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New insights into the evolution of SPX gene family from algae to legumes; a focus on soybean

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Abstract

Background: SPX-containing proteins have been known as key players in phosphate signaling and homeostasis. In Arabidopsis and rice, functions of some SPXs have been characterized, but little is known about their function in other plants, especially in the legumes.

Results: We analyzed SPX gene family evolution in legumes and in a number of key species from algae to angiosperms. We found that SPX harboring proteins showed fluctuations in domain fusions from algae to the angiosperms with, finally, four classes appearing and being retained in the land plants. Despite these fluctuations, Lysine Surface Cluster (KSC), and the third residue of Phosphate Binding Sites (PBS) showed complete conservation in almost all of SPXs except few proteins in *Selaginella moellendorffii* and *Papaver somniferum*, suggesting they might have different ligand preferences. In addition, we found that the WGD/segmentally or dispersed duplication types were the most frequent contributors to the SPX expansion, and that there is a positive correlation between the amount of WGD contribution to the SPX expansion in individual species and its number of EXS genes. We could also reveal that except SPX class genes, other classes lost the collinearity relationships among Arabidopsis and legume genomes. The sub- or neo-functionalization of the duplicated genes in the legumes makes it difficult to find the functional orthologous genes. Therefore, we used two different methods to identify functional orthologs in soybean and Medicago. High variance in the dynamic and spatial expression pattern of GmSPXs proved the new or sub-functionalization in the paralogs.

Conclusion: This comprehensive analysis revealed how SPX gene family evolved from algae to legumes and also discovered several new domains fused to SPX domain in algae. In addition, we hypothesized that there different phosphate sensing mechanisms might occur in *S. moellendorffii* and *P. somniferum*. Finally, we predicted putative functional orthologs of AtSPXs in the legumes, especially, orthologs of AtPHO1, involved in long-distance Pi transportation. These findings help to understand evolution of phosphate signaling and might underpin development of new legume varieties with improved phosphate use efficiency.

Keywords: Phosphate homeostasis, Evolution, Gene family, Legumes

Background

Phosphorus (P) as an essential macronutrient serves as a structural element for many organic compounds, involved in multiple biosynthetic and metabolic processes [1, 2]. P containing molecules play a central role in various physiological processes, including respiration,

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photosynthesis, membrane transport, regulation of enzyme activity, oxidation-reduction reactions and signal transduction throughout plant growth, and development [3, 4]. Therefore, plants have evolved a number of mechanisms to ensure that P is readily available for all these processes. In particular, a wide range of responses are induced by phosphate (Pi) starvation [5, 6]. The regulation occurs at both transcriptional and posttranscriptional levels and many components of the regulatory network are known. The central regulator of the Pi starvation response and signaling network is the MYB transcription factor, AtPHR1 or OsPHR2 [7–9]. The PHR factors are negatively regulated through interaction with SPX domain proteins, which serve as sensors of P-status of the cells. In high P availability, inositol polyphosphates (PP-InsPs) bind to the basic surface of SPX domain proteins and facilitate their binding to PHR. This interaction may sequester PHR1 in the cytosol or prevent its association with DNA in the nucleus [10]. In low P supply, low availability of PP-InsPs-SPX results in the release of PHR1 to translocate to nucleus and to activate Pi starvation induced (PSI) genes [8]. Additionally, SPX domain proteins were shown to be involved in nitrate-phosphate signaling crosstalk in rice where nitrate-dependent interaction with NRT1.1B caused ubiquitination and degradation of OsSPX4 and consequently translocation of OsPHR2 and OsNLP3 into nucleus to induce PSI genes and nitrate inducible genes, respectively [11].

