Characterization of poplar growth-regulating factors and analysis of their function in leaf size control

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Research article

Keywords: Growth-regulating factor, phylogenetic relationship, miR396, leaf development, Populus

DOI: https://doi.org/10.21203/rs.3.rs-30608/v5

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Abstract

Background: Growth-regulating factors (GRFs) are plant-specific transcription factors that control organ size. Nineteen GRF genes were identified in the Populus trichocarpa genome and one was reported to control leaf size by regulating cell expansion. In this study, we further characterize the roles of the other poplar GRFs in leaf size control in a similar manner.

Results: The 19 poplar GRF genes were clustered into six groups according to their phylogenetic relationship with Arabidopsis GRFs. Bioinformatic analysis, degradome, and transient transcription assays showed that 18 poplar GRFs were regulated by miR396, with GRF12b the only exception. The functions of PagGRF6b (Pag, Populus alba × P. glandulosa), PagGRF7a, PagGRF12a, and PagGRF12b, representing three different groups, were investigated. The results show that PagGRF6b may have no function on leaf size control, while PagGRF7a functions as a negative regulator of leaf size by regulating cell expansion. By contrast, PagGRF12a and PagGRF12b may function as positive regulators of leaf size control by regulating both cell proliferation and expansion, primarily cell proliferation.

Conclusions: The diversity of poplar GRFs in leaf size control may facilitate the specific, coordinated regulation of poplar leaf development through fine adjustment of cell proliferation and expansion.

Background

Growth-regulating factors (GRFs) are plant-specific transcription factors that regulate the growth and development of leaves, roots, stems, flowers, and seeds by regulating cell proliferation or cell expansion, leading to the formation of larger organs [1-4]. GRFs form a multigene family found in the reported plant genomes: six genes in Camellia sinensis, eight genes in Vitis vinifera, nine genes in Arabidopsis thaliana, nine genes in Citrus sinensis, 10 genes in Pyrus bretschneideri, 12 genes in Oryza sativa, 13 genes in Solanum lycopersicum, 14 genes in Zea mays, 17 genes in Brassica rapa, 19 genes in Populus trichocarpa, and 25 genes in Nicotiana tabacum [5-14]. The Glu-Leu-Glu (QLQ) and Trp-Arg-Cys (WRC) domains are essential for GRF function in protein–protein interactions [15] and DNA binding [16], respectively. Genome-wide analyses revealed that GRFs and a few bZIP transcription factors are the major targets of miR396 [17].

GRFs are important for leaf size control [1-4]. Overexpression of AtGRF1 (At, Arabidopsis thaliana), rAtGRF2 (with mutations in the miR396 target sites, miR396-resistant version), rAtGRF3, rAtGRF7, rAtGRF9, AtGRF5, BnGRF2 (Bn, Brassica napus), or BrGRF8 (Br, Brassica rapa) in Arabidopsis thaliana and overexpression of rZmGRF1 (Zm, Zea mays) in Zea mays all resulted in larger leaves, while grf mutations or overexpression of miR396 led to smaller leaves [6, 10, 18-22]. Interestingly, most reported GRFs (AtGRF2, AtGRF3, AtGRF5, BnGRF2, BrGRF8, and ZmGRF1) control leaf size by regulating cell proliferation [10, 15, 18, 20-22], except ZmGRF10, which modulates leaf size via both cell proliferation and cell expansion, but mainly through cell proliferation [23]. Previously, we found that overexpression of one of the poplar GRFs, GRF15, also led to larger leaves and further analysis revealed that this GRF
controlled leaf size mainly by regulating cell expansion [24], which differed from the reported GRFs. Since 19 GRFs have been identified in the Populus trichocarpa genome [11] and leaf development is important for poplar biomass production in species like P. nigra and for drought/salt tolerance in species like P. euphratica, we wondered whether and how the other poplar GRFs function in leaf size control.

Here, we renamed the poplar GRFs according to their phylogenetic relationship with Arabidopsis GRFs and clustered them into six groups, and characterized the functions of PagGRF6b, PagGRF7a, PagGRF12a, and PagGRF12b from three different groups. We found that not all poplar GRFs regulate leaf development and their mechanisms of leaf size control are diverse.

