Potential roles of melatonin and ABA on apple dwarfing in semi-arid area of Xinjiang China

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ABSTRACT

Dwarfing is a typic breeding trait for mechanical strengthening and relatively high yield in modern apple orchards. Clarification of the mechanisms associated with dwarfing is important for use of molecular technology to breed apple. Herein, we identified four dwarfing apple germplasms in semi-arid area of Xinjiang, China. The internodal distance of these four germplasms were significantly shorter than non-dwarfing control. Their high melatonin (MT) contents are negatively associated with their malondialdehyde (MDA) levels and oxidative damage. In addition, among the detected hormones including auxin (IAA), gibberellin (GA), brassinolide (BR), zeatin-ribose (ZR), and abscisic acid (ABA), only ABA and ZR levels were in good correlation with the dwarfing phenotype. The qPCR results showed that the expression of melatonin synthetic enzyme genes MdASMT1 and MdSNAT5, ABA synthetic enzyme gene MdAAO3 and degradative gene MdCYP707A, ZR synthetic enzyme gene MdIPT5 all correlated well with the enhanced levels of MT, ABA and the reduced level of of ZR in the dwarfing germplasms. Furthermore, the significantly higher expression of ABA marker genes (MdRD22 and MdRD29) and the lower expression of ZR marker genes (MdRR1 and MdRR2) in all the four dwarf germplasms were consistent with the ABA and ZR levels. Considering the yearly long-term drought occurring in Xinjiang, China, it seems that dwarfing with high contents of MT and ABA may be a good strategy for these germplasms to survive against drought stress. This trait of dwarfing may also benefit apple production and breeding in this semi-arid area.

INTRODUCTION

Apple (Malus domestica) is one of the top four fruits with the largest planting areas globally (Cohen et al., 2007). For apples, dwarfing is the major breeding objective to
shorten juvenile period, increase yield and maintain mechanical strengthening for the high yield (Bulley et al., 2005; Smolka et al., 2010). Although years of crossing and selection to breed apple cultivar for ideal tree architecture, the elite of spur-type apple cultivar combined with other good horticultural traits is still a goal for apple breeders (Zheng et al., 2018). Many efforts have been made to uncover dwarfing mechanism, however, the complex underlying interactions between plants and environments remain unclear (Cohen et al., 2007).

The regulatory effects of hormones, cytokines (CKs), auxin (IAA), brassinosteroids (BRs), and gibberellins (GAs) on dwarfing have been extensively studied (Waldie & Leyser, 2018; Chen et al., 2019). Many genes involved in GA, IAA and BR synthesis, degradation, transportation and signaling have been identified (Davière et al., 2014; Ma et al., 2016). In apple, these hormones also play critical roles in dwarfing (Tong et al., 2014; Unterholzner et al., 2015; Guo et al., 2020). However, different from herbal plants, which flower earlier under severe stresses to end the life cycle, apple trees have developed more complex dwarfing mechanisms (Hatayama & Takeno, 2003; Xu et al., 2014; Minh-Thu et al., 2018) to better suit the long lasting drought stresses. Apple trees usually retard their normal-growth to enhance defense, which leads to a stress-defensive dwarfing phenotype (Liu et al., 2021).

Drought stress is increasing with the global warming, which severely affects apple yield. To adapt the upcoming global warming, it is important to breed dwarfing apple to balance the growth and drought tolerance to maintain its yield. In recent years, melatonin has been frequently reported to exhibit important roles involved in both drought tolerance and plant development (Zhang et al., 2015; Sun et al., 2021; Ahmad et al., 2021). The mechanism may attributed to its antioxidative activity. For example, long-term drought stress in the main apple production area of China, unavoidably results in secondary oxidative damage indicated by increased Reactive Oxygen Species (ROS) and malondialdehyde (MDA) level (Zuo et al., 2014; Huang et al., 2020). The ROS burst induces the biosynthesis of melatonin (MT), a well-known strong ROS scavenger in response to drought stress (Zuo et al., 2014; Wang et al., 2017). When apple trees are long-term-stressed, the induced MT might mediate a stress-defensive dwarfing. However, more evidence needs to be explored.

Melatonin is an amphiphilic molecule (Tan et al., 2012), which can freely cross cell membranes to any organelles including the chloroplast, nucleus, and mitochondria. MT metabolites can also synergistically scavenge ROS, resulting in amplified antioxidant capacity (Sun et al., 2021; Siddiqui et al., 2019). The biosynthesis pathway of melatonin in plants has been well-documented. An amino acid tryptophan is the precursor of melatonin synthesis. The tryptophan is catalyzed successively by tryptophan decarboxylase (TDC), tryptopine-5-hydroxylase (T5H), serotonin-N-acetyltransferase (SNAT) and N-acetylserotonin methoxy transferase (ASMT) to form melatonin, among which SNAT and ASMT are the rate-limiting enzymes in plant (Tan et al., 2012; Zhang et al., 2019; Sun et al., 2021).

ABA, a so-called stress hormone, functions in promoting stoma closure to reduce water lost through transpiration regulation (Shu et al., 2016; Long et al., 2019; Dou et al., 2021).
Therefore, ABA inhibits plant growth via down-regulating photosynthesis (Jiang et al., 2021). ABA level is induced by drought stress. Its metabolic pathway has been well studied (Shu et al., 2016; Ren et al., 2018; Pang et al., 2019). Shi et al. (2021) reported that the drought-induced OsAAO3 regulates rice growth and development. The dehydration responsive CYP707A, encoding a key enzyme in ABA catabolism limits the plant growth for the purpose to survive under drought stress via reducing the expression of genes such as RD22 and RD29 (Matilla, Carrillo-Barral & Rodríguez-Gacio, 2015; Yu et al., 2020).

