Early Flowering Induction in Golden Camellia Seedlings and Effects of Paclobutrazol

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Abstract. Camellia chrysantha flowers are in great market demand as a result of their high ornamental and medicinal values. To induce early flowering in 4-year-old juvenile C. chrysantha seedlings, three levels of paclobutrazol (PBZ) concentration (100, 200, and 300 ppm) were applied to the roots. PBZ is a triazole-type cytochrome P450 inhibitor that was found successful in inducing flowering in juvenile C. chrysantha grafted plants in a prior report. The current study shows that all three PBZ concentrations were equally effective in induction of floral buds, resulting in an average of 20 floral buds per treated plant. In comparison, none of the untreated plants flowered. Although the induced flowers were smaller than the ones from mature trees, PBZ treatment did not affect C. chrysantha flowers’ medical values, because there was no significant change in the content of pharmacologically active compounds (poly saccharide, polyphenols, flavonoids, and saponins). None of the PBZ treatments had a negative effect on the current year’s growth in height and basal diameter, photosynthesis, and levels of water-soluble sugars and nutrients [phosphorus (P), nitrogen (N), potassium (K), and carbon (C)]. It is concluded that PBZ is an effective flowering inducer for juvenile C. chrysantha plants. It was also found that PBZ-treated plants experienced defoliation, and there existed a strong correlation between severity of defoliation and PBZ concentration. This might be attributed by the stress induced by PBZ, as demonstrated by the increased activities of some of the stress-related enzymes [ascorbate peroxidase (APX), catalase (CAT), superoxide dismutase (SOD), and peroxidase (POD)], and the level of malondialdehyde (MDA). Considering that severe defoliation can cause stunted or malformed plants and reduce aesthetic value, 100 ppm is the optimal PBZ concentration for flowering induction in C. chrysantha seedlings.

Camellias are well-known worldwide as ornamental plants. The common flower colors are exclusively limited to red, pink, and white. In contrast, yellow camellias, which have golden yellow petals and are also collectively known as golden camellias, are rare in the world. This group of yellow-flowering camellias consists of 42 species and five varieties, distributed naturally in a narrow region of southern China and northern Vietnam (Zhuang, 2008). In addition to their unique flower color, golden camellia flowers contain diverse flavonoids, saponins, polyphenols, amino acids, and trace elements. These compounds have various bioactivities, including antitumor, antioxidant, hypolipidemic, hypoglycemic, immunomodulatory, antibacterial, anti-inflammatory, and antidepressant properties, as evidenced by numerous studies (e.g., He et al., 2018b; Wang et al., 2015). C. chrysantha is the first-discovered and best-known golden camellia species, and the Ministry of Health of China has listed it as a new food source (Guo, 2010). Currently, a wide range of health-oriented teas and beverages are made from C. chrysantha flowers in China. Products such as flower teas, oral liquids, buccal tablets, and shampoo are not only sold in Southeast Asian, but also in Europe, and America (reviewed in He et al. (2018a)). Although mature golden camellias can produce as many as 300 flowers per tree, it generally takes 6 to 8 years for seedlings and 5 to 6 years for grafted plants to start setting flower buds (Chai et al., 2009; Jiang and Zhao, 1997). During the first few years after flowering starts, only a few floral buds can be produced. This long juvenile phase not only hinders conventional breeding, but also has an adverse impact on economic incomes for commercial producers and breeders. Driven by its scarcity, the price of golden camellia flower tea is about 30,000 Yuan (≈$4600) per kilogram, whereas floral bud tea is about 20,000 Yuan (≈$3000), according to Su (2010) and a report in China Daily (China Daily, 2014). It is noteworthy that C. chrysantha has been listed as a first-grade endangered species in China as a result of its dramatic decline in natural population size (Xu, 1995). Since the establishment of the state-level natural reserve, which is in the primary distribution areas of the species, raw materials for manufacturing camellia products are mainly from cultivated plants. Therefore, there is strong interest in inducing early flowering in golden camellias for earlier harvest.