Results

Names of poplar GRFs according to their Arabidopsis orthologs

Nineteen candidate GRF genes were found in the Populus trichocarpa genome [11]. To enable the comparison of PtrGRFs (Ptr, Populus trichocarpa) with the well-studied AtGRFs, the 19 identified PtrGRFs were renamed according to their Arabidopsis orthologs (Fig. 1, Fig. S1). According to the phylogenetic tree, the PtrGRFs could be classified into six groups (Fig. 1a, Fig. S2), with Group VI as a supplementary group to the reported classification of AtGRFs [6]. In Group I, four PtrGRFs clustered with AtGRF1 and AtGRF2 and were named PtrGRF1/2a, PtrGRF1/2b, PtrGRF1/2c, and PtrGRF1/2d (Fig. 1a). In Group II, only one PtrGRF gene corresponded to AtGRF3 and AtGRF4 and was named PtrGRF3/4 (Fig. 1a). In Group III, AtGRF5 and AtGRF6 each have two poplar orthologs, which were named accordingly (Fig. 1a). In Group IV, three PtrGRFs were named according to their sequence similarity to AtGRF7 and AtGRF8 (Fig. 1a). In Group V, although three PtrGRFs clustered with AtGRF9, only one PtrGRF with two WRC domains was named PtrGRF9 (Fig. 1a and b). In addition, four PtrGRFs with no close Arabidopsis orthologs were clustered in Group VI and named PtrGRF10a, PtrGRF10b, PtrGRF11a, and PtrGRF11b (Fig. 1a). The two PtrGRFs that clustered with AtGRF9, but lacked WRC domains were renamed PtrGRF12a and PtrGRF12b (Fig. 1a and b). Table S1 shows the complete gene information for PtrGRFs and AtGRFs.

The regulation of PtrGRFs by miR396

Since GRFs are the major targets of miR396 [17], the relationship between miR396 and PtrGRFs was investigated. First, the sequences of PtrGRFs and the mature sequences of poplar miR396b were uploaded to RNAhybrid [25, 26] to analyze whether PtrGRFs are targets of miR396. This showed that all of the PtrGRFs, except PtrGRF12b, have the potential to hybridize with miR396b with a minimal free energy hybridization value less than -33 kcal/mol, suggesting that these PtrGRFs could be targets of miR396 (Fig. 2a). For PtrGRF12b and miR396, the number of mismatches exceeded the other hybridization pairs and the hybridization energy was -28 kcal/mol, which exceeded the values observed for most endogenous miRNA targets [27], suggesting that PtrGRF12b is not a target of miR396 (Fig. 2a). Then, we aligned the target sequences of PtrGRFs to the mature miR396b sequences (Fig. S3). The sequences of PtrGRF1/2a-PtGRF12a and miR396 matched perfectly, while a thymine to adenine change
in the 3' terminal of *PtrGRF12b* led to a mismatch, indicating that *PtrGRF12b* is the only *PtrGRF* not targeted by *miR396* (Fig. S3).

In addition, degradome sequencing data [28] were analyzed to identify *miR396* cleavage sites in the *PtrGRFs* (Fig. 2b). As expected, the *miR396* cleavage sites of most *PtrGRFs* were found in the degradome data and no such a site was found in the *GRF12b* transcript (Fig. 2b, Table S2), proving the *in vivo* regulation of the expression of *PtrGRFs* by *miR396* was consistent with the RNAhybrid analysis.

Furthermore, transient expression was used to investigate the regulation of poplar *GRFs* by *miR396*. On fusing *PagGRF1/2c, PagGRF9, PagGRF10b, PagGRF11b*, and *PagGRF12b*, genes isolated from poplar 84K (see Methods), with YFP (Yellow Fluorescent Protein) and expressing them transiently in tobacco leaves, the fluorescence signals of all of the *PagGRF-YFP* fusion proteins were very weak (Fig. S4), except that of *PagGRF12b* (Fig. 3a). Considering the functional conservation of plant miRNAs, the weak fluorescence signal may be due to the cleavage of *PagGRFs* by tobacco *miR396*. To test this, *miR396*-resistant versions of the *GRFs*, which contained six point mutations within the *miR396*-complementary domain of the *GRF* sequence to increase the number of mismatches without altering the amino acid sequence, were constructed and transiently expressed in tobacco leaves (Fig. S5). As expected, the fluorescence signals of the m*PagGRF-YFP* fusion proteins were strong and merged with the DAPI signals (Fig. 3a), indicating that *miR396* targeted all of the *PagGRFs*, except *PagGRF12b*. Furthermore, transient co-expression assays were performed and *PtrmiR408* was used as a negative control to evaluate the regulation of *PagGRF* by *PagmiR396b* (Fig. 3b). Similar to the fluorescent signals of *GRF1/2d* [previously named *GRF15* by Cao *et al.* (2016)] in our published results [24], the fluorescent signals of *GRF12a-YFP* were weak when co-expressed with *PtrmiR408*, but were faint and difficult to detect when co-expressed with *PagmiR396b*, indicating that *PagmiR396b* could downregulate the expression of *PagGRF12a*. By contrast, comparable strong fluorescence of m*GRF12a-YFP*, the mutated version, was detected when co-expressed with *PagmiR396a* or *PtrmiR408*. These results confirmed that *PagmiR396b* could target *PagGRF* directly *in vivo*.