Despite the importance of SPX domain proteins in Pi signaling and nitrogen-dependent phosphate homeostasis, the functionality of all these proteins is still unclear. SPX domain proteins are important components of plant Pi homeostasis and can be divided into four classes based on the presence of extra domains: while class 1 only includes SPX domain, other three classes (SPX-EXS, SPX-MFS, SPX-RING), contain extra EXS, MFS, or RING domains, respectively [6]. There are four and six members of the SPX class 1 in *Arabidopsis* and rice, respectively [12, 13] as AtSPX3 and OsSPX1 act as negative regulators of Pi starvation signaling [12, 13]. Indeed, AtSPX1, localized in the nucleus, has a high binding affinity for AtPHR1 under high P condition and prevents it from activation of the downstream Pi starvation-induced (PSI) genes [8]. The rice OsSPX4 protein involved in the nitrate dependent regulation of Pi uptake [11] also belongs to this class. The most functional variation was observed in the EXS class members, including AtPHO1 and AtPHO1;1 involved in long-distance Pi transport from roots to shoots [14, 15], AtPHO1;4 with a role in response of hypocotyls to blue light [16], seed size and flowering [17–19] and AtPHO1;10 being induced by numerous stresses, such as local wounding [20, 21]. The Major Facilitator Superfamily (MFS) domain confers

transport activity, therefore, SPX-MFS class are involved in both transport and signaling [22]. Finally, members of SPX-RING class are also called Nitrogen Limitation Adaptation (NLA) proteins due to their first identified role in nitrogen starvation resistance [23].

Recently, two other classes of SPX proteins, SPX-SLC and SPX-VTC, were characterized in algae as involved in polyphosphate synthesis and its transportation into vacuoles [24]. These two classes seem to be lost during the evolution of plants with shifting the type of phosphate storage from polyP in algae to Pi in the later-diverging Streptophytes [24]. It seems there have been some extra domains fused with SPX domain that might have been lost during the evolution of SPX proteins and that have not been comprehensively explored yet [25].

Legumes (Fabaceae) are the second most important family of crop plants economically [26]. Characterization of the SPX gene family in legumes can be helpful to gain insights into mechanisms of Pi homeostasis and thus underpin development of P efficient varieties. In this study, we performed a comprehensive analysis of SPX proteins from several legume crops (soybean, alfalfa, and common bean), and compared with species of more basal taxonomic groups such as mosses (*Phacomitrella patens*), liverworts (*Marchantia polymorpha*), lycophytes (*Selaginella moellendorffii*), basal angiosperms (*Papaver somniferum*, *Amborella trichopoda*, and *Nymphaea colorata*), Rhodophytes (*Cyanidioschyzon merolae*, *Galdieria sulphuraria*, and *Chondrus crispus*), chlorophytes (*Chlamydomonas reinhardtii* and *Ostreococcus lucimarinus*), and charophytes (*Chara braunii*). We analyzed SPX protein evolution through phylogenetic analysis, conserved motif changes, and identification of ancestral motifs. In addition, because of only a partial functional characterization of SPX in legumes [27–31], we identified their functional orthologs with well-characterized SPXs from *Arabidopsis thaliana*. Since sequence-based orthology identifications alone have weakness in the one-to-many or many-to-many orthologs, expressologs identification was used as a complementary approach for functional ortholog identification [32]. With the combination of these two methods, we identified the functional orthologs of key regulators AtPHO1 (*GLYMA_02G003700*, *GLYMA_10G004800*), AtSPX4 (*GLYMA_06G069000*), AtPHO1;H10 (*GLYMA_02G110600*), and AtNLA2 (*GLYMA_19G203000*) in soybean. Also, we assessed the *GmSPX* gene expression in response to phosphate and nitrogen deficiency to compare their expression patterns with well-characterized SPXs from rice and *Arabidopsis*. In addition, we identified novel domains in SPX proteins of algae and functionally characterized SPX proteins in soybean and Medicago.