Cytokinin which is also in response to drought resistance via triggering RRI and RR2-involved signaling (Nishiyama et al., 2011), is reported to negatively regulate apple dwarfing via regulating cell division. The Arabidopsis mutants of IPT5b gene, encoding for a ZR synthetase, exhibit reduced level of the endogenous active forms and severe dwarfism (Tokunaga et al., 2012). In apple, the low expression of IPT5b leads to poor zeatin biosynthesis and dwarfing (Feng et al., 2017). The over-expression ZR degradation gene of CKX1 in transgenic Arabidopsis and tobaccos decreased ZR level, enhanced stress tolerance and retarded shoot growth (Werner et al., 2010; Macková et al., 2013).

Herein, we have collected four apple dwarfing germplasms with shortened nodule length in the semi-arid area of Xinjiang, China. We have not found a good correlation between these dwarfing germplasms with the well-known hormones involved in dwarfing, such as GA, BR and auxin. However, the contents of stress induced hormone ABA and melatonin exhibit positive correlation with these four dwarfing apple germplasms. In addition, ZR content is also decreased in all the germplasms and this suggests its involvement in dwarfing. Considering a typical long-term drought season in this area, apples grown in this area will experience high level oxidative tissue damage and they will increase the levels of stress-responsive molecules, MT and ABA by upregulating the expression of their synthetic and signaling genes, it infers a stress-defensive dwarfing mechanism mediated by melatonin and ABA, as an important strategy to survive drought stress in the semi-arid area of Xinjiang.

MATERIALS AND METHODS

Plant materials
The leaves and shoots were collected from the annual branches of the four dwarfing apple germplasms (Dwarf1-4) and a non-dwarfing germplasm (GB2), all the germplasms grafted onto 8-year-old Xinjiang Wild Apple (Malus sieversii (Lebed.) Roem.) in northern Xinjiang, China (82°51’46″E and 43°15’49″N, altitude 1,365 m) which belongs to semi-arid area, with an annual precipitation of proximately 200 mm (Chen et al., 2015). The germplasms in this semi-arid area are under long-term drought stress. The internodal distance of four spur-type apple germplasms were measured, with the non-dwarfing germplasm ‘GB2’ used as control, and the second node of the annual branch was the basal node for node counting. The leaf-samples were quickly frozen with liquid nitrogen, and then were kept at −80 °C till to use. The cell number of stem cortical parenchyma cells were observed using paraffin sections made by Servicebio Co. (Wuhan, Hubei, China) and scanned by CaseViewer2.3. Three different regions (250 × 75 μm) of the middle of internodes were selected for observation.
Detection of malondialdehyde (MDA)
A total of 1 g leaves from the five apple germplasms (GB2, Dwarf1-4) were homogenized for MDA detection, respectively according to Zhao et al. (2013). Each experiment was independently repeated three times.

Extraction and analysis of phytohormones of abscisic acid (ABA), indole-3-acetic acid (IAA), Gibberellin A3 (GA3), trans-zeatin-riboside (ZR) and brassinosteroids (BR)
A total of 0.5 g leaves from the five apple germplasms (GB2, Dwarf1-4) were used for extraction of ABA, IAA, GA3 and ZR, as described by Yang et al. (2001), and BR as described by Xin et al. (2013), respectively. ELISA analysis of ABA, IAA, GA3, ZR and BR were carried out according to Yang et al. (2001). Each experiment was independently repeated three times.

Extraction and analysis of melatonin
A total of 0.4 g leaves from the five apple germplasms (GB2, Dwarf1-4) were ground to a fine powder with liquid nitrogen for melatonin extraction and analysis by HPLC as described by Zhao et al. (2013), respectively. Each experiment was independently repeated three times.

RNA extraction and qRT-PCR analysis
To detect the expression of melatonin synthetic enzyme genes (MdASMT1 and MdSNAT5), ABA synthetic enzyme gene (MdAAO3), degradative enzyme gene (MdCYP707A) and ABA signaling genes (MdRD22 and MdRD29), ZR synthetic enzyme gene (MdIPT5) and ZR signaling genes (MdRR1 and MdRR2) by qPCR analysis, total RNA was isolated from 0.1 g leaves of the five apple germplasms (GB2, Dwarf1-4), respectively with EASY spin Plant RNA Kit (Aidlab Biotechnologies, Beijing, China). cDNA was obtained by reverse transcription, using TUREscript RT MasterMix (Aidlab Biotechnologies, Beijing, China) following the protocol, as template for qRT-PCR. Each experiment was independently repeated three times.

The specific primers of MdAAO3, MdCYP707A, MdIPT5, MdRR1, MdRR2, MdRD22 and MdRD29 genes were designed by Primer 5 software according to the coding sequences from GDR (https://www.rosaceae.org/species/malus/all), and the specific primers of MdASMT1 and MdSNAT5 genes were used as described by Zuo et al. (2014) and Wang et al. (2017), respectively. The Actin gene was used as the internal control. All the primers are shown in Table S2. qRT-PCR was performed with TB Green® Premix Ex Taq (Takara, Dalian, China) on an ABI 7500 real-time PCR machine (ABI, Carlsbad, CA, USA) according to Zheng et al. (2018). Each experiment was independently repeated three times.