**Overexpression of *PagGRF6b, PagGRF7a, PagGRF12a*, and *PagGRF12b* led to diverse changes in leaf size in transgenic poplar**

The result in our previous study [24] showed that *GRF1/2a, GRF1/2b, GRF1/2c, GRF1/2d, GRF5a, GRF5b, GRF6b, GRF7a, GRF7b, GRF8, GRF9, GRF10a, GRF11a, GRF11b*, and *GRF12a* were all highly expressed in young leaves, suggesting that these GRFs may have a role in leaf size control. In addition, although its expression was relative low in all tissues, the *miR396* independent GRF, *GRF12b*, had higher relative expression in young leaves. Therefore, to investigate the function of poplar GRFs in leaf size control, *PagGRF6b* representing group III, *PagGRF7a* from group IV, and *PagGRF12a* and *PagGRF12b* from group V were chosen to generate transgenic plants for functional characterization (Figs. 1 and 4). The mutated versions of *PagGRF6b, PagGRF7a*, and *PagGRF12a*, with synonymous mutations in the *miR396* target sites, were used to avoid degradation by *miR396* (Fig. S6). Three overexpression (OE) lines each for *mPagGRF6b, mPagGRF7a, mPagGRF12a*, and *PagGRF12b* with moderately increased expression of the
corresponding gene were chosen for further investigation (Fig. S7). The leaf size of the \textit{mPagGRF6b} OE plants did not differ significantly (Fig. 4a), while \textit{mPagGRF7a} OE plants had 26.8% smaller leaves than those of the control (CK) (Fig. 4b). By contrast, \textit{mPagGRF12a} and \textit{PagGRF12b} OE plants had 16.1% and 28.1% larger leaves, respectively, in comparison with CK (Fig. 4c and d).

The leaf epidermis cell area was measured and leaf cell numbers were calculated for \textit{mPagGRF6b}, \textit{mPagGRF7a}, \textit{mPagGRF12a}, and \textit{PagGRF12b} OE plants and compared with the CK. The leaf epidermis cell area of \textit{mPagGRF6b} did not change significantly (Fig. 4a), while it decreased in \textit{mPagGRF7a}, \textit{mPagGRF12a}, and \textit{PagGRF12b} OE plants (Fig. 4b-d). The number of leaf cells in the \textit{mPagGRF6b} and \textit{mPagGRF7a} OE plants did not differ significantly (Fig. 4a and b), but increased significantly in the \textit{mPagGRF12a} and \textit{PagGRF12b} OE plants (Fig. 4c and d).

Furthermore, expression of the cell proliferation marker genes \textit{CYCLINB1;1a} and \textit{CYCLINB1;1b} and cell expansion marker genes \textit{EXPA11a} and \textit{EXPA11b} (Zhou et al. 2019) was examined in the fifth leaves from \textit{mPagGRF6b}, \textit{mPagGRF7a}, \textit{mPagGRF12a}, and \textit{PagGRF12b} OE plants. Consistent with our observations, expression of \textit{CYCLINB1;1a} and \textit{CYCLINB1;1b} was unaltered in \textit{mPagGRF6b} and \textit{mPagGRF7a} OE plants, but upregulated in the \textit{mPagGRF12a} and \textit{PagGRF12b} OE plants (Fig. 5, Fig. S8). Meanwhile, the expression of \textit{EXPA11a} and \textit{EXPA11b} did not change much in the \textit{mPagGRF6b} OE plants, but was downregulated significantly in \textit{mPagGRF7a}, \textit{mPagGRF12a}, and \textit{PagGRF12b} OE plants (Fig. 5, Fig. S8).

These results indicate that \textit{PagGRF6b} has no function in leaf size control; \textit{PagGRF7a} functions as a negative regulator of leaf size, mainly by regulating cell expansion; and \textit{PagGRF12a} and \textit{PagGRF12b} positively regulate leaf size through both cell proliferation and cell expansion, but mainly through cell proliferation.