Results

Identification of SPX domain proteins from algae to legumes

While in several plant species four families of SPX proteins were characterized, much less is known about these proteins in legumes: in soybean and common bean just 10 and 3 members of class 1 were characterized and no SPX proteins in *M. truncatula*. Therefore, we intended to characterize this protein family in these legume species and set it into evolutionary context by analysis of SPX proteins from algae and basal plants. Sequences of SPX proteins were obtained by BLASTP searches at EnsemblPlants from the legumes (*G. max*, *P. vulgaris*, and *M. truncatula*), moss (*P. patens*), liverwort (*M. polymorpha*), lycophyte (*S. moellendorffii*), basal angiosperms (*P. somniferum*, *A. trichopoda*, and *N. colorata*) rhodophytes (*C. merolae*, *G. sulphuraria*, and *C. crispus*), chlorophytes (*C. reinhardtii* and *O. lucimarinus*), and charophytes (*C. braunii*) protein databases using full-length amino acid sequences of SPXs from Arabidopsis (20 proteins). After removing sequences lacking the SPX domains and redundant and partial sequences, we compiled all SPX proteins in the latest version of protein database in EnsemblPlants for these 15 species. Some proteins were shorter than 200 aa and were excluded from further analyses, including four short proteins of soybean (*GLYMA_12G154800*, *GLYMA_10G097000*, *GLYMA_09G098200*, *GLYMA_20G032200*), two partial proteins of common bean (*PHAVU_010G0720001g*, *PHAVU_010G0720000g*), one protein of *M. truncatula* (*MTR_8g058603*). In addition, we excluded one protein of *M. truncatula* (*MTR_0262S0060*), where its corresponding gene is located on a scaffold but not chromosomes, and one protein of a common bean (*PHAVU_007g1245000g*), which had different structure from other SPX genes. Finally, 34 SPX proteins in *G. max*, 19 in *M. truncatula*, 17 in *P. vulgaris*, 22 in *P. patens*, 10 in *M. polymorpha*, 2 in *C. merolae*, 4 in *G. sulphuraria*, 2 in *C. crispus*, 5 in *C. reinhardtii*, 2 in *O. lucimarinus*, 4 in *C. braunii*, 42 in *P. somniferum*, 11 in *A. trichopoda*, 16 in *N. colorata*, and 31 in *S. moellendorffii* were identified (Supplemental Table S1). Furthermore, the proteins were classified into the four subfamilies based on their additional domains. Interestingly, in some algae and basal plants, we found extra domains that have not been previously reported (Fig. 1). Totally, among these species, class EXS was with 88 proteins the largest, followed by SPX class with 48 proteins and MFS and RING classes containing 29 and 26 proteins, respectively. Subsequently, the corresponding SPX genes in soybean, *M. truncatula* and

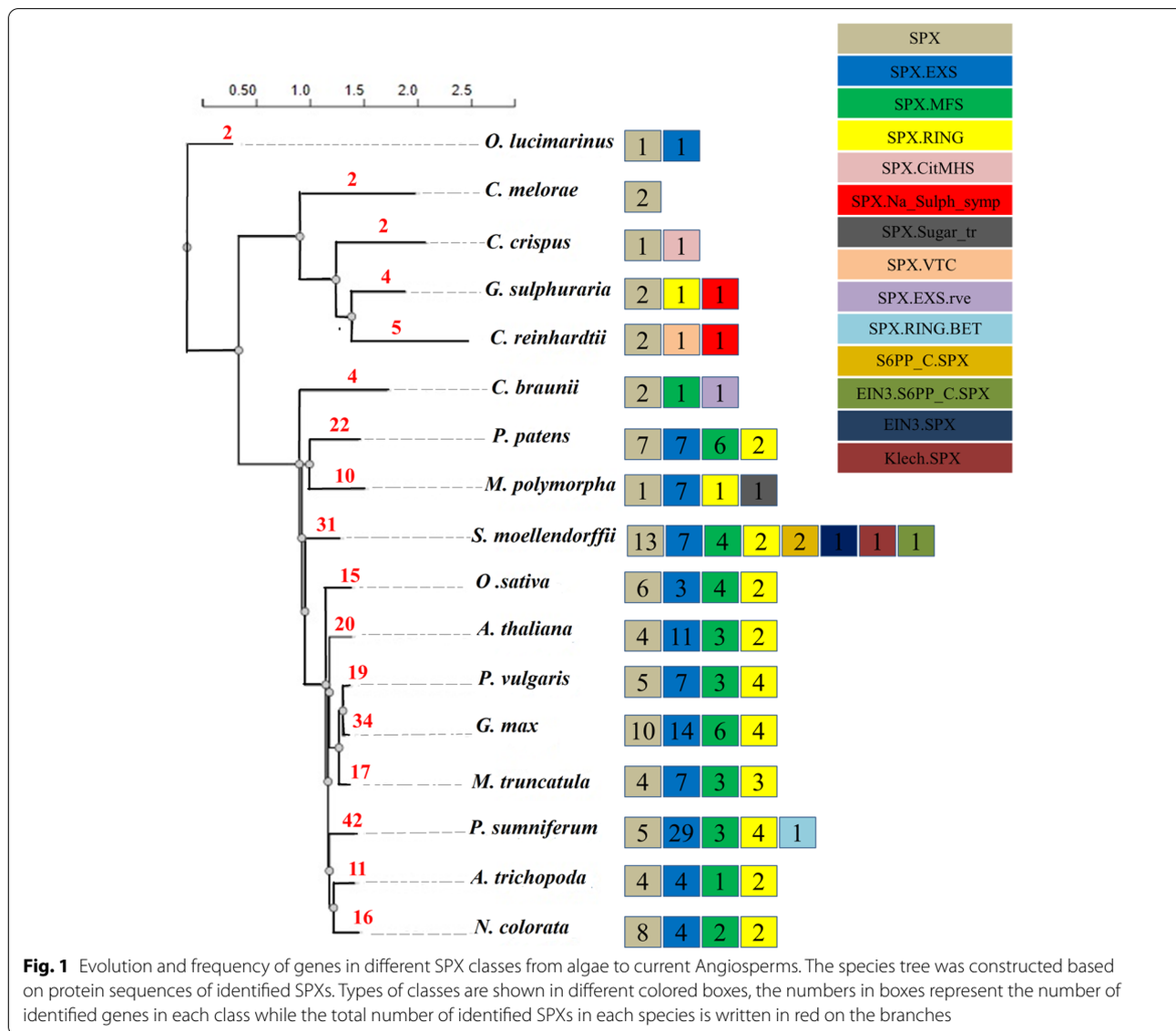
common bean were named in each subfamily based on their chromosomal positions (Supplemental Table S1).