Statistical analysis
The data were expressed as mean ± SEM and analyzed by one-way analysis of variance (ANOVA) followed by Fisher’s Least Significant Difference (LSD) test. p < 0.05 was
considered statistically significant. The SPSS 25.0 software (IBM, Armonk, NY, USA) was used for statistical computations.

**RESULTS**

**The variation of nodular distance between non-dwarfing and dwarfing apple germplasms**

To investigate the underlying molecular mechanism of dwarfing in apple, we selected four dwarfing apple germplasms and a non-dwarfing apple germplasm used as control. Compared with GB2, the annual branches of the dwarfing apple germplasms exhibited significantly short nodular length (Fig. 1A). As the branch growing, differences of the nodular distance between the non-dwarfing and dwarfing apple germplasms became more apparent. The greatest differences were observed between nodes 3 and 5. At the third internode, GB2 had an internodal length of 2.97 cm, whereas that of dwarfing apple germplasms (Dwarf1-4) were 1.37, 1.53, 1.87 and 1.50 cm, respectively (Fig. 1B), indicating that dwarfing is caused by the shortened internodes.

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**Figure 1** The phenotypes of the four dwarfing and control apple germplasms. (A) Shoot lengths of the four dwarfing and control apple germplasms. Bar, 3 cm. (B) Internodal distance of the four dwarfing and control apple germplasms. (C) Paraffin sections of shoot of the four dwarfing and control apple germplasms. (D) Statistical chart of cell number per unit area. Data are means ± SD of triplicate studies. Different lower-case letters indicate significant differences among germplasms according to Fisher’s LSD test (p < 0.05). 

Full-size [DOI: 10.7717/peerj.13008/fig-1]
To further determine whether the decrease in cell number or the inhibition of cell elongation is the cause of the dwarfi ng, we analyzed the longitudinal sections of stems with an optical microscope (Figs. 1C, 1D). Longitudinal sections of stems between the normal and dwarfi ng apple germplasms revealed clear difference in the cell number of cortical parenchyma cells. The cell number of dwarfi ng apple germplasms were signifi cantly less than GB2. Therefore, shorter internodal lengths of dwarfi ng apple germplasms is the result of fewer cells.

The annual branches of dwarfi ng apple germplasms have higher MT and MDA levels than that of GB2

We measured the content of MT and MDA in GB2 and dwarfi ng germplasms. The MT contents in all the four dwarfi ng germplasms (Dwarf1-4) were signifi cantly higher than that of non-dwarfi ng germplasms (Fig. 2A). Their contents were at least 1.7 times higher than that of the non-dwarfi ng germplasm, and the highest MT level could reach 196.827 ng·g\(^{-1}\), which is about three times higher than that of the GB2.

The relative expressions of *MdASMT1* and *MdSNAT5* were also signifi cantly higher in the four dwarfi ng germplasms than that in the non-dwarfi ng germplasm. As shown in Fig. 2B, the relative expression of *MdASMT1* in four dwarfi ng germplasms were 1.66, 1.93, 2.87 and 2.07 times of higher that in GB2, respectively. The relative expression of *MdSNAT5*, a rate-limiting enzyme for melatonin synthesis in mitochondria, showed a similar trend, and the level in four dwarfi ng germplasms were 1.64, 1.95, 1.43 and 1.40 times higher than that in GB2, respectively (Fig. 2B).

In addition, the content of MDA, indicating the degree of lipid oxidative damage, was also detected. Similarly, the MDA content in dwarfi ng germplasms were significantly higher than that of GB2. (A) The melatonin level of the four dwarfi ng apple germplasms and control. (B) Relative expression level of *MdASMT1* and *MdSNAT5* in the four dwarfi ng and control apple germplasms. (C) The MDA content in the four dwarfi ng and control apple germplasms. Data are means ± SD of triplicate studies. Different lower-case letters indicate signifi cant differences among germplasms according to Fisher’s LSD test (\(p<0.05\)).

![Image](DOI: 10.7717/peerj.13008/fig-2)
higher than that of GB2 (0.00568 \mu\text{mol} \cdot \text{g}^{-1} \text{ FW}). Among the dwarfing germplasms, the MDA content in Dwarf1 (0.01255 \mu\text{mol} \cdot \text{g}^{-1} \text{ FW}) was the highest, which was 2.2 times higher than that of non-dwarfing germplasm, and the MDA content in Dwarf4 (0.00669 \mu\text{mol} \cdot \text{g}^{-1} \text{ FW}) was the lowest, which was 1.2 times higher than that of GB2 (Fig. 2C).

The ABA level is higher in dwarfing apple germplasms than that of GB2
Likewise, the ABA content in all the four dwarfing apple germplasms were significantly higher than that in GB2. The ABA content in Dwarf2-4 were 49.949, 54.766, and 56.410 ng·g$^{-1}$, respectively, while it was only 39.174 ng·g$^{-1}$ in the GB2. Although the ABA content in Dwarf1 (41.848 ng·g$^{-1}$) was lower than the other three dwarfing apple germplasms, it was still higher than that in GB2 (Fig. 3A). The expression of MdRD22 and MdRD29 (ABA inducible genes involved in ABA signaling) in four dwarfing apple germplasms were significantly higher than that in GB2 (Fig. 3C).

