Sex-specific floral morphology, biomass, and phytohormones associated with altitude in dioecious Populus cathayana populations

Jundong He¹ | Tingfa Dong¹ | Kechao Huang² | Yanxia Yang¹ | Dadong Li¹ | Xiao Xu¹ | Xinhua He³,⁴

¹Key Laboratory of Southwest China Wildlife Resources Conservation (China West Normal University), Ministry of Education, Nanchong, Sichuan, China
²Guangxi Institute of Botany, Chinese Academy of Sciences, Guilin, Guangxi, China
³Centre of Excellence for Soil Biology, College of Resources and Environment, Southwest University, Beibei, Chongqing, China
⁴School of Plant Biology, University of Western Australia, Crawley, WA, Australia

Abstract
Relationships between sex-specific floral traits and endogenous phytohormones associated with altitude are unknown particularly in dioecious trees. We thus examined the relationships between floral morphology or biomass and phytohormones in male and female flowers of dioecious Populus cathayana populations along an altitudinal gradient (1,500, 1,600, and 1,700 m above sea level) in the Xiaowutai Nature Reserve in northern China. The female and male flowers had the most stigma and pollen at 1,700 m, the largest ovaries and least pollen at 1,500 m, and the smallest ovaries and greater numbers of anthers at 1,600 m altitude. The single-flower biomass was significantly greater in males than in females at 1,600 or 1,700 m, but the opposite was true at 1,500 m altitude. The biomass percentages were significantly higher in anthers than in stigmas at each altitude, while significantly greater gibberellin A3 (GA₃), zeatin riboside (ZR), indoleacetic acid (IAA), and abscisic acid (ABA) concentrations were found in female than in male flowers. Moreover, most flower morphological traits positively correlated with IAA in females but not in males. The biomass of a single flower was significantly positively correlated with ABA or IAA in males but negatively with ZR in females and was not correlated with GA₃ in both females and males. Our results demonstrate a distinct sexual adaptation between male and female flowers and that phytohormones are closely related to the size, shape, and biomass allocation in the pollination or fertilization organs of dioecious plants, although with variations in altitude.

Keywords
abscisic acid, female and male flowers, floral morphology, gibberellin A3, indoleacetic acid, zeatin riboside

1 | INTRODUCTION

As the defining organ of the angiosperms, the flower performs a reproductive function and exhibits relatively high variability in response to environmental changes (Espírito-Santo et al., 2003; Humeau, Pailler, & Thompson, 2000; Malaspina et al., 2007). For example, calyx lengths vary significantly along a latitudinal transect (Olsson & Ågren, 2002), and petal size and ovary weight greatly increase under higher soil
temperatures (Poerwanto & Inoue, 1990). Furthermore, the staminal column and standard petal lengths markedly decrease with increased UV-B radiation (Koti, Reddy, Reddy, Kakani, & Zhao, 2005). These findings indicate that floral morphology is sensitive to abiotic factors.

Altitude is an important abiotic factor associated with temperature, precipitation, light, and soil physicochemical properties (Körner, 2007), and studies have shown that floral characteristics are significantly affected by altitude (Bodson & Outlaw, 1985; Nagano et al., 2014). For instance, Duan, He, and Liu (2005) demonstrated that the floral display in Gentiana straminea increased with increasing altitude, and Baonza and Malo (1997) found that the floral size of Cytisus scoparius showed a clinal variation with larger flowers at higher altitudes ranging from 700 m to 1,500 m. Kudo and Molau (1999) observed that the floral size (as well as anthesis) in Astragalus alpinus was significantly greater in higher population. In contrast, Nagano et al. (2014) found that the floral size of Campanula punctata var. hondoensis decreased with increasing altitude. These inconsistent variations in the floral characteristics of different species at different altitudes may be related to changes in biomass allocation (Li, Xu, Zang, Koripelainen, & Berninger, 2007; Pickering, 2000; Zhao, Du, Zhou, Wang, & Ren, 2006) as plants may allocate more carbon to reproductive organs (i.e., the flower; Fabbro & Körner, 2004; Hautier, Randin, Stöcklin, & Guisan, 2009) or to flower physiological traits at higher altitudes (Chandler, 2011; van Doorn & van Meeteren, 2003).