Considerable studies have demonstrated that phytohormones can induce early flowering in many plant species. However, the effect varies with the plant species, its age, and concentration of phytohormones used. For instance, gibberellic 3 treatment can activate SUPPRESSOR OF CONSTANS1 (SOC1) to induce flowering in Arabidopsis (Moon et al., 2003), and reduce apple flowering rate (Zhang et al., 2016). With ethephon, Luo et al. (2013) successfully increased flower number in mature 15- to 20-year-old C. chrysantha seedlings. However, the optimal concentration for flower induction, 2.50 mg·L⁻¹, caused slower flowers and leaf area. Sprouting of new shoots was also severely affected with this concentration of 2.50 mg·L⁻¹. PBZ [(2RS,3RS)-1(4-chlorophenyl)-4,4-dimethyl-2-(1H-1,2,4-triazol-1-yl)pentan-3-ol] is a triazole-type cytochrome P450 inhibitor that is used extensively in horticulture as a plant growth retardant and fungicide. This growth regulator has been found to be predominantly effective in inducing and manipulating flowering/fruiting in plant species such as mango (Blakie et al., 2004; 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). According to a report on guava (Psidium guajava L.), PBZ was found more effective in increasing fruit number, fruit yield, yield efficiency, and fruiting density than ethephon (Brag and Bal, 2016).

In a previous study with 4-year-old C. chrysantha-grafted plants with C. osmantha as rootstock, we tested the effect of PBZ and urea fertilizer on flowering induction with a 4 × 4 factorial design: four dosages of urea [0, 3, 5, or 8 g/plant] and four concentrations of PBZ (50, 150, 350, or 750 ppm) (Wei et al., 2017). It was found that 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 without severe defoliation. In the current study, the goal was to elucidate the effects of PBZ on floral induction in juvenile *C. chrysantha* seedlings.

**Materials and Methods**

*Plant materials and experiment design.* Seeds of *C. chrysantha* were sown in Mar. 2013 and grown in nonwoven fabric garden bags (height, 30 cm; diameter, 25 cm) containing yellow podzolic soil, with a pH range of 4.5 to 6.0. Healthy seedlings with uniform growth (height, 0.85–1.3 m; basal diameter, 10.3–15.5 mm) were selected for the study (Fig. 1A). Three levels of PBZ concentrations (100, 200, or 300 ppm) were prepared with tap water, then 1 L of solution was applied to the garden bags on the mornings of 23 Mar. 2017 and 14 Apr. 2017. The control plants were managed in the same way as the treated group except with no PBZ applied to the garden bags. PBZ (CAS no. 76738-62-0) was purchased from Anyang Quanfeng Biological Technology Co. Ltd. (He Nan Province, China) and contained 95% active components.

*Study site.* The study was conducted in the Camellia Nursery of Guangxi Forestry Research Institute, China (lat. 22°56′N, long. 108°21′E, 95 m.a.s.l.). With a subtropical monsoon climate, the area has distinct dry and wet seasons. The average annual temperature is 21.8 °C, whereas the average in January is 11.8 °C, and 27.6 °C in July. The recorded coldest and hottest temperatures are −1.5 °C and 39.4 °C, respectively. Annual accumulated temperature of at least 10 °C is 7,200 °C. The rainy season is during May to September, with an annual precipitation of more than 1300 mm. All plants were maintained under a shade canopy that was 2.9 m aboveground and blocked 50% to 60% sunlight.

*Sample and data collection.* When floral buds started to open in Jan. 2018, floral buds and flowers were counted. Defoliation percentage was estimated and categorized into five groups: A, <1%; B, (1%–5%; C, 6%–30%; D, 31%–40%; E, 41%–50%; and F, >50%. Size (length and width) of at least four fully opened flowers (if available) and fresh weight were measured for each plant. Healthy young leaves (freshly opened from current-year shoots) and old leaves from the past year’s growth were harvested from each plant and used for analyses of activities of stress-related enzymes [APX, CAT, polyphenol oxidase (PPO), SOD, and POD], and levels of MAD, chlorophylls, water-soluble sugar, and nutrients (N, P, K, and C).

