Pervasive duplication, biased molecular evolution and comprehensive functional analysis of the PP2C family in Glycine max

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Abstract

Background: Soybean (Glycine max) is an important oil provider and ecosystem participant. The protein phosphatase 2C (PP2C) plays important roles in key biological processes. Molecular evolution and functional analysis of the PP2C family in soybean are yet to be reported.

Results: The present study identified 134 GmPP2Cs with 10 subfamilies in soybean. Duplication events were prominent in the GmPP2C family, and all duplicated gene pairs were involved in the segmental duplication events. The legume-common duplication event and soybean-specific tetraploid have primarily led to expanding GmPP2C members in soybean. Sub-functionalization was the main evolutionary fate of duplicated GmPP2C members. Meanwhile, massive genes were lost in the GmPP2C family, especially from the F subfamily. Compared with other genes, the evolutionary rates were slower in the GmPP2C family. The PP2C members from the H subfamily resembled their ancestral genes. In addition, some GmPP2Cs were identified as the putative key regulator that could control plant growth and development.

Conclusions: A total of 134 GmPP2Cs were identified in soybean, and their expansion, molecular evolution and putative functions were comprehensively analyzed. Our findings provided the detailed information on the evolutionary history of the GmPP2C family, and the candidate genes can be used in soybean breeding.

Keywords: Protein phosphatase 2C, Soybean, Molecular evolution, Expansion, Functional analysis

Background

Reversible protein phosphorylation by protein kinases (PKs) and protein phosphatases (PPs) can regulate numerous biological processes [1]. The PKs can phosphorylate serine (Ser), threonine (Thr), and tyrosine (Tyr), whereas PPs can reverse these reactions by removing the phosphate groups [2]. Based on substrate specificity, the PPs can be classified into three major groups, namely, Ser/Thr phosphatases (STPs), protein Tyr phosphatases (PTPs), and dual-specificity phosphatases (DSPTPs) [2, 3]. PTPs can be further divided into phosphor-protein phosphatases (PPPs) and phosphoprotein metallophosphatases (PPMs) [4]. Many PPs belongs to the PPP family, whereas the PPM family includes protein phosphatase 2C (PP2Cs) and pyruvate dehydrogenase phosphate [2].

The PP2C enzyme is a monomer Mg²⁺/Mn²⁺-dependent PP [2]. The PP2C family has been found in archaea, bacteria, fungi, plants and animals [4]. In plants,
the PP2C family is the largest PP family [5, 6]. Over 70 PP2C members have been identified in Arabidopsis [7], rice [8], Brachypodium distachyon [9], Medicago truncatula [10], maize [11], and cotton [12]. In addition, the PP2C family can regulate plant growth and development, including root development [13], organ initiation [14], stem cell polarity [15], seed dormancy [16], and cell expansion [17]. Moreover, the PP2C members can respond to many biotic and abiotic stresses, including bacterial [18], wound [19], drought [20], salt [21], and cold stresses [22]. PP2C members are very important in hormonal signaling, such as abscisic acid (ABA) [23], jasmonic acid [19] and salicylic acid (SA) [24] pathways.

Soybean (Glycine max) is an important oil provider and a natural nitrogen-fixing plant. The release of the soybean genome can help comprehensively characterize the important gene families and analyze their putative functions [25]. Some PP2Cs have been previously identified in soybean. The PP2C-1 can be related to seed weight and size [26], and GmPP2C1 can interact with GmPYL1 in an ABA-dependent manner [27]. GmPP2Cs have already been previously found by searching the NCBI protein database [28], but a comprehensive expansion, molecular evolution and functional analysis of the PP2C family are still lacking in soybean. In this study, a comparative genomic analysis of the PP2C family revealed the species-specific molecular evolution and expansion in soybean. In addition, the expression profiles of the GmPP2C family in growth and development were systematically performed. The gene co-expression network (GCN) and gene regulatory network (GRN) in the GmPP2C members were constructed to unravel certain tissue-specific candidate genes and their putative regulatory mechanisms.

**Results**

**Identification and phylogenetic analysis of the PP2C family in soybean, grape, Arabidopsis, Amborella trichopoda, and Nymphaea colorata**

The Hidden Markov Model (HMM) profile of the PP2C domain (PF00481) was used as the query to identify the PP2C members in soybean, grape, Arabidopsis, and two sequenced basal angiosperms through the HMMER program. After confirming the presence of the PP2C domain using the CDD program, we found 134 GmPP2Cs in soybean, 62 VvPP2Cs in grape, 70 PP2C members in Arabidopsis, 43 AtrPP2Cs in A. trichopoda, and 40 AcPP2Cs in N. colorata (Fig. 1, Additional files 1, 2, 3, 4, 5, 6).