On the other hand, endogenous phytohormones or plant growth regulators (e.g., abscisic acid [ABA], auxins, cytokinins) are able to regulate floral initiation and development, and changing the level of any phytohormone could affect floral formation during the transition from vegetative to reproductive growth (Chandler, 2011; Law, Lebel-Hardenack, & Grant, 2002; van Doorn & van Meeteren, 2003; Villacorta, Fernández, Prinsen, Bernad, & Revilla, 2008). For example, ABA has a molecular effect on downstream events in the autonomous floral pathway and, consequently, on the transition to flowering (Razem, El-Kereamy, Abrams, & Hill, 2006; Su, Huang, Shen, & Chen, 2002); thus, ABA concentrations are dramatically increased during the flower development (Domagalska, Samowska, Nagy, & Davis, 2010). Gibberellic acid, or gibberellin A3 (GA3), plays an essential role in the development of floral organs (Goto & Pharis, 1999; Sawhney, 1983) and increases the numbers of petal, stamens, carpels and locules (Carrera, Ruiz-Rivero, Peres, Atares, & Garcia-Martinez, 2012), and flowers (Chen, Henny, McConnell, & Caldwell, 2003). The variation in indoleacetic acid (IAA) correlates with early floral initiation (Ding et al., 1999), and the application of IAA may induce flowering (Brcko et al., 2012; Wang & Guo, 2015). As a high activity of the cytokinin, zeatin riboside (ZR) can promote cell division, stimulate floral formation, and prevent leaf senescence by activating gene expression and metabolic activity (Galoch, Czaplewskia, Burakcka-Lauatkaitis, & Kopcewicz, 2002; Singh, Palni, & Latham, 1992; Subbaraj, Funnell, & Woolley, 2010), and its concentrations are significantly increased in the leaf, leaf exudate, and shoot apical meristem during early floral transition events (Corbesier et al., 2003). However, the function of these phytohormones in regulating floral formation in dioecious plants is less well known, particularly along an altitudinal gradient.

Dioecious plants constitute 6% of the total angiosperm species worldwide and play important roles in maintaining the sustainability of terrestrial ecosystems (Renner & Ricklefs, 1995). In general, more and larger flowers are produced in the male plants of Diospyros pentamera, Litsea leefean, and Neolitsea dealbata than in female plants (Houe, 1992). Meanwhile, the male plants of Borderea pyrenaica also display significantly greater variation in flower size during flowering (Thomas & Lafrankie, 1993). In contrast, female plants allocate more biomass to growth during the early flowering period than male plants (Delph, 1990; Gross & Soule, 1981; Koripelainen, 1992) and also contribute more carbon to floral performance (Laporte & Delph, 1996). Furthermore, the male and female plants of dioecious trees exhibit significant differences in sex ratio, physiological processes, or antioxidant defense enzymes under changes in numerous environmental conditions, including altitude elevation (Lei, Chen, Jiang, Yu, & Duan, 2017; Li et al., 2007), increased temperature (Xu et al., 2008), elevated CO2 concentration (Wang & Griffin, 2003; Zhao, Xu, Zhang, Koripelainen, & Li, 2011), enhanced UV-B radiation (Chen et al., 2016; Xu et al., 2010), nitrogen status (Chen, Dong, & Duan, 2014; Li, Dong, Guo, & Zhao, 2015; Li & Koripelainen, 2015), and competition (Chen, Duan, Wang, Koripelainen, & Li, 2014), but limited information is available on how sexual differences in floral traits in woody species are affected by altitude.

*Populus cathayana* Rehd., a dioecious woody tree, is widely distributed in northern, central, and southwestern China, including mountainous areas at altitudes from 1,000 to 3,000 m above sea level (a.s.l.). Our previous studies addressed the different growth and floral performance responses to elevated temperatures and UV-B radiation (Xu et al., 2008, 2010) and relationships among twig components between male and female *P. cathayana* saplings (Yang, He, Xu, & Yang, 2015). To further address whether sexual differences in the floral traits of *P. cathayana* could vary with altitude, this study aimed to determine (1) how sex-related differences in the morphology, biomass, and phytohormones of flowers could respond to altitude changes and (2) what the possible intrinsic relationships between morphological traits or biomass and phytohormone levels could be in male and female flowers. The expected results could provide insights into the adaptive physiological responses of flowers or reproductive organs to variations in altitude and the contribution of phytohormones to the morphological traits and biomass production of flowers in dioecious trees.

2 | MATERIALS AND METHODS

2.1 | Study site

The study site is located in the Xijin River Valley of Xiaowutai Mountain Nature Reserve in Hebei, China (39°50′−40°07′N, 114°47′−115°30′E; 1,142−2,882 m a.s.l.). This site area is characterized by a warm-temperate continental monsoon climate with mean annual precipitation of 528 mm and a mean annual temperature of 3.5°C. The major soil types are Alfisols, Aridisols, and Inceptisols (USDA soil taxonomy). There are five distinct forest zones along the
western slope of Xiaowutai Mountain: the deciduous shrub zone, the deciduous broad-leaved forest zone, the mixed coniferous and broad-leaved forest zone, the conifer forest zone, and the subalpine meadow zone (Liu, Zheng, & Fang, 2004). The forest vegetation is dominated by species in the Acer, Birch, Cerasus, Corylus, Quercus, Populus, Tilia, or Ulmus genus. The natural secondary Ulmus zone (Liu, Zheng, & Fang, 2004). The forest vegetation is dominated across the deciduous broad-leaved forest zone (1,400–1,800 m a.s.l.), but it has been gradually replaced by Betula platyphylla above 1,700 m a.s.l. (Yu, Liu, & Cui, 2002).