Furthermore, the relative expression of key genes involved in ABA biosynthesis and degradation were also analyzed in non-dwarfing and dwarfing apple germplasms. Compared with the GB2, the MdAAO3 (a key gene in ABA biosynthesis) level in four dwarfing apple germplasms were significantly high, while the MdCYP707A (a key gene in ABA degradation) level was significantly lower than that of GB2 (Fig. 3B), which is another mechanism to lead to the enhanced ABA concentrations.

The ZR level is lower in dwarfing apple germplasms than that in GB2
Plant height is primarily regulated by hormones, including GA, IAA, ZR and BR. We measured the contents of GA, IAA, ZR and BR in dwarfing and GB2 germplasms. Between them, the concentrations of IAA, GA3 and BR had no significan differences (Table S1). By contrast, the concentration of ZR was significantly lower in dwarfing
germplasms than that of GB2. The ZR concentrations in the dwarfishing apple germplasms (Dwarf1-4) were 3.071, 3.486, 7.504 and 6.750 ng·g⁻¹, respectively which were only 28.3%, 32.1%, 69.1% and 62.2% of level of GB2 (10.857 ng·g⁻¹) (Fig. 4A). The expression of \textit{MdRR1} and \textit{MdRR2} (ZR up-regulated genes) in four dwarfishing apple germplasms were significantly lower than that in GB2 (Fig. 4B). We analyzed the expression of \textit{MdIPT5} (a key gene in ZR biosynthesis), the relative expression of \textit{MdIPT5} in the four dwarfishing germplasms were significantly lower than that in GB2, and \textit{MdIPT5} expression level in Dwarf3 was the lowest (Fig. 4B).

**DISCUSSION**

Dwarfishing is the major breeding trait targeted by researchers to increase apple yield and improve mechanical strengthening of the apple tree in the modern orchard (Bulley et al., 2005; Smolka et al., 2010; Yang et al., 2012). Due to the complex of the dwarfishing process involved in the perennial woody apple tree, extensive studies have been carried out by researchers at the temptation to uncover its underlining mechanisms. For this purpose, we have collected four dwarfishing apple germplasms in the semi-arid area of Xinjiang, China to explore the mechanisms of dwarfishing process. Unexpected, the levels of all well-known phytohormones involved in plant dwarfishing including IAA, GA3, and BR (Zheng et al., 2017; Zheng et al., 2018) did not show any correlation with the phenotypes of the four dwarfishing apple germplasms. In contrast, the levels of stress-responsive molecules, MT and ABA, significantly increased in all of the dwarfishing apple germplasms (Figs. 2A, 3A),
inferring the dwarfting phenotype may be an evolutionary adaptation responding to the drought stress (Fig. 5).

The elevated level of MDA in the dwarfting apple germplasms indicates the significantly increased oxidative damage, possibly induced by long-term drought stress in semi-arid area of Xinjiang (Fig. 2C). This oxidative damage caused by the over-produced ROS, which in turn, triggers the synthesis of ROS scavenger, MT (Zheng et al., 2017) and the enhanced MT level has been detected in all dwarfting apple germplasms. The MT level actually is determined by both its synthesis and consumption. The expression of MT synthetic enzyme genes MdASMT1 and MdSNAT5 are significantly upregulated in all the dwarfting apple germplasms (Fig. 2B). The expression pattern of these genes correlate well with the MT level, confirming that MT synthetic genes contribute great to the MT content in the dwarfting apple germplasms. There are some inconsistent reports related to the role of MT in plant dwarfting. For example, the reduced melatonin leads to dwarfting transgenic rice (Hwang & Back, 2018) while the enhanced melatonin level has detected in transgenic tomato dwarfting lines (Wang et al., 2014). These inconsistences suggest the different roles of melatonin between monocotyledon and dicotyledon plants. In the perennial woody apple tree, high melatonin level is first reported to associated with its dwarfting phynotype. As a ROS scavenger, melatonin level is highly regulated by environmental alterations and therefore it has a high chance to manipulate the dwarfting process associated with environmental changes.

The increased ABA level in the dwarfting apple germplasms may also result from the long-seasonal drought stress (Yoshida et al., 2021). The expression of MdAAO3 (ABA synthetase gene), MdRD22 and MdRD29 (ABA signaling genes) are significantly upregulated while ABA degradation gene MdCYP707A, is downregulated in all of the dwarfting apple germplasms. These suggest that the increased ABA level is attributed to both its enhanced synthesis and reduced degradation (Fig. 3B).

Role of the increased ABA in apple dwarfting germplasms can be logically explained, i.e., the long-term drought stress induces ABA accumulation and therefore promotes stomatal closure. This leads to decrease photosynthesis by promoting ABA-mediated chlorophyll degradation (Asad et al., 2019) and reducing carbon dioxide in-take and therefore retards apple tree growth. Similarly, the pear trees which accumulated with ABA also show dwarfting phenotype after knockout of PbPAT14 (Pang et al., 2019). The similar phenomenon of dwarfting phenotype due to ABA accumulation was also found in Gerbera hybridra and Arabidopsis thaliana (Yang, Worley & Udvardi, 2015; Ren et al., 2018). On the contrary, there are also reports on the positive role of ABA in regulating apple tree growth, that is, the reduced ABA and BR synthesis are found in transgenic dwarfting apple trees which overexpress MdNAC1 (Jia et al., 2018). But it is not clear whether this dwarfting phenotype is mainly caused by the reduced ABA or BR. Li et al. (2015), have reported that the exogenous melatonin pre-application reduces drought-inducible ABA biosynthesis. It is likely that the exogenous melatonin decreases ROS burst, thus weakens stress signal and ABA biosynthesis. In our research, the long-term environmental drought stress induced both ABA and melatonin biosynthesis. Apple plants may adapt the stress for survival at the cost of retarded growth. The environmental stress can be converted into
hormone signals including both ABA and melatonin, which triggered the ZT-regulated growth retard.