The phylogenetic analysis of the identified PP2C members was performed using FastTree, Mega, Phylib, and Bayesian (Fig. 2, Additional files 1, 7 and 8). The four different tools constructed the similar phylogenetic trees with high supporting values. The PP2C family was classified into 10 subfamilies, which were designated as A–J subfamilies based on the previous criterion [6]. Each subfamily contained different percentages of PP2C members (Fig. 1 and Additional file 9). In soybean, more than 20 GmPP2Cs were found in the D, E and F subfamilies, whereas only five GmPP2Cs were identified in the B and I subfamilies. The PP2Cs in grape, Arabidopsis, A. trichopoda, and N. colorata had the similar subfamily distributions. In addition, all of the subfamilies had more PP2C members in soybean than in other plants. The number of PP2C members in the other four eudicots was similar in the B, F, H, I and J subfamilies, but other subfamilies had more PP2Cs in grape and Arabidopsis.

**Conserved motifs and gene structure analysis of the GmPP2C members**

All PP2C members had the PP2C domain through the CDD program (Additional files 1 and 10). The locations of the PP2C domain were highly conserved in the PP2C subfamily. Twenty putative motifs were found in the GmPP2C family through the MEME tool (Fig. 2 and Additional file 11). On the basis of the distribution of motifs, the GmPP2C family could be classified into 10 groups, the same classification as the phylogenetic analysis. Eight motifs (motif 1, 2, 3, 4, 5, 6, 7 and 9) were annotated as the PP2C domain through the CDD program. More than 100 GmPP2Cs had 10 motifs (motif 1, 2, 3, 4, 6, 8, 11, 13, 15 and 16), and the other motifs existed in certain specific subfamilies (Additional file 12). Motif 7 existed in all the subfamilies except the C, D and E subfamilies. GmPP2Cs from the A, B, G and I subfamily had motif 9. Moreover, the C subfamily had motif 5, and the D subfamily contained motif 5, 10, 14 and 17. Motif 12 and 18 were enriched in the E subfamily, and the H subfamily had motif 14, 19 and 20. Motif 19 and 20 were only found in the H subfamily. Furthermore, the sequence similarity of GmPP2Cs was analyzed through the Blastp tool (Additional file 13A). GmPP2Cs had a significantly higher sequence similarity in the same subfamily than in the different subfamilies. GmPP2Cs from the H, I and J subfamilies had the highest sequence similarity (~ 600 score), followed by the B, C and D subfamilies (~ 400 score). The F subfamily contained the lowest sequence similarity members (~ 205 score).

Gene structure and intron position were analyzed in the GmPP2C family (Fig. 2 and Additional file 14). Each subfamily had a highly conserved gene structure. Most GmPP2Cs had less than five exons and four introns. However, the highly fragmented members were mainly found in the F, H and I subfamilies. Most GmPP2Cs only contained intron position 0, but GmPP2Cs from the D, E, F, H and I subfamilies had other intron positions.
Gene duplication and molecular evolution analysis of the PP2C family in soybean

Chromosomal location images of the PP2C family were generated in soybean (Additional files 9, 15 and 16). Each chromosome had GmPP2Cs, but these GmPP2Cs were unevenly distributed across the chromosomes.

Gene duplication events were investigated to illustrate the expansion pattern of the PP2C family in soybean. A total of 131 duplicated gene pairs were identified in the GmPP2C family (Fig. 3A and Additional file 17), whereas only 13 duplicated gene pairs were found in the VvPP2C family (Additional file 18). Duplication events occurred in each GmPP2C subfamily. The D and E subfamilies had the most duplication events (seven each), followed by the A, F and G subfamilies (more than five each) and the H subfamily (three each). The B and I subfamilies

| Subfamily | Species          | A. trichopoda | N. colorata | V. vinifera | G. max | A. thaliana |
|-----------|------------------|---------------|-------------|-------------|--------|-------------|
| A         | 5                | 4             | 12          | 18          | 9      |
| B         | 4                | 3             | 4           | 5           | 4      |
| C         | 2                | 1             | 3           | 6           | 5      |
| D         | 4                | 8             | 9           | 21          | 9      |
| E         | 5                | 3             | 7           | 20          | 12     |
| F         | 8                | 9             | 9           | 20          | 12     |
| G         | 4                | 2             | 6           | 16          | 6      |
| H         | 3                | 2             | 3           | 10          | 3      |
| I         | 1                | 2             | 2           | 5           | 2      |
| J         | 2                | 2             | 1           | 6           | 2      |
| No group  | 5                | 4             | 6           | 7           | 6      |
| Total     | 43               | 40            | 62          | 134         | 70     |