Fig. 1. *C. chrysantha* seedlings and their paclobutrazol (PBZ)-induced reproductive buds and flowers. (A) Four-year-old *C. chrysantha* seedlings in January 2017 before PBZ treatment. (B–E) Reproductive buds in (B) late May, (C) mid October, (D) mid December 2017, and (E) early 2018. (F) Fully opened flowers in late Jan. 2018. (G) A 300 ppm-treated plant with severe defoliation in mid Dec. 2017. The scale bars represent 10 cm in each panel.
flowers were from a 35-year-old plant. PBZ treatments increased floral buds. The average number of floral buds per plant induced by PBZ was 20, and there was no statistical difference among the three levels of PBZ concentration. The results are similar to the ones from *C. chrysanth a* grafted plants. When treated with PBZ, *C. chrysanth a* grafted plants produced a range of 9 to 33 floral buds per plant, depending on the PBZ concentration and its combination with urea (Wei et al., 2017). Thus, it is corroborated that PBZ is an effective flower inducer for *C. chrysanth a* juveniles. In the experiment conducted in 2016 (Wei et al., 2017), it was found that 33% of the plants that flowered in 2017 continued to flower in 2018, although they did not receive PBZ treatment in 2017. It is notable that size and fresh weight of PBZ-induced flowers were smaller than the ones from a 35-year-old *C. chrysanth a* plant (31.9 vs. 46.0 mm, 4.75 vs. 6.9 g, respectively), whereas their dry weight was similar (0.82 vs. 0.71 g). There was no statistical differences among the three levels of PBZ concentration in terms of flower size, and fresh and dry weights. Similar to the observations in grafted *C. chrysanth a* plants (Wei et al., 2017), defoliation occurred in PBZ-treated seedlings in this study. Although the 100-ppm-treated plants lost less than 30% of their leaves, the 200- and 300-ppm-treated seedlings lost 40% to 50% and more than 50% of leaves, respectively. There existed a strong correlation between severity of defoliation and PBZ concentration (absolute R value, 0.87; P value, <0.01). 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 25% and 53% increase in 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). However, we did not observe a significant correlation between defoliation severity and flower bud number in the current study (absolute R value 0.41; P value > 0.1). In the previous study with grafted *C. chrysanth a* plants, there existed a weak positive correlation (absolute R values < 0.35; P value < 0.05) (Wei et al., 2017).

**Plant growth, chlorophyll content, water-soluble sugars, and nutrients.** When plant height and basal diameter were compared, a significant difference was not found between the treated and control plants, nor was one found among the treated ones. This is consistent with the previous study on grafted *C. chrysanth a* when treated with comparable PBZ concentrations (Wei et al., 2017). As a growth retardant, PBZ inhibits plant height and diameter, as well as bud, leaf, and fruit size, when applied in high concentrations, as documented in numerous studies (e.g., Mabongwwe et al., 2016; Pal et al., 2016; Wei et al., 2017). Similarly, there was no significant difference in chlorophyll a and b content in young leaves among the treatments and control. However, the 200-ppm treatment decreased both the chlorophyll a and b levels significantly, by 20% and 26%, respectively, in old leaves when compared with the untreated control (Fig. 2).

PBZ seemed to decrease water-soluble sugar content in old leaves; however, the reduction was not statistically significant (P > 0.05). Total C content in both young and old leaves was also similar among control and PBZ treatments, with an average of 454 ± 14.5 g/kg. These results are consistent with the fact that chlorophyll a and b content in young leaves is not significantly affected by the PBZ concentrations used in our study. PBZ-treated plants contained similar levels of C, N, and P contents determined by standard K2Cr2O7/H2SO4 oxidation, Kjeldahl, and NaHCO3 extraction/molybdenum–antimony anticolorimetric methods, respectively. Inductively coupled plasma–atomic emission spectrometry was used for K analysis. Nutrient contents were calculated as milligrams per kilogram dry weight. Sample preparation and measurement were performed by a core facility at Guangxi Forestry Research Institute (Nanning, China).

Fully open flowers were collected from each plant, dried in a lyophilizer (Heto FD3 freeze dryer; Heto-Holten A/S, Copenhagen, Denmark), and ground into a fine powder. Pharmacologically active compounds were analyzed in a core facility at Guangxi Forestry Research Institute (Nanning, China). Total flavonoids were extracted with 70% ethanol and measured using the aluminum nitrate colorimetric method, with absorbance taken at 500 nm (Wang et al., 2012). For crude saponins, flowers were heated in ddH2O at 80 °C and the extracts were purified with ethanol and diethyl ether as described in Shah et al. (2014). Determination of the total saponin content was conducted using the colorimetric method as described in Murakami et al. (2013). The reaction reagents included vanillin, perchloric acid, and glacial acetic acid. Absorbance wavelength was 540 nm. Polysaccharides were measured with the standard phenol–sulfuric acid reaction method, and absorbance was taken at 490 nm (Dubois et al., 1956). Polyphenols were extracted with 70% methanol and determined using the Folin–Ciocalteu method (Turkmen et al., 2006). The absorbance wavelength was 765 nm. The analyses of chemical components in flowers were performed by a facility in the College of Forestry, Guangxi University, Nanning, China. The control flowers were from a 35-year-old *C. chrysanth a* plant.