Kamboj et al. (1999) have observed that ZR, a zeatin precursor, is found to negatively regulate apple dwarfing. Our results showed that ZR content of all the apple dwarfasting germplasms were significantly lower than control plant (Fig. 4A). The ZR content correlates well with the expression of its synthetase gene *MdIPT5* and the signaling genes *MdRR1* and *MdRR2*, which were lower in all of the apple dwarfasting germplasms than control plant (Fig. 4B). This suggests that ZR biosynthesis is probably also involved with the apple dwarfasting. Moreover, it is well documented that growth and development of plant are mainly influenced by CK signaling pathways (Kieber & Schaller, 2018; Mao et al., 2019; Li et al., 2021). In Arabidopsis, overexpression of *RR1* promotes stem growth (Hwang & Sheen, 2001). In the current study, the result indicates that high level of *MdIPT5* expression in non-dwarfasting germplasm increases zeatin level to promote cell division and internode elongation, while low level of *MdIPT5* expression in the dwarfasting germplasms results in defective zeatin and further reduced internode length. To reduce the ZR content might be an important strategy for apple to survive long-term drought stress via decreasing cell division and energy cost, retarding growth and promoting dwarfasting (Ma et al., 2020; Zong et al., 2021).

Whether this stress-induced dwarfasting phenotype can pass to the next generation or maintain in the propagated apple tree is currently unknown. However, many environment-induced methylations for altered gene expression, selected in the evolution, is inheritable (Marfil et al., 2006; Keyte et al., 2006). This stress-tolerance dwarfasting
mechanism may also be a heritable strategy. But further research is needed to draw a conclusion.

CONCLUSIONS

In summary, we have identified a potential molecular mechanism of apple dwarfing to adapt the drought in the semi-arid areas of Xinjiang, China. This mechanism is involved in not only the enhanced levels of melatonin and ABA, but also reduced content of ZR in different apple dwarfing germplasms. These hormone alterations are primarily caused by the long-lasting drought season which the apple dwarfing plants face to in Xinjiang, China. This is the first report to show a drought-associated dwarfing mechanism in apple trees. This dwarfing process is involved in altered gene expressions of phytohormones and antioxidant (ROS scavenger). These observations provide novel dwarfing mechanisms for apple breeding and production. In addition, this drought stress induced dwarfing phenotype in apple tree may be inheritable and to indentify this is the goal of our future study.

ADDITIONAL INFORMATION AND DECLARATIONS

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Competing Interests

The authors declare that they have no competing interests.

Author Contributions

- Tianci Yan performed the experiments, analyzed the data, prepared figures and/or tables, and approved the final draft.
- Chuang Mei performed the experiments, authored or reviewed drafts of the paper, and approved the final draft.
Handong Song performed the experiments, analyzed the data, prepared figures and/or tables, and approved the final draft.

Dongqian Shan analyzed the data, authored or reviewed drafts of the paper, and approved the final draft.

Yanzhao Sun analyzed the data, prepared figures and/or tables, and approved the final draft.

Zehui Hu performed the experiments, prepared figures and/or tables, and approved the final draft.

Lin Wang analyzed the data, prepared figures and/or tables, and approved the final draft.

Tong Zhang analyzed the data, prepared figures and/or tables, and approved the final draft.

Jixun Wang analyzed the data, authored or reviewed drafts of the paper, and approved the final draft.

Jin Kong conceived and designed the experiments, authored or reviewed drafts of the paper, and approved the final draft.

Data Availability
The following information was supplied regarding data availability:

The raw measurements are available in the Supplemental Files.

Supplemental Information
Supplemental information for this article can be found online at http://dx.doi.org/10.7717/peerj.13008#supplemental-information.

REFERENCES

Ahmad S, Muhammad I, Wang GY, Zeeshan M, Yang L, Ali I, Zhou XB. 2021. Ameliorative effect of melatonin improves drought tolerance by regulating growth, photosynthetic traits and leaf ultrastructure of maize seedlings. *BMC Plant Biology* 21(1):368 DOI 10.1186/s12870-021-03160-w.

Asad MAU, Zakari SA, Zhao Q, Zhou L, Ye Y, Cheng F. 2019. Abiotic stresses intervene with ABA signaling to induce destructive metabolic pathways leading to death: premature leaf senescence in plants. *International Journal of Molecular Sciences* 20(2):256 DOI 10.3390/ijms20020256.

Bulley SM, Wilson FM, Hedden P, Phillips AL, Croker SJ, James DJ. 2005. Modification of gibberellin biosynthesis in the grafted apple scion allows control of tree height independent of the rootstock. *Plant Biotechnology Journal* 3(2):215–223 DOI 10.1111/j.1467-7652.2005.00119.x.

Chen C, Zhao K, Ablimitijan A, Li N, Yu B. 2015. Temporal and spatial distributions of hourly rain intensity under the warm background in Xinjiang. *Arid Land Geography* 38(4):692–702.

Chen L, Zhao J, Song J, Jameson PE. 2019. Cytokinin dehydrogenase: a genetic target for yield improvement in wheat. *Plant Biotechnology Journal* 18(3):614–630 DOI 10.1111/pbi.13305.