Fig. 1 Distribution of the PP2C members in grape, soybean, Arabidopsis, A. trichopoda, and N. colorata. The upper phylogenetic tree represents the evolutionary relationship of the five species.
Fig. 2 Phylogenetic relationship, putative motif distribution and gene structure of the GmPP2C family. The phylogenetic tree (left) was constructed using the Bayesian method. The numbers were posterior probability values of each clade, and different subfamilies were indicated by different letters and colors. The putative motifs (middle) were identified by the MEME program. Each motif was presented by the specific colored boxes, and the motif location can be estimated using the scale at the bottom. The gene structures (right) were drawn according to the location of exons and introns. The location of exons and introns can be estimated using the scale at the bottom.
only contained one duplication event. In grape, duplication events were only detected in the A, B, D, F and G subfamilies. Meanwhile, 121 GmPP2Cs were related to duplication events. However, only 13 GmPP2Cs, including five GmPP2Cs from the F subfamily, did not find any duplication event. In addition, 101 syntenic blocks were found in the GmPP2C family (Fig. 3A). The longest syntenic block, including six duplicated gene pairs, was from chromosome 04 and 06, and the syntenic block in chromosome 10 and 20 had five duplicated gene pairs. Furthermore, the sequence similarity of duplicated GmPP2Cs was analyzed through the Blastp tool (Additional file 13B). Duplicated GmPP2Cs from the H subfamilies had the highest sequence similarity (~ 700 score), and the B and F subfamilies contained the lowest sequence similarity members (~ 400 score).

Based on the sequence analysis and chromosomal distribution, all duplicated gene pairs were involved in the segmental duplication events, and no tandem duplication events were found (Additional file 17). Synonymous substitution rate (Ks) value distribution of the GmPP2C duplicated gene pairs was calculated to predict
the burst of duplication (Fig. 3B). The Ks values peaked at approximately 0.125 and 0.535, which were similar to the Ks distributions of all soybean duplicated gene pairs. Meanwhile, nonsynonymous substitution rate (Ka) value and Ka/Ks ratio were lower in the GmPP2C duplicated gene pairs than in all soybean duplicated gene pairs (Fig. 3C-E). Moreover, Ka values and Ks values were lower in the F and H subfamilies than in their averages across the entire PP2C family (Fig. 3F-G). The Ka/Ks ratio was less than 1 for the duplicated gene pairs, and it significantly declined in the D and H subfamilies (Fig. 3H). However, Ka values, Ks values, and Ka/Ks ratios of the other subfamilies were similar to one another. Furthermore, polarizing evolutionary rates were found in the GmPP2C subfamily. GmPP2C004/128 and GmPP2C017/073 had higher Ka values than other D and G duplicated gene pairs, and their corresponding sequences also had a lower similarity (Additional file 13B). The Ks values of GmPP2C004/128, GmPP2C016/105, GmPP2C042/125, and GmPP2C003/027 were more than 2, which were different from other gene pairs in the same subfamily. All of the GmPP2C subfamilies, expect for the D and H subfamilies, had a polarizing Ka/Ks ratio.

**Comparative genomic analysis of the PP2C members in soybean and grape**

The grape genome was analyzed to deconvolute the genomic complex in soybean [29, 30]. In this study, 96 orthologous genes of VvPP2Cs were identified in soybean (Fig. 4A and Additional file 19). Twenty VvPP2Cs can find two orthologous genes in soybean (Fig. 4B and Additional file 20). Eight VvPP2Cs with three soybean orthologous genes were found, mostly from the D and G subfamilies. Seven VvPP2Cs, mostly from the D and H subfamilies, had four orthologous genes in soybean. However, two VvPP2Cs from the F subfamily only detected one orthologous gene in soybean.

Ks value distribution was calculated to estimate the divergent time of the PP2C orthologous genes in soybean and grape (Fig. 4C). The Ks values peaked at 1.015, which was significantly lower in the PP2C orthologous genes than in all orthologous genes. Compared with all orthologous genes, Ka values, Ks values, and Ka/Ks ratios significantly decreased in the PP2C orthologous genes (Fig. 4D-F). In addition, the Ks values were lower in the H and I subfamilies than in their averages across the entire PP2C family (Fig. 4G). The Ka values were low in the F, H and I subfamilies, and the Ka/Ks ratios were low in the F and H subfamilies (Fig. 4H-I). However, Ka values, Ks values, and Ka/Ks ratios of the other subfamilies were similar to one another.

**Sequence similarity analysis of the PP2C members in soybean and basal angiosperms**

*A. trichopoda* and *N. colorata* are sequenced basal angiosperms. The sequence similarity of PP2Cs in soybean and basal angiosperms was calculated through the Blastp tool (Additional files 21 and 22). The duplicated gene pair (GmPP2C002/057) had the highest sequence similarity to its corresponding orthologous genes in *A. trichopoda* and *N. colorata*. Meanwhile, the highest similarity in soybean and *A. trichopoda* enriched in the PP2C members from the H subfamily, followed by the I subfamily (Fig. 5A). GmPP2Cs and NcPP2Cs from the H subfamily had the highest similarity (Fig. 5B).