2.2 | Plant sampling

Five sampling sites were established at each altitude of 1,500, 1,600, and 1,700 m a.s.l. along the Xijin River Valley where the natural P. cathayana population density is relatively high over the northern China, and in the middle of April (the beginning of the P. cathayana flowering season) 2013. Ten (five males and five females) mature trees of similar size were randomly selected at each sampling site. The selected trees were (1) healthy with full-grown crowns reaching the average tree crown height (canopy); (2) located away from the forest edge on a ridge or next to a previously sampled tree; (3) more than 30 years old, approximately 20–25 m high, and 80–100 m apart. During the first 5–7 days after anthesis, five male or female inflorescences (in the same flowering period) on the south or sunny side of trees were randomly chosen from the outer surface of the crown for the measurement of floral traits.

2.3 | Measurements of floral morphological traits

The inflorescence length was measured with a micrometer, and the number of flowers per inflorescence was then counted before the flowers were removed from the inflorescence. Five randomly selected male or female flowers (one middle, two terminal, and two basolateral flowers) per inflorescence were then dissected with the aid of a stereoscope (Leica, M205C; Leica Microsystems, Wetzlar, Germany), and the pedicel, sepal, floral disk, anther, and filament (or pedicel sepal, floral disk, stigma, and ovary) of each flower were then dissected under a stereoscopic microscope equipped with a charge-coupled device (CCD) camera (MoticamPro285A; Motic, Xiamen, China). The number of anthers per flower, pollen grains per anther, and ovules per ovary were recorded, and the sizes (length, width, or diameter) of the individual parts (pedicel, sepal, ovary, and stigma) were measured to the nearest 0.01 mm using an ocular reticle.

To calculate the number of pollen grains, 50 randomly selected undehisced anthers (one anther per flower) were soaked in 1.0 mol HCl solution for 1 hr at 60°C to dispose of the anther wall, and 10.0 ml 0.9% NaCl solution was added after grinding (method modified from Guo, Wang, and Weber (2013)). A 2.0-μl suspension was plated on a hemocytometer (with a blood-cell counting chamber with 400 small, square grids in a central 1.0-mm square), and the pollen grains per anther was calculated. After dissecting the ovary on a slide, ovules were counted under the above-mentioned stereoscopic microscope equipped with a CCD camera.

2.4 | Measurement of flower biomass traits

The biomass production of the male and female flowers measured for their morphological traits was recorded. The samples were oven-dried at 70°C for 48 hr to a constant weight. The biomass of the individual anther, stigma, or flower was then determined, and the weight of the anthers or stigma per single flower was accordingly calculated as a percentage.

2.5 | Phytohormone measurements

The five male or female inflorescences measured for their morphological traits were also used to measure the concentrations of ABA, GA₃, IAA, and ZR. The samples were homogenized in liquid nitrogen and extracted in cold 80% (v/v) methanol with butylated hydroxytoluene (1 mmol/L) overnight at 4°C. The extracts were collected after centrifugation at 10,000 g (4°C) for 20 min, and the extracts were passed through a C₁₅ Sep-Pak cartridge (Waters, Milford, MA) and dried in N₂ to prepare for an enzyme-linked immune absorbent assay according to the method of Yang, Xu, Wang, and Jia (2001). Prior to the phytohormone measurements and after the removal of their floral axis and pedicels, the inflorescences were wrapped with aluminum foil and immersed in liquid nitrogen. The phytohormone measurements were performed in the Key Laboratory of Molecular Plant Pathology, Ministry of Agriculture, Beijing, China.

2.6 | Statistical analysis

Data (means ± SE, n = 5) analyses were performed using SPSS 17.0 (SPSS Inc., Chicago, IL, USA). One-way ANOVA was used to determine differences in the flower morphological traits among altitudes, and Duncan’s multiple range tests were employed to detect significant differences among means at p ≤ .05. Two-way ANOVAs were used to separate the effects of sex, altitude, and their combination. Pearson’s correlation coefficients were calculated to determine the relationships between the biomass and phytohormone concentrations of male or female flowers, and a simple linear regression was used to examine these relationships.