**Data analysis.** A completely randomized design was used in the experiments. There were four plants per treatment type and control. All chemical analyses were conducted individually with each plant unless indicated otherwise. For each biologic sample, there were three technical replicates. All statistical analyses were performed using SPSS (version 17.0; IBM, Armonk, NY), and significance was set at P < 0.05, as previously reported in Wei et al. (2017).
occurring flowers in our study had a greater polysaccharide content (40.8 mg/g) than the ranges reported by Niu et al. (2014) and Tang et al. (2017), which were 32.8 and 21.4 mg/g, respectively. However, the PBZ-induced flowers contained a polysaccharide level (25.5 mg/g) similar to the reports. Niu et al. (2015) reported a level of 114 mg/g saponins in *C. chrysanthemum* flowers, similar to what we detected in PBZ-induced flowers (106 mg/g). As for flavonoids, both PBZ-induced and naturally occurring flowers in our study showed a comparable level to the report by Niu et al. (2015): 180, 159, and 136.3 mg/g respectively. Overall, PBZ did not change significantly the levels of beneficial compounds in the induced flowers.

**Stress caused by PBZ.** PBZ treatment causes defoliation, and there exists a strong correlation between severity of defoliation and PBZ concentration. When commonly used stress-related markers (APX, CAT, PPO, SOD, POD, and MAD) were examined (Fig. 5), it was found that PBZ treatment did not have an effect on POD, regardless of leaf age. Levels of CAT and APX increased in both young and old leaves of plants treated with 200 and 300 ppm. All three PBZ concentrations used resulted in an enhancement in SOD activity in young leaves, but no effect in old leaves was significant. PPO decreased in young leaves of 200- and 300-ppm-treated plants, and decreased in old leaves of all PBZ-treated plants. In comparison, 300 ppm PBZ increased the MDA level in young leaves significantly, whereas 200 and 300 ppm decreased its level in old leaves. Overall, 300-ppm-treated plants suffered the most severe defoliation: Two of the treated plants lost most of their leaves by January and died by May 2018. Consistent with defoliation severity, young leaves of 300-ppm-treated plants contained significantly greater levels of CAT and APX than 100-ppm-treated plants, and their MDA content was significantly greater than both 100- and 200-ppm-treated plant.

Various stresses can induce oxidative damage in plants, in which reactive oxygen species, such as superoxide radical (O$_2^-$), hydroxyl radical (·OH), hydrogen peroxide (H$_2$O$_2$), and alkoxyl radical (RO·), are produced. The toxic superoxide radical is usually dismutated rapidly by SOD to H$_2$O$_2$—a product that is relatively stable and can be detoxified by CAT and peroxidases. SOD, POX, APX, CAT, and PPO are enzymatic antioxidants that participate in alleviation of oxidative damage (Prochazkova et al., 2001), and their activities generally increase under stress (Shigeoka et al., 2002). Our results generally follow this trend, except for POD and PPO, and there existed a difference in reaction between young and old leaves. MDA results from lipid peroxidation of polyunsaturated fatty acids is an indicator of free radical production and consequent tissue damage. Our data indicate that 300 ppm PBZ resulted in significant oxidative lipid damage in young leaves.

To conclude, PBZ is an effective flowering inducer for juvenile *C. chrysanthemum*, and 100 ppm provides the best results in seedlings in comparison with greater concentrations. Although the induced flowers were smaller than the ones from mature trees, there were no significant changes in the contents of pharmacologically active compounds.
(polysaccharide, polyphenols, flavonoids, and saponins). Thus, PBZ treatment does not affect C. chrysantha flowers' medicinal value. PBZ is a growth retardant; however, the concentrations effective in flowering induction in C. chrysantha had no negative effect on the current year’s growth in height and basal diameter. Severe defoliation is a major concern, considering that it not only may cause stunted or malformed plants, and even death to the plant in the long term, but also it reduces the plant’s aesthetic appeal. In addition, leaves of golden camellias are valued for tea. PBZ treatment led to enhanced stress, which might be a key contributing factor for defoliation. Currently, an investigation is being undertaken to study whether reducing treatment frequency (from twice to once) and

![Graphs showing activities of peroxidase (POD), catalase (CAT), ascorbate peroxidase (APX), superoxide dismutase (SOD), and polyphenol oxidase (PPO), and level of malondialdehyde (MDA) in young and old leaves of paclobutrazol-treated and untreated C. chrysantha plants. Samples were collected in Jan. 2018. Within each graph, different letters indicate significant difference at \( P < 0.05 \). FW = fresh weight.](image)
solution volume applied (from 1 L to 0.5 L) will lessen the effect of PBZ on defoliation. In future studies it would be informative to investig- ate the changes of endogenous hormones, such as auxins, gibberellins, cytokinins, ethy- ene, and abscisic acid, during bud differentiation after PBZ treatment.

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