**Expression profiles of the GmPP2C members in different tissues**

The RNA-seq data were analyzed to investigate the expression profiles of the GmPP2C members in different tissues (Fig. 6A and Additional file 23). Data were available for all GmPP2Cs. A total of 117 GmPP2Cs were expressed at medium and high levels (Fragments per Kilobase Million [FPKM] > 5) in one or more tissues. Based on the hierarchical clustering of the expression profiles, GmPP2Cs can be divided into four clusters (Fig. 6A-B). In cluster 1, 23 GmPP2Cs were highly expressed in almost each tissue, whereas 18 GmPP2Cs in cluster 2 had the low expression levels. Cluster 3 included 51 GmPP2Cs with the similar expression level in all tissues. Cluster 4 had 42 GmPP2Cs which were relatively highly expressed in the leaves and roots. In addition, most GmPP2Cs from the same subfamily had the similar expression level (Fig. 6C). Eleven GmPP2Cs from the F subfamily belonged to the cluster 1, and the cluster 2 mainly contained GmPP2Cs from the B, E and G subfamilies. More than half of the GmPP2Cs in cluster 3 came from the A, D and E subfamilies. The GmPP2Cs from the D, E and G subfamilies comprised most of the members in cluster 4. Furthermore, most of the duplicated genes showed the similar expression level in different tissues (Fig. 6A). A total of 102 duplicated gene pairs gathered the same expression clusters. The Pearson Correlation Coefficient (PCC) between GmPP2Cs was also calculated on the basis of the FPKM values (Fig. 6D and Additional file 24). The PCC between non-duplicated GmPP2Cs was a pearshape distribution, whereas the expression levels of most duplicated GmPP2Cs (73 gene pairs) were highly positively related (PCC > 0.5). Then, six duplicated gene pairs with high PCC and sequence similarity were performed through qRT–PCR analysis (Fig. 6E). The PCCs were more than 0.799 in the expression level of six duplicated gene pairs.
Fig. 4 (See legend on next page.)
Gene co-expression network analysis of the GmPP2C members

Through GCN analysis, 13,954 genes were grouped into 22 modules ranging from 234 genes (bisque) to 610 genes (ivory) (Additional file 25). Certain modules had a high-positive correlation, such as blue and cyan, and brown and darkgray (Fig. 7A).

Most GmPP2C members were grouped into five major modules (blue, brown, green, lightcyan1 and yellow) (Fig. 7B and Additional file 26). A total of 14 GmPP2Cs from the F and G subfamilies were clustered in lightcyan1, and blue had six members from the A subfamily. Ten GmPP2Cs from the D, E and F subfamilies were clustered in yellow. Meanwhile, genes in module were closely associated with traits (Fig. 7C and Additional file 27). The correlation was 1 between blue and flower, and green and nodule. A high correlation was observed between yellow and seed, brown and leaf, and lightcyan1 and leaf.

Five subnetworks with GmPP2Cs were extracted in the major modules (Fig. 7D). In blue, GmPP2C133 and GmPP2C104 had the most co-expressed genes enriched in gene ontology (GO):0045490 (pectin catabolic process). GmPP2C024, GmPP2C035 and GmPP2C091 were hub genes in brown, and their co-expressed genes were related to plastid organization (GO:0009657). In green, GmPP2C005, GmPP2C013, and GmPP2C118 were mostly connected with many genes with nodulation (GO:0009877). In lightcyan1, a hydrogen peroxide catalytic process was enriched in the co-expressed genes with GmPP2C092 and GmPP2C038. In yellow, many genes had a co-expressed relationship with GmPP2C031, GmPP2C071, and GmPP2C131, and were associated with lipid storage (GO:0019915).

cis-regulatory element analysis

The cis-regulatory elements are important in the biological functions and regulatory networks. In this study, the promoter regions of the GmPP2C members included six cis-regulatory elements (Fig. 8 and Additional file 28). All GmPP2Cs contained at least one cis-regulatory element. All GmPP2Cs from the A subfamily had the ABA-responsive element in their promoters. ABA-, Gibberellin-, and SA-responsive elements were found in many GmPP2C promoters from the D and F subfamilies. More than half of the Methyl Jasmonate (MeJA)-responsive elements existed in the GmPP2C promoters from the E, G and H subfamilies. Most of the promoter regions of the duplicated gene pairs contained the similar cis-regulatory elements in the GmPP2C family.
Gene regulatory network analysis of the GmPP2C members

The GRN of the GmPP2C members was performed through the MERLIN+Prior method (Fig. 9a). The GmPP2C members can be regulated by 324 transcription factors (TFs) (Additional file 29). Forty and thirty-eight TFs belonged to the WRKY and bHLH families, respectively (Fig. 9b). More than 20 TFs belonged to the ERF, MYB and C2H2 families. Meanwhile, six and five GmPP2Cs were target genes of Glyma.02G051100 and Glyma.15G232000, respectively (Fig. 9c). Twenty-four target genes of Glyma.02G051100 (~59%) belonged to the lightcyan1 module, and 37 target genes of Glyma.15G232000 (~79%) belonged to the blue module. In addition, 14 duplicated gene pairs can be regulated by the same TF in the GmPP2C family (Additional file 30). Four duplicated gene pairs can be regulated by the same TFs in the A and H subfamilies, respectively. The related target genes contained half of the GmPP2Cs (five members) in the H subfamily.