3 | RESULTS

3.1 | Variations in the morphological traits of female and male flowers

Almost all tested flower morphological traits were significantly affected by altitude (Table 1, Figure 1). Among the female flowers, the number of flowers per inflorescence and stigma width increased with altitude elevation, while flowers at 1,600 m had the lowest values for inflorescence length, pedicel length, sepal size, ovary diameter, and number of ovules per ovary compared to their counterparts at other two altitudes (Table 1). Among the male flowers, the sepal size, single-anther biomass, and number of pollen grains per anther significantly increased with altitude (p < .05), and plants at 1,700 m had the highest...
**Table 1** Morphological traits of female and male flowers of *Populus cathayana* at three altitudes (1,500, 1,600, and 1,700 m a.s.l.) in the Xiaowutai Nature Reserve, northern China

| Traits                              | Altitude       |                          |                          |                          |                          | \(p\_S\) | \(p\_A\) | \(p\_S&A\) |
|-------------------------------------|----------------|--------------------------|--------------------------|--------------------------|--------------------------|---------|---------|-----------|
|                                     | 1,500 m        |                          |                          |                          |                          |         |         |           |
|                                     | Female         | Male                     | Female                   | Male                     | Female                   | Male    |         |           |
| Inflorescence length (mm)           | 50.73 ± 3.28 ab, x | 53.78 ± 2.69 \(\alpha\), x | 45.15 ± 1.06 b, x       | 42.83 ± 0.91 \(\beta\), x | 54.44 ± 2.09 a, x       | 49.62 ± 2.55 \(\alpha\), x | .468 ns | .001*** | .228 ns   |
| No. of flowers per inflorescence    | 30.00 ± 4.34 b, y | 42.40 ± 3.01 \(\alpha\), x | 42.00 ± 0.22 ab, x      | 37.60 ± 1.30 \(\alpha\), y | 43.40 ± 2.66 a, x       | 42.40 ± 1.36 \(\alpha\), x | .270 ns | .046*   | .007**    |
| Sepal size (mm)                     | 0.94 ± 0.04 a, x | 0.99 ± 0.02 \(\gamma\), x | 0.69 ± 0.01 c, y         | 1.47 ± 0.03 \(\alpha\), x | 0.76 ± 0.02 b, y         | 1.20 ± 0.04 \(\beta\), x | <.001*** | .001***  | <.001***  |
| Ovary length (mm)                   | 2.21 ± 0.03 a, y | 2.73 ± 0.05 \(\beta\), x | 2.48 ± 0.01 c, y         | 2.85 ± 0.05 \(\alpha\), x | 1.70 ± 0.05 b, y         | 3.17 ± 0.04 \(\alpha\), x | <.001*** | <.001***  | <.001***  |
| Ovary diameter (mm)                 | 2.68 ± 0.12 a   | –                        | 2.26 ± 0.02 b            | –                        | 2.38 ± 0.07 ab           | –        | –        | .010*     |
| Stigma length (mm)                  | 2.86 ± 0.10 a   | –                        | 2.26 ± 0.02 c            | –                        | 2.34 ± 0.04 bc           | –        | –        | <.001***  |
| Stigma width (mm)                   | 2.38 ± 0.09 a   | –                        | 2.43 ± 0.01 a            | –                        | 2.71 ± 0.17 a            | –        | –        | .105 ns   |
| No. of ovules per ovary             | 3.32 ± 0.16 b   | –                        | 3.49 ± 0.02 ab           | –                        | 4.09 ± 0.20 a            | –        | –        | .008**    |
| No. of anthers per flower           | 33.62 ± 0.32 ab | –                        | 27.38 ± 0.25 b           | –                        | 34.43 ± 0.75 a           | –        | –        | <.001***  |
| Single-anther biomass (µg)          | –              | 26.98 ± 1.08 \(\beta\)   | 36.04 ± 1.30 \(\alpha\)  | –                        | 33.62 ± 0.98 \(\alpha\)  | –        | –        | <.001***  |
| No. of pollen grains per anther     | –              | 64.13 ± 2.82 \(\beta\)   | 64.66 ± 1.52 \(\alpha\)  | –                        | 73.44 ± 2.95 \(\alpha\)  | –        | –        | .039*     |

Different letters following the data (means ± SE, \(n = 5\)) in the same row denote significant differences between altitudes among females (\(a, b, c\)) and males (\(\alpha, \beta, \gamma\)) or between sexes at the same altitude (\(x, y\)), respectively, according to Duncan’s multiple range test at \(p < .05\). The significance values of the factorial analysis (ANOVA) for \(p\_S\), sex effect; \(p\_A\), altitude effect; and \(p\_S&A\), sex and altitude interaction effects are denoted as ns, not significant.

\(\ast p < .05; \ast\ast p < .01; \ast\ast\ast p < .001\).
values for these traits (Table 1). However, no significant effects of altitude were observed on the number of male flowers per inflorescence ($p = .17$). In addition, compared with other altitudes, male plants at 1,600 m had the shortest inflorescence length, the longest pedicel, and the greatest number of anthers per flower (Table 1). Moreover, compared with the females, male flowers had a significantly larger sepal size at the same altitude, a longer pedicel length at 1,600 or 1,700 m, and a higher number of flowers per inflorescence at 1,500 m ($p < .05$; Table 1).