**Discussion**

The expansion of GRFs in \textit{Populus} and their functional diversification in leaf development have drawn our attention. We have re-grouped the 19 GRFs identified in the \textit{P. trichocarpa} genome into six groups according to their phylogenetic relationships with their \textit{Arabidopsis} counterparts and renamed these poplar GRFs based on orthology. This facilitates the comparison of the evolution and functional diversity of GRF members in \textit{Populus} and \textit{Arabidopsis}. We found that as a result of the divergence of GRF sequences in \textit{Populus}, one of the 19 \textit{PtrGRFs}, \textit{PtrGRF12b}, was not targeted by miR396 and that \textit{PagGRF6b}, \textit{PagGRF7a}, \textit{PagGRF12a}, and \textit{PagGRF12b} worked differently in leaf size control.

Previously, we reported that PagGRF15 (which named as PagGRF1/2d in this study) could work as a positive regulator on leaf size through mainly regulating cell expansion [24]. Here, we found that PagGRF7a acts distinctly as a negative regulator on leaf size while PagGRF6b has no effect on leaf development, though PagGRF12a and PagGRF12b are similar to PagGRF1/2d functioning as positive regulators, indicating that different members of poplar GRFs have distinct roles in leaf size control. In addition, we found that PagGRF7a regulates leaf size through negatively affecting cell expansion while PagGRF12a and PagGRF12b though positively affecting cell proliferation and negatively regulating cell
expansion. All these GRFs exhibit differences with PagGRF1/2d that positively affects cell expansion. The unexpected diversified regulation on leaf size control by various poplar GRFs provides more and additional information than our previous report about PagGRF1/2d.

miR396 regulates the expression of GRFs through direct cleavage of complementary sequences in the GRF genes [18]. Here, we found that 18 PtrGRFs were regulated by miR396, with PtrGRF12b the only exception, based on sequence comparison, the cleavage sites of transcripts, and in vivo miRNA-target analysis. In comparison, miR396 did not target two GRFs in A. thaliana, AtGRF5 and AtGRF6 [29, 30]. PtrGRF12b belongs to group V, while AtGRF5 and AtGRF6 belong to group III, suggesting that the miR396 regulation pattern of GRFs has different features in the two species. It was recently reported that AtGRF5 plays roles in chloroplast development, nitrogen signaling, and senescence, apart from its function in leaf development [31]. Therefore, the loss and gain of miR396 regulation of GRFs may cause functional shifts in their roles in plant growth and development. Further studies are needed to investigate whether PtrGRF12b has functions in addition to those of other miR396-regulated GRFs.

Additionally, our study also suggests the importance of the regulation of GRF genes by miR396 in poplar, which is found in Arabidopsis [18, 21, 32]. Firstly, the miR396 regulation on GRFs using miR396-resistant version have been tested through transient expression assays in order to obtain the overexpression of GRFs. The fluorescence signals of the cells expressing PagGRF6b-YFP, PagGRF7a-YFP and PagGRF12a-YFP were faint and hardly detected, while the signals expressed their miR396-resistant version were strong, exampled as PagGRF12a-YFP in Fig. 3, which indicated that these GRFs were regulated by the existing miR396 in cells. Secondly, although PagGRF12a and PagGRF15 (PagGRF1/2d in this study) [24] are positive regulators on leaf size, the PagmiR396b OE plants showed a phenotype of smaller leaves [24], suggesting the “epistasis” effect of miR396b on the regulation of leaf size. Therefore, the interaction of miR396 and GRFs is important for leaf development.

GRFs are important regulators of leaf development [3, 4], and we found the functional divergence of GRFs in this study. PagGRF7a is a negative regulator, while PagGRF12a and PagGRF12b are positive regulators of leaf size, and PagGRF6b has no effect on leaf size. In Arabidopsis, AtGRF1, AtGRF2, AtGRF3, AtGRF5, and AtGRF7 all function as positive regulators of leaf size [15, 18, 21, 32], while only AtGRF9 functions as a negative regulator of leaf size [33]. Therefore, GRFs from poplar and Arabidopsis show diverse regulation of leaf size and their functions need to be assessed individually. In addition, it is interested that poplar GRF12a and GRF12b, like rice and maize GRF10 [23, 30], have truncated C-terminal end. It was proposed that overexpression of ZmGRF10 may break the homeostasis of distinct GRF/GIF complexes and result in the altered representation of other GRF/GIF complex to affect leaf growth [23], whether poplar GRF12a and GRF12b also work in this way needs to be investigated.