Cohen S, Naor A, Bennink J, Grava A, Tyree M. 2007. Hydraulic resistance components of mature apple trees on rootstocks of different vigours. *Journal of Experimental Botany* 58(15–16):4213–4224 DOI 10.1093/jxb/erm281.
Davière J, Wild M, Regnault T, Baumberger N, Eisler H, Genschik P, Achard P. 2014. Class I TCP-DELLA interactions in inflorescence shoot apex determine plant height. *Current Biology* **24**(16):1923–1928 DOI 10.1016/j.cub.2014.07.012.

Dou L, He K, Peng J, Wang X, Mao T. 2021. The E3 ligase MREL57 modulates microtubule stability and stomatal closure in response to ABA. *Nature Communications* **12**(1):2181 DOI 10.1038/s41467-021-22455-y.

Feng Y, Zhang X, Wu T, Xu X, Han Z, Wang Y. 2017. Methylation effect on *IPT5b* gene expression determines cytokinin biosynthesis in apple rootstock. *Biochemical and Biophysical Research Communications* **482**(4):604–609 DOI 10.1016/j.bbrc.2016.11.080.

Guo W, Chen L, Herrera-Estrella L, Cao D, Tran LP. 2020. Altering plant architecture to improve performance and resistance. *Trends in Plant Science* **25**(11):1154–1170 DOI 10.1016/j.tplants.2020.05.009.

Hatayama T, Takeno K. 2003. The metabolic pathway of salicylic acid rather than of chlorogenic acid is involved in the stress-induced flowering of *Pharbitis nil*. *Journal of Plant Physiology* **160**(5):461–467 DOI 10.1078/0176-1617-01041.

Huang D, Ma M, Wang Q, Zhang M, Jing G, Li C, Ma F. 2020. Arbuscular mycorrhizal fungi enhanced drought resistance in apple by regulating genes in the MAPK pathway. *Plant Physiology and Biochemistry* **149**:245–255 DOI 10.1016/j.plaphy.2020.02.020.

Hwang OJ, Back K. 2018. Melatonin is involved in skotomorphogenesis by regulating brassinosteroid biosynthesis in rice plants. *Journal of Pineal Research* **65**(2):e12495 DOI 10.1111/jpi.12495.

Hwang I, Sheen J. 2001. Two-component circuitry in arabidopsis cytokinin signal transduction. *Nature* **413**(6854):383–389 DOI 10.1038/35096500.

Jiang Y, Tong S, Chen N, Liu B, Bai Q, Chen Y, Bi H, Zhang Z, Lou S, Tang H, Liu J, Ma T, Liu H. 2021. The PalWRKY77 transcription factor negatively regulates salt tolerance and abscisic acid signaling in Populus. *Plant Journal* **105**(5):1258–1273 DOI 10.1111/tpj.15109.

Jia D, Gong X, Li M, Li C, Sun T, Ma F. 2018. Overexpression of a novel apple NAC transcription factor gene, MdNAC1, confers the dwarf phenotype in transgenic apple (*Malus domestica*). *Genes (Basel)* **9**(5):229 DOI 10.3390/genes9050229.

Keyte AL, Percifield R, Liu B, Wendel JF. 2006. Intraspecific DNA methylation polymorphism in cotton (*Gossypium hirsutum* L.). *Journal of Heredity* **97**(5):444–450 DOI 10.1093/jhered/esl023.

Kieber J, Schaller G. 2018. Cytokinin signaling in plant development. *Development* **145**(4):845 DOI 10.1242/dev.149344.

Kamboj JS, Blake PS, Quinlan JD, Baker DA. 1999. Identification and quantitation by gc-ms of zeatin and zeatin riboside in xylem sap from rootstock and scion of grafted apple trees. *Plant Growth Regulation* **28**(3):199–205 DOI 10.1023/A:1006292309765.

Li S-M, Zheng H-X, Zhang X-S, Sui N. 2021. Cytokinins as central regulators during plant growth and stress response. *Plant Cell Reports* **40**(2):271–282 DOI 10.1007/s00299-020-02612-1.

Liu X, Zhao C, Gao Y, Xu Y, Wang S, Li C, Xie Y, Chen P, Yang P, Yuan L, Wang X, Huang L, Ma F, Feng H, Guan Q. 2021. A multifaceted module of BRI1 ethylmethane sulfonate supressor1 (BES1)-MYB88 in growth and stress tolerance of apple. *Plant Physiology* **185**(4):1903–1923 DOI 10.1093/plphys/kiab116.

Li C, Tan D-X, Liang D, Chang C, Jia D, Ma F. 2015. Melatonin mediates the regulation of ABA metabolism, free-radical scavenging, and stomatal behaviour in two Malus species under drought stress. *Journal of Experimental Botany* **66**(3):669–680 DOI 10.1093/jxb/eru476.
Long H, Zheng Z, Zhang Y, Xing P, Wan X, Zheng Y, Li L. 2019. An abscisic acid (ABA) homeostasis regulated by its production, catabolism and transport in peanut leaves in response to drought stress. PLOS ONE 14(6):e213963 DOI 10.1371/journal.pone.0213963.

Ma X, Yu J, Zhuang L, Shi Y, Meyer W, Huang B. 2020. Differential regulatory pathways associated with drought-inhibition and post-drought recuperation of rhizome development in perennial grass. Annals of Botany 126(3):481–497 DOI 10.1093/aob/mcaa099.