Fig. 6 Expression profiles of GmPP2Cs in different tissues. A Hierarchical cluster analysis of GmPP2C expression in seven tissues. B Expression profiles of four clusters in seven tissues. The red lines represented the mean expression level of GmPP2Cs in the corresponding cluster. C Gene numbers of the four clusters in different subfamilies. D The PCC distribution of expression levels in the duplicated and non-duplicated GmPP2Cs. E Expression of the duplicated GmPP2Cs in three tissues.
**Discussion**

In the present study, 134 GmPP2Cs were identified in soybean, whereas less than 70 members were found in grape, Arabidopsis, and two basal angiosperms (Fig. 1, Additional files 1, 2, 3, 4, 5, 6). The increased GmPP2Cs were due to the larger genome (1150 Mb) in soybean [25, 30]. The GmPP2C family could be phylogenetically classified into 10 subfamilies (Figs. 1 and 2, Additional files 1, 2, 3, 4, 5, 6).
The D, E, and F subfamilies had the most GmPP2Cs, whereas the B and I subfamilies had the least members (Fig. 1 and Additional file 9). The greater number was associated with the higher number of GmPP2Cs in the D, E, and F subfamilies. The PP2C members had the similar distribution in grape, Arabidopsis, and two basal angiosperms. Similar distribution of the PP2C family has been reported in Arabidopsis, *M. truncatula* and cotton [7, 10, 12]. Thus, the PP2C family had the conserved subfamily distribution in plants. Meanwhile, the duplicated gene pairs of the PP2C members in soybean (131) were 10 times more than those in grape (13) (Fig. 3A and Additional files 17 and 18). A greater number of the duplicated gene pairs could result in the increased number of the PP2C family (Fig. 1). The duplication events of the GmPP2C family could be found in each subfamily, but were unevenly distributed across the GmPP2C subfamilies. Most duplication events in the GmPP2C family were identified in the A, D, E, F and G subfamilies, whereas each subfamily in grape contained one duplication event at most. The duplication events in the A, D, E, F and G subfamilies led to the expansion of the GmPP2C members.

Fig. 8 Distributions of certain responsive-regulatory elements in the promoter regions of the GmPP2C members. The PP2C subfamilies in soybean were marked by the brackets. The colored box represented the responsive regulatory elements.
All duplication events were segmental duplication events in the GmPP2C family (Fig. 3A and Additional files 15 and 17). No tandem duplication events in the GmPP2C family might be related to the genomic fractionation from transposon activities, duplicating and relocating individual genes [31–34]. Moreover, all of the duplicated genes were located in the 101 syntenic blocks (Fig. 3A). The GmPP2C expansion was primarily caused by whole genome duplications [25]. The occurrence of the GmPP2C duplicated gene pairs corresponded to the legume-common duplication event and soybean-specific tetraploid through the Ks analysis (Fig. 3B), and the duplication events occurred simultaneously in GmPP2Cs and other genes (Fig. 3D). Our Ks values were a little lower than those in the previous reports because of the absence evolutionary rate correction [30]. Meanwhile, in our study, detecting the Ks peak of the core eudicot-common hexaploidy was difficult. No identification of the corresponding peak might be due to the Ks peak decline with increasing Ks values or the widespread gene losses after recursive polyploidizations [30, 35].

Based on the Ka/Ks ratio of the duplicated genes, the GmPP2C members mainly have experienced purifying selection with limited functional divergence after whole-genome duplications (Fig. 3E). The similar function in the duplicated genes was further supported by their similar expression profiles (Fig. 6 and Additional file 24) and distribution of responsive regulatory elements (Fig. 8). However, most of the duplicated genes cannot be regulated by the same TF through gene regulatory networks (Fig. 9 and Additional files 29 and 30). Thus, sub-functionalization was the main evolutionary fate of duplicated GmPP2Cs. Most of these duplicated genes have kept the original or similar functions of the ancestral genes after duplication events. In particular, many duplicated genes are known to have similar evolutionary
fates in plants [36, 37]. Moreover, the evolutionary rates of the duplicated genes were various in different GmPP2C subfamilies (Fig. 3F-H). Compared with other subfamilies, the H subfamily evolved slowest (Fig. 3F-H), had a high sequence similarity (Additional file 13) and lost less genes (Fig. 4B). The slow evolutionary rates of the H subfamily resulted from low gene loss rate after polyploidization. Furthermore, some polarizing evolutionary rates were found in the same subfamily (Fig. 3F-H). The high Ka values could result from the low sequence similarity (Additional file 13B), and the high Ks values were related to the loss of the orthologous genes (Fig. 4B). Sequence similarity and gene loss affected the Ka/ Ks ratio.