### 3.2 Variations in single-flower biomass and its allocation in the two sexes

Significantly greater single-flower biomass was observed among altitudes in the order of 1,500 m > 1,700 m > 1,600 m for the female flowers, whereas the order was 1,700 m > 1,600 m > 1,500 m for the male flowers (Figure 2a). Moreover, males had greater single-flower biomass than females at altitude of 1,600 or 1,700 m but less biomass at 1,500 ($p < .05$). In addition, a significantly greater stigma biomass percentage was observed among altitudes in the order of 1,600 m > 1,700 m > 1,500 m, whereas the biomass percentage of the anthers was similar among altitudes (Figure 2b). The anther biomass percentage was always significantly higher than the stigma biomass percentage at all altitudes ($p < .001$; Figure 2b).

### 3.3 Variations in flower phytohormone concentrations in the two sexes

For both male and female flowers, ABA concentrations were significantly increased with altitude ($p < .001$) in the order of 1,700 m > 1,600 m > 1,500 m (Figure 3a), while flowers at 1,600 m had the highest GA$_3$ and ZR concentrations compared to the other two altitudes (Figure 3b,d). As a general rule, there were sex-related differences in the tested phytohormones as the concentrations of ABA, GA$_3$, IAA, and ZR were significantly higher ($p < .001$) in female than in male flowers at each of the three altitudes (Figure 3).
3.4 | Relationships between morphological traits and phytohormone concentrations in the two sexes

In female flowers, significantly positive correlations were observed between ABA concentrations and the sepal size or the number of ovules per ovary as well as between IAA concentrations and the pedicel length, sepal size, ovary length, ovary diameter, or the number of ovules per ovary (bold values in Table 2). In contrast, significantly negative correlations were exhibited between ZR concentrations and the pedicel length or ovary length, and no correlations were observed between GA₃ and any of the female flower trait examined (Table 2). In male flowers, significantly positive correlations were found between ABA concentrations and sepal size, between GA₃ concentrations and pedicel length, and between ZR concentrations and the pedicel length or number of anthers per flower (bold values in Table 3). However, no correlations were observed between IAA and any of the tested female flower traits (Table 3). In addition, GA₃ significantly positively correlated with ZR in both female and male flowers (Tables 2 and 3). In female flowers, ABA significantly positively correlated with ZR (Table 2) but significantly negatively correlated with GA₃ or ZR in male flowers (Table 3).

3.5 | Relationships between flower biomass and phytohormone concentrations in the two sexes

The biomass of a single flower was significantly positively related to concentrations of ABA and IAA in male but not in female flowers \((p < .01; \text{Figure 4a or c})\), while was negatively related to ZR concentrations in female but not in male flowers \((p < .05; \text{Figure 4d})\). In addition, no relationships were observed between the biomass of a single
**TABLE 2** Correlation coefficients among morphological traits and phytohormone concentrations (ng/g FW) in female flowers of *Populus cathayana*

|                 | Pedicel length | Sepal size | Ovary length | Ovary diameter | Stigma length | Stigma width | No. of ovules per ovary | GA3  | ZR  | IAA  | ABA  |
|-----------------|----------------|------------|--------------|----------------|---------------|--------------|------------------------|------|-----|------|------|
| Pedicel length  | -              | .718**     | .866**       | .850**         | -.060         | -.339        | -.519*                 | -.037| -.659**| .760**| .387 |
| Sepal size      | -.755**        | -          | .632*        | .329           | .160          | .734**        | .159                   | -.359| .756**| .699**| .399 |
| Ovary length    | -.855**        | -.175      | -            | -.209          | -.386         | .391         | .161                   | -.383| .563*| .369 |
| Ovary diameter  | -.175          | -          | -.209        | -.386          | .391          | .161         | -.383                  | .563*| .369 |
| Stigma length   | -.155          | -.805**    | -            | -.07           | .024          | .052         | .269                   | .391 |
| Stigma width    | -.353          | .198       | .106         | .217           | .021          | .391         | -                      |      |
| No. of ovules per ovary | -.107      | .349       | -.111        | -.915**        | .878**        | -            |                        |      |
| GA3             | -.567*         | -          | .286         | .309           |               |              |                        |      |
| ZR              | -.324          | -          | .071         |               |              |              |                        |      |
| IAA             | -.784**        | -          |              |               |              |              |                        |      |
| ABA             | -              | -          |              |               |              |              |                        |      |

ABA, abscisic acid; GA3, gibberellin A3; IAA, indoleacetic acid; ZR, zeatin riboside. The bold values is mentioned in the text.

*.01 < p < .05; **.001 < p < .01.
On the one hand, compared with the females at the single-flower level, male plants produced both significantly greater flower biomass at higher altitudes (1,600 and 1,700 m) and pollination organ (anthers) biomass percentage at all three altitudes (Figure 2). These results suggested that male *P. cathayana* plants at a higher altitude invest more mass in flower and allocate more biomass to the anthers than the females, and these may be beneficial to enhance pollen dispersal.