Macková H, Hronková M, Dobrá J, Turečková V, Novák O, Lubovská Z, Motyka V, Haisel D, Hájek T, Prášil IT, Gaudinová A, Štorchová H, Ge E, Werner T, Schmülling T, Vanková RV. 2013. Enhanced drought and heat stress tolerance of tobacco plants with ectopically enhanced cytokinin oxidase/dehydrogenase gene expression. Journal of Experimental Botany 64(10):2805–2815 DOI 10.1093/jxb/ert131.

Ma Y, Xue H, Zhang L, Zhang F, Ou C, Wang F, Zhang Z. 2016. Involvement of auxin and brassinosteroid in dwarfi sm of autotetraploid apple (Malus × domestica). Scientific Reports 6(1):452 DOI 10.1038/srep26719.

Marfil CF, Masuelli RW, Davison J, Comai L. 2006. Genomic instability in Solanum tuberosum × Solanum kurtzianum interspecific hybrids. Genome 49(2):104–113 DOI 10.1139/g05-088.

Minh-Thu PT, Kim JS, Chae S, Jun KM, Lee GS, Kim DE, Cheong JJ, Song SI, Nahm BH, Kim YK. 2018. A WUSCHEL homeobox transcription factor, OsWOX13, enhances drought tolerance and triggers early flowering in rice. Molecules and Cells 41(8):781–798 DOI 10.14348/molcells.2018.0203.

Matilla AJ, Carrillo-Barral N, Rodríguez-Gacio M. 2015. An update on the role of NCED and CYP707A ABA metabolism genes in seed dormancy induction and the response to after-ripening and nitrate. Journal of Plant Growth Regulation 34(2):274–293 DOI 10.1007/s00344-014-9464-7.

Mao J, Zhang D, Meng Y, Li K, Wang H, Han M. 2019. Inhibition of adventitious root development in apple rootstocks by cytokinin is based on its suppression of adventitious root primordia formation. Physiologia Plantarum 166(2):663–676 DOI 10.1111/ppl.12817.

Nishiyama R, Watanabe Y, Fujita Y, Le DT, Kojima M, Werner T, Vankova R, Yamaguchi-Shinozaki K, Shinozaki K, Kakimoto T, Sakakibara H, Schmülling T, Tran LP. 2011. Analysis of cytokinin mutants and regulation of cytokinin metabolic genes reveals important regulatory roles of cytokinins in drought, salt and abscisic acid responses, and abscisic acid biosynthesis. The Plant Cell 23(6):2169–2183 DOI 10.1105/tpc.111.087395.

Pang H, Yan Q, Zhao S, He F, Xu J, Qi B, Zhang Y. 2019. Knockout of the S-acyltransferase gene, PbPAT14, confers the dwarf yellowing phenotype in first generation pear by ABA accumulation. International Journal of Molecular Sciences 20(24):6347 DOI 10.3390/ijms20246347.

Ren G, Li L, Huang Y, Wang Y, Zhang W, Zheng R, Zhong C, Wang X. 2018. GhWIP2, a WIP zinc finger protein, suppresses cell expansion in Gerbera hybrida by mediating crosstalk between gibberellin, abscisic acid, and auxin. The New Phytologist 219(2):728–742 DOI 10.1111/nph.15175.

Shi X, Tian Q, Deng P, Zhang W, Jing W. 2021. The rice aldehyde oxidase OsAO3 gene regulates plant growth, grain yield, and drought tolerance by participating in ABA biosynthesis.
Biochemical and Biophysical Research Communications 548:189–195
DOI 10.1016/j.bbrc.2021.02.047.

Siddiqui MH, Alamri S, Al-Khaishany MY, Khan MN, Al-Amri A, Ali HM, Alaraidh IA, Alsahl AA. 2019. Exogenous melatonin counteracts NaCl-induced damage by regulating the antioxidant system, proline and carbohydrates metabolism in tomato seedlings. International Journal of Molecular Sciences 20(2):353 DOI 10.3390/ijms20020353.

Sun C, Liu L, Wang L, Li B, Jin C, Lin X. 2021. Melatonin: a master regulator of plant development and stress responses. Journal of Integrative Plant Biology 63(1):126–145 DOI 10.1111/jipb.12993.

Tan DX, Hardeland R, Manchester LC, Korkmaz A, Ma S, Rosales-Corral S, Reiter RJ. 2012. Functional roles of melatonin in plants, and perspectives in nutritional and agricultural science. Journal of Experimental Botany 63(2):577–597 DOI 10.1093/jxb/err256.

Tong H, Xiao Y, Liu D, Gao S, Liu L, Yin Y, Jin Y, Qian Q, Chu C. 2014. brassinosteroid regulates cell elongation by modulating gibberellin metabolism in rice. The Plant Cell 26(11):4376–4393 DOI 10.1105/tpc.114.132092.

Tokunaga H, Kojima M, Kuroha T, Ishida T, Sugimoto K, Kiba T, Sakakibara H. 2012. Arabidopsis lonely guy (LOG) multiple mutants reveal a central role of the LOG-dependent pathway in cytokinin activation. Plant Journal 69(2):355–365 DOI 10.1111/j.1365-313X.2011.04795.x.

Unterholzner SJ, Rozhon W, Papacek M, Ciomas J, Lange T, Kugler KG, Mayer KF, Sieberer T, Poppenberger B. 2015. brassinosteroids are master regulators of gibberellin biosynthesis in arabidopsis. The Plant Cell 27(8):2261–2272 DOI 10.1105/tpc.15.00433.