We also noticed that poplar and Arabidopsis GRFs classified in the same group could function in different ways in leaf size control. For instance, PagGRF6b from Group III has no effect on leaf size, while AtGRF5 from Group III is a positive regulator of leaf size [15]. PagGRF7a from Group IV functions as a negative regulator, while AtGRF7 from the same group functions as a positive regulator of leaf size [32].
Similarly, *PagGRF12a* and *PagGRF12b* from Group V work as positive regulators, while *AtGRF9* from Group V is a negative regulator of leaf size [33]. In comparison, although *PagGRF1/2d* from Group I is similar to *AtGRF1* and *AtGRF2* from Group I [18] and all three function as positive regulators of leaf size [24], *PagGRF1/2d* functions mainly by regulating cell expansion, while *AtGRF1* and *AtGRF2* function mainly by regulating cell proliferation. Therefore, the ways in which GRFs control leaf size in poplar cannot be simply inferred from their orthologs in *Arabidopsis*.

Previously, we reported that *PagGRF1/2d* regulated leaf size mainly by regulating cell expansion in poplar [24], which is different from all reported *Arabidopsis* GRFs, including *AtGRF1*, *AtGRF2*, *AtGRF5*, *AtGRF7*, and *AtGRF9*, which mainly act by regulating cell proliferation [15, 18, 21, 32, 33]. In this study, we found that *PagGRF12a* and *PagGRF12b* are involved in leaf size control mainly through regulating cell proliferation, while *PagGRF7a* and *PagGRF1/2d* negatively or positively, respectively, regulate leaf size mainly by regulating cell expansion. Therefore, the underlying mechanisms by which GRFs regulate leaf size are more diverse in poplar than in *Arabidopsis*. Leaf size is important for biomass production in woody plants [34] and should be under tight control. Poplar has more than twice the number of GRFs than *Arabidopsis* (19 vs. 9), so the diverse regulation in leaf size of these GRFs in poplar will facilitate the specific and coordinated regulation of leaf development through fine-tuning of cell proliferation and expansion.

**Conclusions**

In conclusion, we analyzed the phylogenetic relationship of GRF genes in *Populus* with their counterparts in *Arabidopsis* and functionally characterized *PagGRF6b*, *PagGRF7a*, *PagGRF12a*, and *PagGRF12b*, which work differently in leaf size control in transgenic poplar. This diversity may facilitate the specific, coordinated regulation of poplar leaf development through fine adjustment of cell proliferation and expansion. Our findings provide an abundant resource for genetic engineering leaf size in trees.

**Methods**

**Phylogenetic tree construction**

*Populus* GRF gene sequences were downloaded from the Poplar Genome Database (http://www.phytozome.net/poplar.php, release 3.0). All sequences were confirmed according to the annotation of the QLQ and WRC domains. WoLF PSORT (http://wolfpsort.org) was used to predict the protein subcellular localization. The pI and molecular weight were estimated using Lasergene. The full-length protein sequences were aligned using ClustalX2 (ver. 2.1) [35]. A neighbor-joining phylogenetic tree was constructed using MEGA (v5.0) with the bootstrap method (1000 bootstrap replicates, Poisson model, uniform rates, and pairwise deletion) [36]. Functional motifs or domains of *PtrGRF* sequences were analyzed using the reported FFD, GPL, and TQL motifs [8] as queries to find the corresponding sequences.
Degradome sequencing

The degradome data was from our previous study [28]. In brief, the degradome libraries of *P. tomentosa* were constructed from the poly(A) tail-containing fraction of total RNA samples pooled from the regenerating tissues after girdling to identify target genes of miRNAs. Then, data were analyzed using the CleaveLand pipeline and psRNATarget (http://plantgrn.noble.org/psRNATarget/) to predict the targets of miRNAs against the transcript sequences of *P. trichocarpa* genome (V3.0).