Also compared with females as a wind-pollinated plant, most and/or more male *P. cathayana* plants are distributed at higher-altitude sites (Wang, Xu, Li, Yang, & Yuan, 2011), which would benefit their exposure to higher wind speeds and longer pollen dispersal distances (Hesse & Pannell, 2011; Van Drunen & Dorken, 2014). Consequently, maintaining an adequate quantity of pollen could improve the rate of successful fertilization. Similar results from *Dombeya ciliata* showed that the flower size was larger in males than in females at a higher altitude (Humeau et al., 2000). On the other hand, the female plants had a greater single-flower biomass at 1,500 m altitude than at the two other higher altitudes (Figure 2a), indicating that a female plant at a lower altitude might invest more resources in the development of reproductive organs or have a higher biomass accumulation. Consistent with this phenomenon, male plants at the 1,500 m altitude had a relatively small amount of pollen than at a higher altitude (Figure 2a). As a result, a well-developed reproductive organ in a female flower would capture more pollen at a lower altitude. These results are consistent with the view that the sexual allocation in wind-pollinated plants involves an evolutionary strategy for promoting effective pollen dispersal and capture (Burd & Allen, 1988; Friedman & Barrett, 2009).

Meanwhile, it is noteworthy that *P. cathayana* plants at 1,600 m exhibited obvious changes in floral morphology when compared with their counterparts at either the lower 1,500 m or higher 1,700 m altitude (Table 1, Figure 1). At this middle-altitude site, the plants presented the smallest inflorescence length, female flower, and ovule number per ovary but the highest stigma biomass percentage and number of anthers per flower (Table 1, Figure 2b). This phenomenon suggested that other abiotic or biotic environment factors, such as population density or intraspecific competition, could also affect the morphological development of flowers. For example, studies have reported that population density could affect plant reproductive outputs and survival by increasing resource competition (Knight, 2003; Oleques & de Avila, 2014), and plants in high-density populations have fewer ovules per flower and smaller inflorescences (Weber & Kolb, 2011). At the same experimental sites, our previous work showed that...
**P. cathayana** plants could best reproduce and survive at the middle altitude of 1,600 m (Wang et al., 2011), resulting in a higher population density. We thus assumed that these distinctively morphological traits in flowers at this altitude might result from adaptive evolution to achieve maximal reproductive efficiency under high-intensity competition in the population.

Moreover, consistent with the observed morphological changes, the concentrations of ABA, GA₃, IAA, and ZR in the male and female flowers of **P. cathayana** plants varied with altitude, and different response patterns were presented between the two sexes (Figure 3). These results suggested that intrinsic relationships between floral morphology and endogenous phytohormones could lead to sex-specific morphological changes in response to altitude. As low-molecular mass-signaling substances in plants, phytohormones (e.g., ABA, GA₃, IAA, and ZR) are known to function in intercellular regulation in multicelled organisms (Sonnewald, 2013). Studies have reported that the levels of these endogenous hormones in flower tissues vary during flowering (Chen, Du, Zhao, & Zhou, 1996; Villacorta et al., 2008) and are related to the initiation and development of floral organs (Arrom & Munné-Bosch, 2012; Meilan, 1997). Consistent with these findings, morphological traits were closely related to phytohormone contents in both male and female flowers (e.g., between sepal size and ABA or between pedicel length and ZR, see Tables 2 and 3), which indirectly confirmed the role of phytohormones in flower development.

Sexual differences were also detected between phytohormones and biomass of flowers or a number of morphological traits (Tables 2 and 3, Figure 4). Our results indicated that phytohormones might be involved in the formation of male or female flowers, and changes in the concentration of these phytohormones would significantly affect the development of flower organs. Indeed, sexual organogenesis in dioecious plants is realized through the action of genes that use phytohormones as modulators to initiate the development of flower’s generative structures (Gerashchenkov & Rozhnova, 2013; Khryanin, 2002). Hence, sexual differences in the phytohormone concentrations as well as the relationships among phytohormones, morphological traits, and flower biomass were observed in this study (Figure 3, Tables 2 and 3). Similar results were reported in plant **Populus tomentosa** that genes related to phytohormone synthesis were significantly differentially expressed between the sexes and resulted in quite different endogenous GA, IAA, ABA, and CT (cytokinins) contents during all of floral development (Song et al., 2013).

In conclusion, as a general rule, our results demonstrated that morphological traits, biomass allocation, and phytohormone levels of the flowers of both male and female **P. cathayana** plants significantly differed with altitude. Along the investigated altitudinal gradient from 1,500 m to 1,700 m, the floral morphology of dioecious plants exhibited sex-specific differentiation, such as the largest stigma and greater pollen at the highest altitude, the smallest ovaries and higher number of anthers at the middle altitude, and the largest ovaries and lowest quantity of pollen at the low altitude. Additionally, the phytohormone levels in male and female flowers varied with altitude and were closely related to flower morphology, resulting in different morphological responses of flower organs to altitude between sexes. Our findings thus provided direct evidence of reproductive adaptation to altitude by dioecious plants.

