     | T14| T15 | T16 | T17 |

This was expected because it usually takes 5 or 6 years and a basal stem diameter of at least 2.5 cm for grafted plants to set flower buds (Chai et al., 2009; Jiang and Zhao, 1997). While all trees included in the study were close to 5 years old, their basal diameters were smaller than 2 cm by the Spring of 2017. The fact that the three control plants, which received a single foliar spray of 800 ppm in May 2016, all produced flower buds for the first time in 2016 reaffirms the efficacy of PBZ. Because the mode of action of PBZ is the inhibition of GA synthesis in plants, our results suggest that decreasing GA level benefits flower bud initiation in *C. chrysantha*.

There are reports showing effects of nitrogenous compounds, such as urea, in increasing floral initiation and flowering intensity in both deciduous and evergreen species (Edwards, 1986; George and Nissen, 1993; Monselise, 1986; Nunez-Elisea, 1985). In particular, a study by Qi et al. (2012) compared nitrogen content in *C. chrysantha* shoots that had abundant, few, or zero flowers. It was found that shoots bearing abundant flowers contained more nitrogen, suggesting a role of nitrogen in flowering of *C. chrysantha*. In our study, urea, in combination with PBZ, effectively induced flowering in young *C. chrysantha* plants. Among the 16 urea and PBZ combinations, nine produced significantly more flower buds than the control (Fig. 2). A low PBZ concentration of 50 ppm was effective only in combination with 5 g urea. When the PBZ concentration was increased to 150 ppm, combinations with 1 and 5 g urea were effective. At the concentration of 350 ppm, PBZ was effective with all four dosages of urea included in the study, i.e., 1, 3, 5, and 8 g. At a higher concentration of 750 ppm, PBZ worked effectively with low dosages of urea (1 and 3 g) in flower bud induction. These results indicate interactions of PBZ and urea: a low concentration of PBZ can be effective if it is in combination with a higher dosage of urea, whereas high concentrations of PBZ and lower dosages of urea are better combinations.

Flower bud abscission is normally high in camellias, with up to 50% in some cultivars (McElwee, 1952). This phenomenon was observed in all the flowered *C. chrysantha* plants in our study, with 22% of the plants dropping more than 50% of their flower buds. We suspect that the young age and the small size of our plants contributed to the high abscission rate considering that balancing vegetative and reproductive development is essential to survival and the fact that abscission rate was higher in plants producing more flower buds (Pearson R value of 0.54 and a P value of 0.03). After taking abscission into account, four treatments types, T2, T3, T7, and T11, produced an average of 11, 11, 16, and 14 flowers per tree, respectively (Fig. 2). These numbers are significantly higher than the control, which only had two flowers per tree. Among these effective treatment combinations, 350 ppm PBZ was included in T3, T7, and T11 and 150 ppm PBZ in T2. As for urea application, T2 and T3 contained 1 g of urea, whereas T7 contained 3 g and T11 5 g.

Urea is a commonly used nitrogenous fertilizer because of its high nitrogen content and low cost. In our study, however, a high dosage of 8 g urea inhibited the flowering induction effect of PBZ at the concentrations of 50, 150, and 750 ppm. Whereas the combination of 8 g urea and 350 ppm PBZ was initially effective in inducing more flower buds than the control, most of the induced flower buds dropped before they could open, leading to no significant difference between this treatment combination and the control. Therefore, combinations of low-dosage urea (1–3 g) with 150–350 ppm PBZ are most effective when taking flower bud abscission into account.
Similar to experiments using ethephon (Luo et al., 2013), a high concentration of 750 ppm PBZ in our experiments significantly decreased either flower bud length, diameter, or both regardless of the dosage of urea (Fig. 3). Combinations including 350 ppm also affected flower bud size except for T3 (350 ppm PBZ+1 g urea). The effect of PBZ on stem diameter depends on the concentration and species. For instance, Yelenosky et al. (1995) reported a reduction by 12% to 50% in citrus rootstock seedlings, whereas Berova and Zlatev (2000) and Tsegaw et al. (2005) observed an increase in tomato and potato. In contrast, PBZ consistently reduces the plant height in a range of species (Baion et al., 2005; Mabvongwe et al., 2016; Pal et al., 2016; Terri and Millie, 2000). None of our treatment types had an influence on basal stem diameter. Neither did they change relative chlorophyll content, suggesting that photosynthesis is unaffected by the treatments. However, combinations that included either 750 ppm PBZ or 8 g of urea significantly reduced the plant height, with the exception of T8, which was 750 PBZ+3 g urea. The concentration 750 ppm PBZ also caused severe defoliation (>50%), with the exception of T16 (750 ppm PBZ+8 g urea) (Fig. 4). This may suggest that inclusion of an appropriate amount of urea can mitigate the effects of high concentrations of PBZ on the plant height and defoliation. When applied together, PBZ and urea may also have an additive effect on stem height reduction because it took a lower concentration of PBZ than 750 ppm and a lower dosage of urea than 8 g to have a height reduction effect in T11 (350 ppm PBZ and 5 g urea). In addition, a lower concentration of 350 ppm PBZ in combination with 3 g urea resulted in severe defoliation, indicating the existence of PBZ and urea interaction on defoliation. It is well known that environmental stresses, such as drought, poor nutrition, and defoliation, can stimulate flowering. For instance, 50% and 75% defoliation in kiwifruit vines could lead to about 75% and 53% increase of flower number, respectively (Cruz-Castillo et al., 2010). This stress-induced flowering may involve salicylic acid, the flowering gene FLOWERING LOCUS T, or both (Takeno, 2012). The fact that there exists positive correlation of defoliation severity with the flower bud number in our study (absolute R values were lower than 0.35, P value < 0.05) suggests the role of defoliation in precocious flowering of C. chrysantha.

To conclude, treatment types T2 (150 ppm PBZ+1 g urea) and T3 (350 ppm PBZ+3 g urea) can induce and sustain significant flowering in juvenile C. chrysantha grafted plants without negative effects on vegetative growth and flower bud size. Although defoliation was observed in the T2- and T3-treated plants, their severity was less than 50%. This approach can be explored for both field and indoor golden camellia plants for flowering manipulation. Currently, we are in the process of observing the growth of the 2016-treated plants and examining if they will continue to flower in 2018 without further PBZ treatments in 2017. The carryover effect of PBZ seems to depend on multiple factors, such as species, tree age, method of application, and concentration, because so far, reports from various sources are not consistent. For instance, responses of two Eucalyptus species to high levels of trunk injection and collar drenching persisted for up to six growing seasons, yielding both increases in frequency of flowering and heaviness of bud crop (Griffin et al., 1993). In contrast, response to sprays and media drenches did not carry over to subsequent years in the two red camellia hybrids studied by Wilkinson and Richards (1988). In the future, it is important to study whether PBZ treatments change the chemical composition in flowers and leaves because they are popularly used as tea and to identify an optimized solution to minimize defoliation. It is also informative to investigate the potential flowering induction effect of PBZ on juvenile C. chrysantha seedlings.

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