Waldie T, Leyser O. 2018. Cytokinin targets auxin transport to promote shoot branching. Plant Physiology 177(2):803–818 DOI 10.1104/pp.17.01691.

Wang L, Feng C, Zheng X, Guo Y, Zhou F, Shan D, Liu X, Kong J. 2017. Plant mitochondria synthesize melatonin and enhance the tolerance of plants to drought stress. Journal of Pineal Research 63(3):e12429 DOI 10.1111/jpi.12429.

Wang L, Zhao Y, Reiter RJ, He C, Liu G, Lei Q, Zuo B, Zheng XD, Li Q, Kong J. 2014. Changes in melatonin levels in transgenic ‘Micro-Tom’ tomato overexpressing ovine AANAT and ovine HIOMT genes. Journal of Pineal Research 56(2):134–142 DOI 10.1111/jpi.12105.

Werner T, Nehnevajova E, Köllmer I, Novák O, Strnad M, Krámer U, Schmülling T. 2010. Root-specific reduction of cytokinin causes enhanced root growth, drought tolerance, and leaf mineral enrichment in arabidopsis and tobacco. The Plant Cell 22(12):3905–3920 DOI 10.1105/tpc.109.072694.

Xin P, Yan J, Fan J, Chu J, Yan C. 2013. An improved simplified high-sensitivity quantification method for determining brassinosteroids in different tissues of rice and arabidopsis. Plant Physiology 162(4):2056–2066 DOI 10.1104/pp.113.221952.

Xu MY, Zhang I, Li WW, Hu XL, Wang M, Fan YL, Zhang CY, Wang L. 2014. Stress-induced early flowering is mediated by miR169 in Arabidopsis thaliana. Journal of Experimental Botany 109(1):89–101 DOI 10.1093/jxb/ert353.

Yang DL, Yao J, Mei CS, Tong XH, Zeng LJ, Li Q, Xiao LT, Sun TP, Li J, Deng XW, Lee CM, Thomashow MF, Yang Y, He Z, He SY. 2012. Plant hormone jasmonate prioritizes defense over growth by interfering with gibberellin signaling cascade. Proceedings of the National Academy of Sciences 109(19):E1192–E1200 DOI 10.1073/pnas.1201616109.

Yang J, Worley E, Udvardi M. 2015. A NAP-AAO3 regulatory module promotes chlorophyll degradation via ABA biosynthesis in arabidopsis leaves. The Plant Cell 26(12):4862–4874 DOI 10.1105/tpc.114.133769.
Yang J, Zhang J, Wang Z, Zhu Q, Wang W. 2001. Hormonal changes in the grains of rice subjected to water stress during grain filling. *Plant Physiology (Bethesda)* 127(1):315–323 DOI 10.1104/pp.127.1.315.

Yu Y, Kim HS, Ma P, Jia Z, Guo X, Xie Y, Kwak SS, Zhang P, Bian X. 2020. A novel ethylene-responsive factor IbERF4 from sweetpotato negatively regulates abiotic stress. *Plant Biotechnology Reports* 14(4):397–406 DOI 10.1007/s11816-020-00612-x.

Yoshida T, Fernie AR, Shinozaki K, Takahashi F. 2021. Long-distance stress and developmental signals associated with abscisic acid signaling in environmental responses. *The Plant Journal* 105(2):477–488 DOI 10.1111/tpj.15101.

Zhang H, Wang L, Shi K, Shan D, Zhu Y, Wang C, Bai Y, Yan T, Zheng X, Kong J. 2019. Apple tree flowering is mediated by low level of melatonin under the regulation of seasonal light signal. *Journal of Pineal Research* 66(2):e12551 DOI 10.1111/jpi.12551.

Zheng X, Tan DX, Allan AC, Zuo B, Zhao Y, Reiter RJ, Wang L, Wang Z, Guo Y, Zhou J, Shan D, Li Q, Han Z, Kong J. 2017. Chloroplastic biosynthesis of melatonin and its involvement in protection of plants from salt stress. *Scientific Reports* 7(1):2827 DOI 10.1038/srep41236.

Zheng X, Zhao Y, Shan D, Shi K, Wang L, Li Q, Wang N, Zhou J, Yao J, Xue Y, Fang S, Chu J, Guo Y, Kong J. 2018. *MdWRKY9* overexpression confers intensive dwarfing in the M26 rootstock of apple by directly inhibiting brassinosteroid synthetase *MdDWF4* expression. *The New Phytologist* 217(3):1086–1098 DOI 10.1111/nph.14891.

Zuo B, Zheng X, He P, Wang L, Lei Q, Feng C, Zhou J, Li Q, Han Z, Kong J. 2014. Overexpression of *MzASMT* improves melatonin production and enhances drought tolerance in transgenic Arabidopsis thaliana plants. *Journal of Pineal Research* 57(4):408–417 DOI 10.1111/jpi.12180.

Zhang N, Sun Q, Zhang H, Cao Y, Weeda S, Ren S, Guo YD. 2015. Roles of melatonin in abiotic stress resistance in plants. *Journal of Experimental Botany* 66(3):647–656 DOI 10.1093/jxb/eru336.

Zong X, Lv J, Anjum SA, Wu X, Wu C, Li Y, Wang S. 2021. Effects of NPK and ALA on endogenous hormones of *Leymus chinensis* (Trin.) Tzvel under drought stress. *Bangladesh Journal of Botany* 50(1):113–117 DOI 10.3329/bjb.v50i1.52678.