**ACKNOWLEDGMENTS**

This research was supported by the National Natural Science Foundation of China (31170389 and 31370596) and the Innovative Team Foundation of the Sichuan Provincial Department of Education (14TD0015), China.
CONFLICT OF INTEREST

None declared.

REFERENCES

Arron, L., & Munné-Bosch, S. (2012). Hormonal changes during flower development in floral tissues of Lilium Planta. 236, 343–354.

Baonza, J., & Malo, J. E. (1997). Floral size variability of Cyntis scoparius along an altitudinal gradient. Lagasca, 19, 845–850.

Bodson, M., & Outlaw, W. H. (1985). Elevation in the sucrose content of the shoot apical meristem of Sinapis alba at floral evocation. Plant Physiology, 79, 420–424.

Brcko, A., Pencik, A., Magnus, V., Prebeg, T., Minaric, S., Antunovic, J., ... Salopek-Sondi, B. (2012). Endogenous auxin profile in the Christmas rose (Helleborus niger L) flower and fruit: Free and amide conjugated IAA. Journal of Plant Growth Regulation, 31, 63–78.

Burd, M., & Allen, T. F. H. (1988). Sexual allocation strategy in wind-pollinated plants. Evolution, 42, 403–407.

Carrera, E., Ruiz-Rivero, O., Peres, L. E. P., Atares, A., & Garcia-Martinez, J. L. (2012). Characterization of the procera tomato mutant shows novel functions of the SIDEALLA protein in the control of flower morphology, cell division and expansion, and the auxin-signaling pathway during fruit-set and development. Plant Physiology, 160, 1581–1596.

Chandler, J. (2011). The hormonal regulation of flower development. Journal of Plant Growth Regulation, 30, 242–254.

Chen, L., Dong, T., & Duan, B. (2014). Sex-specific carbon and nitrogen partitioning under N deposition in Populus cathayana. Trees, 28, 793–806.

Chen, J. G., Du, X. M., Zhao, H. Y., & Zhou, X. (1996). Fluctuation in levels of endogenous plant hormones in ovules of normal and mutant cotton during flowering and their relation to fiber development. Journal of Plant Growth Regulation, 15, 173–177.

Chen, J., Duan, B., Wang, M., Korpelainen, H., & Li, C. (2014). Intra- and inter-sexual competition of Populus cathayana under different watering regimes. Functional Ecology, 28, 124–136.

Chen, J., Henny, R. J., McConnell, D. B., & Caldwell, R. D. (2003). Gibberellic acid affects growth and flowering of Philodendron 'Black Cardinal'. Plant Growth Regulation, 41, 1–6.

Chen, M., Huang, Y., Liu, G., Qin, F., Yang, S., & Xu, X. (2016). Effects of enhanced UV-B radiation on morphology, physiology, biomass, leaf anatomy and ultrastructure in male and female mulberry (Morus alba) saplings. Environmental and Experimental Botany, 129, 85–93.

Corbesier, L., Prinsen, E., Jacqmard, A., Lejeune, P., Van Onckelen, H., Périlleux, C., & Bernier, G. (2003). Cytokinins in leaves, leaf exudate and shoot apical meristem of Arabidopsis thaliana during floral transition. Journal of Experimental Botany, 54, 2511–2517.

Delph, L. F. (1990). Sex-differential resource allocation patterns in the sub- dioecious shrub Hebe subalpina. Ecology, 71, 1342–1351.

Ding, S. F., Chen, W. S., Su, C. L., Du, B. S., Twitchin, B., & Bhaskar, V. K. (1999). Changes in free and conjugated indole-3-acetic acid during early stage of flower bud differentiation in Polianthes tuberosa. Plant Physiology and Biochemistry, 37, 161–165.

Domagalska, M. A., Sarnowska, E., Nagy, F., & Davis, S. J. (2010). Genetic analyses of interactions among gibberellin, abscisic acid, and brassinosteroids in the control of flowering time in Arabidopsis thaliana. PLoS ONE, 5, e14012.

Duan, Y., He, Y., & Liu, J. (2005). Reproductive ecology of the Qinghai-Tibet Plateau endemic Gentiana straminea (Gentianaceae), a hermaphrodite perennial characterized by herkogamy and dichogamy. Acta Oecologica, 27, 225–232.

Espírito-Santo, M. M., Madeira, B. G., Neves, F. S., Faria, M. L., Fagundes, M., & Fernandes, G. W. (2003). Sexual differences in reproductive phenotype and their consequences for the demography of Baccharis dracunculifolia (Asteraceae), a dioecious tropical shrub. Annals of Botany, 91, 13–19.