The legume-common duplication event and soybean-specific tetraploid provided an expected 1:4 ratio of the orthologous regions between grape and soybean [25, 29, 30]. Only seven VvPP2Cs were in line with the expected ratio (Fig. 4A-B, and Additional files 19 and 20). Thus, the large-scale gene losses were recognized in the GmPP2C family. The widespread gene losses might be due to the transposon movements or poor assemblies and annotations [32, 33]. The gene losses could help in diploidization, including a series of chromosome breakages and fusions [34, 38, 39]. In addition, the gene losses were unevenly distributed across the GmPP2C subfamily (Fig. 4B). The H subfamily in soybean lost less GmPP2Cs, whereas the F subfamily had a higher gene loss rate. The gene loss rate was associated with the sequence similarity in the GmPP2C subfamily (Additional file 13). Natural selection might be a reason for the unbalanced gene retention rates [32].

The divergent time of GmPP2Cs and VvPP2Cs was significantly later than that of other gene families in soybean and grape (Fig. 4C-D). The evolutionary rates of the PP2C family were significantly lower than that of other gene families in soybean (Fig. 4C-D). Moreover, compared with other subfamilies, the H subfamily evolved slowest (Fig. 3F-H), had a high sequence similarity (Additional file 13) and lost less genes (Fig. 4B). The slow evolutionary rates of the H subfamily resulted from low gene loss rate after polyploidization. Furthermore, some polarizing evolutionary rates were found in the same subfamily (Fig. 3F-H). The high Ka values could result from the low sequence similarity (Additional file 13B), and the high Ks values were related to the loss of the orthologous genes (Fig. 4B). Sequence similarity and gene loss affected the Ka/ Ks ratio.

The spatial expression analysis of the GmPP2C members revealed tissue-specific expression profiles (Fig. 6 and Additional file 23). Most of the GmPP2Cs had a very broad expression spectrum, suggesting their very important roles in the regulation of plant growth and development. The PP2C family had the similar results in rice, Arabidopsis, B. distachyon, maize and cotton [7, 9, 11, 12]. Moreover, GCN analysis indicated certain key putative genes in growth and development (Fig. 6 and Additional files 25, 26, 27). GmPP2C104, GmPP2C123 and GmPP2C133 might participate in flower cell morphogenesis, and an orthologous gene of GmPP2C123 (AT5G02760) could regulate the stamen filaments in Arabidopsis [41]. GmPP2C024, GmPP2C035 and GmPP2C091 directly co-expressed with certain plastid-related genes in chloroplast development, and AT4G27800, an orthologous gene of GmPP2C024 and GmPP2C035, reportedly played key roles in LHClII dephosphorylation [42]. GmPP2C029, GmPP2C038 and GmPP2C117 might be very important to leaf development. The mutant of AT2G28890 and AT1G07630 (the GmPP2C117 orthologous genes) had abnormally shaped leaves [43]. Meanwhile, GmPP2C031, GmPP2C071 and GmPP2C131 might be involved in seed development, and an orthologous gene of GmPP2C071 (AT5G51760) could regulate seed germination in Arabidopsis [44]. GmPP2C005, GmPP2C103, GmPP2C059 and GmPP2C118 might play important roles in the nodulation development stage, and LjNPP2C1 (a GmPP2C005 orthologous gene) in Lotus japonicus was related to both early and late stages of nodule development [45]. In addition, most GmPP2Cs could respond to the hormone signaling through the cis-regulatory element analysis (Fig. 7 and Additional file 28). The GmPP2Cs from the A subfamily might participate in the ABA pathway. Some PP2Cs from the A subfamily reportedly regulated the ABA signaling in soybean and other plants [23, 27, 46]. Moreover, through the GRN analysis, 40 TFs from the WRKY family could regulate the GmPP2C expression (Fig. 8 and Additional file 29), indicating that the evolution of the GmPP2C family was directly or indirectly associated with the WRKY family.

Conclusions

A total of 134 GmPP2C members with 10 subfamilies were found in the present study. The number of duplication events from the legume-common duplication event and soybean-specific tetraploid led to the GmPP2C expansion. All of the duplication events were the segmental duplication events. Sub-functionalization was the main evolutionary fate of duplicated PP2C members in soybean. Meanwhile, the GmPP2C family lost massive genes, especially in the F subfamily. The GmPP2C members evolved more slowly than other genes, and the
PP2C members from the H subfamily resembled their ancestral genes. In addition, some genes were considered as the putative key candidates that could control plant growth and development (GmPP2C104, GmPP2C123, and GmPP2C135 in flower cell morphogenesis; GmPP2C024, GmPP2C035, and GmPP2C091 in chloroplast development; GmPP2C029, GmPP2C038, and GmPP2C117 in leaf development; GmPP2C031, GmPP2C071, and GmPP2C131 in seed development; GmPP2C005, GmPP2C013, GmPP2C059 and GmPP2C118 in nodulation development).