Transient expression assay

The transient expression assay was conducted according to our previous report [24]. *GRF1/2c, GRF9, GRF10b, GRF11b*, and *GRF12b* were cloned from the hybrid poplar clone 84K (*Populus alba × P. glandulosa, Pag*) reserved by State Key Laboratory of Tree Genetics and Breeding, Chinese Academy of Forestry. pEarleyGate 101 vector was used to generate the PagGRF-YFP construct, while the pMDC32 vector was used to overexpress *PagmiR396b* and *PtrmiR408*. The various construct combinations were introduced into 1-month-old *Nicotiana benthamiana* (reserved by State Key Laboratory of Tree Genetics and Breeding, Chinese Academy of Forestry) leaves through *Agrobacterium*-mediated infiltration. Fluorescence signals were probed using LSM 510 AX70 (Zeiss).

Plant transformation

Plant transformation was done as previously reported [24]. *GRF6b, GRF7a, GRF12a*, and *GRF12b* were cloned from 84K using the primers listed in Table S3. pMDC32 vector was used to overexpress *PagGRF6b, PagGRF7a, PagGRF12a*, and *PagGRF12b*. All vectors were transformed into 84K leaf discs via *Agrobacterium*-mediated transformation. Tissue-cultured plants were grown under long-day conditions (16 h light/8 h dark). Transgenic plants were confirmed by examining the expression of the corresponding genes.

Leaf phenotyping

Leaf phenotyping was performed as described in our previous study [24]. Briefly, the first completely uncurled leaf was defined as the first leaf. The fifth leaves of OE and CK plants were detached, fixed with FAA (formaldehyde: acetic acid: 96% alcohol: water; 10:5:50:35), cleared with chloral solution (200 g chloral hydrate, 20 g glycerol, and 50 mL dH₂O), and surveyed using a confocal Zeiss LSM 510 AX70 microscope. The cell number in the lower epidermis was calculated by dividing the leaf area by the area of epidermal cells. At least six leaves were used for the leaf area measurements and more than 100 epidermal cells in each leaf were used for cell area measurements.

RNA isolation and quantitative RT-PCR

The expression of *CYCLINB1;1a, CYCLINB1;1b, EXPA11a*, and *EXPA11b* in OE and CK plants was analyzed using quantitative RT-PCR (qRT-PCR) according to our previous study [24]. Briefly, the fifth leaves were collected from 2-month-old OE and CK plants and total RNA was extracted using the easy-
spin plus RNeasy Plant Mini Kit (Aidlab, Beijing, China). First-strand cDNA was synthesized using the SuperScript III reverse transcription kit (TaKaRa, Dalian, China) and oligo dT primers. All primer sequences are listed in Table S3. Real-time PCR was conducted on a LightCycler 480 (Roche, Basel, Switzerland) using SYBR Premix Ex Taq™ Kit (TaKaRa, Dalian, China). Actin and UBQ were used as internal controls.

**Abbreviations**

GRF: Growth-regulating factor; QLQ: Glu-Leu-Glu; WRC: Trp-Arg-Cys; Pag, *Populus alba × P. glandulosa*; At *Arabidopsis thaliana*; Br: *Brassica napus*, Br: *Brassica rapa*, Zm: *Zea mays*, Ptr, *Populus trichocarpa*; YFP: Yellow Fluorescent Protein; OE: overexpression; qRT-PCR: quantitative real time polymerase chain reaction.

**Declarations**

**Ethics approval and consent to participate**

Not applicable.

**Consent for publication**

Not applicable.

**Availability of data and materials**

All data generated or analyzed during this study are included in this article (and its supplementary information files) or are available from the corresponding author on reasonable request. Protein sequences have been deposited in GenBank (PagGRF6b, MW014326; PagGRF7a, MW015994; PagGRF9, MW015997; PagGRF10b, MW015995; PagGRF11b, MW015996; PagGRF12a, MW014327; PagGRF12b, MW014328).

**Competing interests**

The authors declare that they have no competing interests.

**Funding**

This work was supported by the Basic Research Fund of Research Institute of Forestry, Chinese Academy of Forestry (RIF-2014-08) to X.S, the National Natural Science Foundation of China (31570676) to X.S. The funders did not participate in study design, data collection and analysis, decision to publish, or preparation of the manuscript.

**Authors’ contributions**
M.L. and X.S. designed the experiments and wrote the main manuscript text. J.W. performed phenotype analysis, RNA isolation and qRT-PCR. H.Z. performed transgenic plants generation and transient expression assay. Y.Z. participated in transgenic plants cultivation. F.T. participated in degradome data analysis. P.S. participated in paper writing. All authors have reviewed the manuscript.

Acknowledgements

We thank Prof. Shanfa Lu (The Institute of Medicinal Plant Development, CAMS) for kindly providing the vector harboring PtrMiR408.

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