Fabbro, T., & Körner, C. (2004). Altitudinal differences in flower traits and reproductive allocation. Flora, 199, 70–81.

Friedman, J., & Barrett, S. C. (2009). Wind of change: New insights on the ecology and evolution of pollination and mating in wind-pollinated plants. Annals of Botany, 103, 1515–1527.

Galoeh, E., Czaplewskas, J., Burkacka-Łaukajtys, E., & Kopcewicz, J. (2002). Induction and stimulation of in vitro flowering of Pharbitis nil by cytokinin and gibberellin. Plant Growth Regulation, 37, 199–205.

Gerashchenko, G. A., & Rozhnova, N. A. (2013). The involvement of phytohormones in the plant sex regulation. Russian Journal of Plant Physiology, 60, 597–610.

Goto, N., & Pharis, R. P. (1999). Role of gibberellins in the development of floral organs of the gibberellin-deficient mutant, ga1-1, of Arabidopsis thaliana. Canadian Journal of Botany, 77, 944–954.

Gross, K., & Soule, J. (1981). Differences in biomass allocation to reproductive and vegetative structures of male and female plants of a dioecious, perennial herb, Silene alba (Miller) Krause. American Journal of Botany, 68, 801–807.

Guo, Y. F., Wang, Y. Q., & Weber, A. (2013). Floral ecology of Oreocharis acaulis (Gesneriaceae): An exceptional case of “preanthetic” protogyny combined with approach herkogamy. Flora, 208, 58–67.

Hautier, Y., Randin, C. F., Stöcklin, J., & Guisan, A. (2009). Changes in reproductive investment with altitude in an alpine plant. Journal of Plant Ecology, 2, 125–134.

Hesse, E., & Pannell, J. R. (2011). Density-dependent pollen limitation and reproductive assurance in a wind-pollinated herb with contrasting sexual systems. Journal of Ecology, 99, 1531–1539.

House, S. M. (1992). Population density and fruit set in 3 dioecious species in Australian tropical rain forest. Journal of Ecology, 8, 57–69.

Humeau, L., Pailler, T., & Thompson, J. D. (2000). Variation in gender and flower-size dimorphism in the dioecious tree Dombeya ciliata, an endemic to La Reunion Island. Biotropica, 32, 463–472.

Khryanin, V. N. (2002). Role of phytohormones in sex differentiation in plants. Russian Journal of Plant Physiology, 49, 545–551.

Knight, T. M. (2003). Floral density, pollen limitation, and reproductive success in Trillium grandiflorum. Oecologia, 137, 557–563.

Körner, C. (2007). The use of “altitude” in ecological research. Trends in Ecology & Evolution, 22, 569–574.

Korpelainen, H. (1992). Patterns of resource allocation in male and female plants of Rumex acetosa and R. acetosella. Oecologia, 89, 133–139.

Koti, S., Reddy, K. R., Reddy, V. R., Kakani, V. G., & Zhao, D. (2005). Interactive effects of carbon dioxide, temperature, and ultraviolet-B radiation on soybean (Glycine max L) flower and pollen morphology, pollen production, germination, and tube lengths. Journal of Experimental Botany, 56, 725–736.

Kudo, G., & Molau, U. (1999). Variations in reproductive traits at inflorescence and flower levels of an arctic legume, Astragalus alpinus L.: Comparisons between a subalpine and an alpine population. Plant Species Biology, 14, 181–191.

Laporte, M. M., & Delph, L. F. (1996). Sex-specific physiology and source-sink relations in the dioecious plant Silene latifolia. Oecologia, 106, 63–72.

Law, T. F., Lebel-Hardenack, S., & Grant, S. R. (2002). Silver enhances stamen development in female white campion (Silene latifolia [Caryophyllaceae]). American Journal of Botany, 89, 1014–1020.

Lei, Y., Chen, K., Jiang, H., Yu, L., & Duan, B. (2017). Contrasting responses in the growth and energy utilization properties of sympatric Populus and Salix to different altitudes: Implications for sexual dimorphism in Salicaceae. Physiologia Plantarum, 159(1), 30–41.

Li, J., Dong, T., Guo, Q., & Zhao, H. (2015). Populus deltoids females are more selective in nitrogen assimilation than males under different nitrogen forms supply. Trees, 29, 143–159.

Li, C., & Korpelainen, H. (2015). Transcriptomic regulatory network underlying morphological and physiological acclimation to nitrogen starvation and excess in poplar roots and leaves. Tree Physiology, 35, 1279–1282.
He J, Dong T, Huang K et al. Sex-specific floral morphology, biomass, and phytohormones associated with altitude in dioecious *Populus cathayana* populations. Ecol Evol 2017;7:3976–3986. https://doi.org/10.1002/ece3.2808