Methods

Sequence retrieval and sequence analysis
The genome sequences of G. max, V. vinifera, A. thaliana and A. trichopoda were downloaded from the Phytozome database (https://phytozome.jgi.doe.gov/pz/portal.html). The genome sequences of N. colorata were obtained from the Waterlily Pond (http://waterlily.eplant.org). The HMM profile of the PP2C domain (PF00481) from the Pfam database (https://pfam.xfam.org/) was used to identify the PP2C members in soybean, grape, Arabidopsis, A. trichopoda and N. colorata with hmmsearch tool in HMMER 3.0 program [47]. Then, all putative PP2C members were further confirmed through the CDD program (https://www.ncbi.nlm.nih.gov/cdd). The conserved motifs of the PP2C family in soybean were identified by the MEME program (http://meme-suite.org/tools/meme). The parameters were set according to the previous report [48]. The chromosomal distribution images and the exon–intron structures of the GmPP2C genes were visualized by the TBtools software [49]. “Intron phases” were defined through the intron–exon location. Phase 0, 1 and 2 defined introns inserted between codons, between the first and second nucleotides of the codon and between the second and third nucleotides of the codon, respectively [11]. Further, the sequence similarity of the PP2C family in soybean and two basal angiosperms were compared through the Blastp program with default parameters.

Phylogenetic analysis
The PP2C proteins were aligned using the ClustalX software with default configurations [50]. The phylogenetic trees were constructed by four different tools. The maximum likelihood method was used with the FastTree version 2.1.3 using the Jones-Taylor-Thornton amino acid substitution model and the bootstrapping value [51]. The Bayesian analysis was performed through MrBayes version 3.1.2 with the mixed model and Bayesian posterior probabilities [52]. The neighbor-joining trees were analyzed by the PHYMLIP and MEGA tool, and the reliability was assessed by a 1000 bootstrapping test [53, 54]. All trees were visualized with Figtree version 1.4.3.

Syntenic and evolutionary rate analysis
Syntenic blocks were found by the MCScan software [55], and the syntenic relationship was visualized by the Circos program. The Ka, Ks and their ratio (Ka/Ks) were calculated with the Nei–Gojobori method by the TBtools software [49].

Expression analysis
The RNA-seq transcriptome dataset of soybean in different tissues (pod, leaf, root, stem, nodule, seed, and flower) were downloaded from the Phytozome database, and more details about experimental materials and RNA-seq data analysis are described in the previous report [56]. The expression level was log-transformed via the log2(FPKM+1) function [57]. The expression data were hierarchically clustered through the hclust function in R.

G. max cv. Tianlong-1 was used to construct the expression profiles of the PP2C family in soybean. Roots, stems, and leaves were harvested from 4-week-old seedlings. RNA extraction and qRT-PCR analysis were performed on the basis of previous studies [48]. The GmHELIC gene was used as the reference gene, and gene-specific primers were designed on the basis of their coding sequences (Additional file 31). qRT–PCR analysis was conducted by three biological replications. Moreover, the PCCs of the RNA-seq and qRT-PCR analyses were calculated by cor function in R.

Gene co-expression network analysis
The co-expression network was constructed by the weighted gene co-expression network analysis (WGCNA) in R [58]. The expression pattern in soybean used the same RNA-seq transcriptome dataset in the expression analysis. Genes with the low coefficient of variation (CV < 0.6) and the low maximum expression value (FPKM<10) were discarded and all GmPP2C genes were added into the analysis. The remaining 13,954 genes were used for the WGCNA analysis. The co-expression networks were performed with default settings, except for those whose soft-threshold power was 12, minModuleSize was 30, and mergeCutHeight was 0.25. The node and edge information of GmPP2Cs with edge weight > 0.6 was extracted from the interested modules, and only the top 1000 weight values were selected. The sub-networks were analyzed and visualized using the Cytoscape software.

GO and cis-element analysis
GO enrichment was performed using the GO Term Enrichment tool with P value ≤0.01 in PlantRegMap (http://plantregmap.cbi.pku.edu.cn/). Meanwhile, the 1000 bp upstream of the initiation codon (ATG) were extracted from the Phytozome database genes in the GmPP2C family. The potential responsive-regulatory
elements in the GmPP2C promoters were identified by the PlantCARE database (http://bioinformatics.psb.ugent.be/webtools/plantcare/html/). The distributions of the responsive-regulatory elements were visualized by the TBtools software [49].

**Gene regulatory network analysis**

GRN was performed by the MERLIN+Prior method [59]. The expression file used the same expression matrix in the WGCNA analysis. The soybean TFs as the regulator list were downloaded from the PlantTFDB database (http://planttfdb.cbi.pku.edu.cn/index.php). The TF binding site database in soybean as the prior network was obtained from the PlantRegMap database. The WGCNA module was set as the initial cluster assignment. The GRN networks were analyzed and visualized by the Cytoscape software.

**Supplementary information**

Supplementary information accompanies this paper at https://doi.org/10.1186/s12864-020-06877-4.

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**Additional file 1**

Phylogenetic and sequence analysis of the PP2C members in soybean, Arabidopsis, grape, A. trichopoda and N. colorata. The inner circle was the phylogenetic tree using the FastTree method. The numbers in the clades were the FastTree bootstrap values. The PP2C subfamilies were labeled using different letters and were indicated using different colors. The outer circle was the sequence analysis of PP2Cs. The conserved PP2C domain was represented by the blue box. The location of each PP2C protein can be estimated using the scale at the end of the circle.

**Additional file 2**

GmPP2C proteins identified in G. max.

**Additional file 3**

VvPP2C proteins identified in V. vinifera.

**Additional file 4**

AtPP2C proteins identified in A. thaliana.

**Additional file 5**

AtrPP2C proteins identified in A. trichopoda.

**Additional file 6**

NcPP2C proteins identified in N. colorata.

**Additional file 7**

Phylogenetic relationship of the GmPP2C members. The neighbor-joining (NJ) tree was constructed through the MEGA program, and the numbers in the clades are the bootstrap values. The PP2C subfamilies are grouped by colors.

**Additional file 8**

Phylogenetic analysis of the GmPP2C members. The neighbor-joining (NJ) tree was generated through PHYLIAP package, and the numbers in the clades are the bootstrap values. The PP2C subfamilies are grouped by colors.

**Additional file 9**

Percentage of GmPP2C numbers in the PP2C subfamily (upper) and its location distribution on each chromosome (bottom).

**Additional file 10**

Locations of the conserved PP2C domain in GmPP2Cs, VvPP2Cs, AtrPP2Cs, and NcPP2Cs.

**Additional file 11**

Motif information of GmPP2Cs through the MEME tool.

**Additional file 12**

Motif distributions in the GmPP2 subfamily.

**Additional file 13**

Sequence similarity of GmPP2Cs. (A) Sequence similarity of GmPP2Cs from the same and different subfamily. (B) Sequence similarity of duplicated GmPP2Cs in each subfamily.

**Additional file 14**

Structural analysis of GmPP2C in this study.

**Additional file 15**

Chromosomal locations of GmPP2Cs on all 20 chromosomes. The scale used is base pair (bp). Parentheses after the gene names show the PP2C subfamily.

**Additional file 16**

Genomic locations of the GmPP2C genes.

**Additional file 17**

Ka/Ks analysis for the duplicated PP2Cs in soybean.

**Additional file 18**

Ka/Ks analysis for the duplicated PP2Cs in grape.

**Additional file 19**

Ka/Ks analysis for GmPP2Cs and its corresponding orthologous genes in grape.

**Additional file 20**

Number of VvPP2C orthologous genes in soybean.

**Additional file 21**

Sequence similarity between GmPP2Cs and its corresponding AtrPP2Cs.

**Additional file 22**

Sequence similarity between GmPP2Cs and its corresponding NcPP2Cs.

**Additional file 23**

The FPKM values of GmPP2Cs in seven tissues.

**Additional file 24**

PCC of the FPKM values for the duplicated GmPP2Cs pairs.

**Additional file 25**

The constructed co-expression modules in soybean by hierarchical clustering and dynamic tree cut.

**Additional file 26**

Module information of GmPP2C through WGCNA analysis.

**Additional file 27**

Scatterplot of gene significance and module membership in certain significant modules.

**Additional file 28**

Number of the responsive-regulatory elements in the promoter regions of GmPP2Cs.

**Additional file 29**

Regulatory relationship between TFs and duplicated GmPP2Cs through the MERLIN+Prior method.

**Additional file 30**

Regulatory relationship between TFs and duplicated GmPP2Cs through the MERLIN+Prior method.

**Additional file 31**

PCR primers used in the study.

**Abbreviations**

ABA: Abscisic Acid; CV: Coefficient of variation; DSPTP: Dual-specificity phosphatase; FPKM: Fragments per Kilobase Million; GCN: Gene co-expression network; GO: Gene ontology; GRN: Gene regulatory network; HMM: Hidden Markov Model; Ka: Nonsynonymous substitution rate; Ks: Synonymous substitution rate; MeJA: Methyl Jasmonate; PP2C: Protein phosphatases 2C; PCC: Pearson Correlation Coefficient; PK: Protein kinase; PP: Protein phosphatase; PPM: Phosphoprotein metallophosphatase; PPP: Phosphor-protein phosphatase; PTP: Protein Tyr phosphatase; SA: Salicylic Acid; Ser: Serine; STP: Ser/Thr phosphatase; TF: Transcription factor; Thr: Threonine; Tyr: Tyrosine; WGCNA: Weighted gene co-expression network analysis

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Not applicable.

**Authors’ contributions**

KF, JH and WXL designed the research. KY, ZM, YF and ZL performed the experiments. KF, YC, WWL, YZ and JL analyzed the data. KF, JH and WXL wrote the paper with contributions from all the authors. All authors have read and approved the manuscript.

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**Availability of data and materials**

All data in this study are included in this article and its additional files.
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