As can be seen in the Fig. 1, all basal and current angiosperms possess only the four main classes of SPX proteins. On the other hand, some additional domains were observed in liverwort, lycophyte, and algae based on Pfam and CDD scanning of sequences; SPX-VTC (vacuolar transporter chaperone), EIN3-SPX (Ethylene intensive 3), SPX-CitMHS (Citrate transporter), SPX-Na_sulph_symp (sodium sulphate symporter), SPX-RING-BET (Bromodomain extra-terminal-transcription regulation), S6PP_C-SPX (Sucrose-6F-phosphate phosphohydrolase C-terminal), EIN3-S6PP_C-SPX, Kelch-SPX (Galactose oxidase), SPX-EXS-rve, and SPX-Sugar_tr (Fig. 1). The exact roles of these additional domains in the basal plants and algae are not completely known. It was previously reported that in some SPX proteins, SPX domain was located at C terminal instead of N terminal [33]. Indeed, we observed this structure in 4 different classes in *S. moellendorffii*, including EIN3-S6PP_C-SPX, Kelch-SPX, EIN3-SPX, and S6PP_C-SPX.

Predicted physicochemical and biochemical parameters of these SPX proteins in legume crops are listed in Supplemental Table S1. Indeed, members of the same subfamily have similar properties. The most variation in physicochemical parameters was observed in EXS class, while MFS class was the most similar. For example, lengths of all SPX-MFS proteins in the three species ranged from 691 to 700 aa, but the corresponding SPX-EXS proteins ranged from 475 to 1570 aa with the MtEXSs having the largest proteins in comparison with soybean and common bean. SPX-EXS and SPX-RING classes have the highest isoelectric point (pI), above 9 and 8, respectively. The calculated values for aliphatic index of SPX proteins show that the SPX-MFS subfamily have most thermostability, with a range of 105 to 111. GRAVY value (grand average of hydrophobicity) is the sum of the hydrophobicity values of all amino acids divided by the protein length. Except for the proteins in the SPX-MFS subfamily, nearly all of the GmSPXs are hydrophilic, with a GRAVY value less than 0. Subcellular localization prediction performed with Wolf PSORT revealed that most of the GmSPX proteins are located in the plasma membrane or endomembrane system, followed by nucleus and chloroplast. In PSORT results, all members of SPX-EXS and SPX-MFS subfamilies were located in the plasma membrane, and all members of SPX-RING were located in nucleus, corresponding to the known functions of representatives of these subfamilies in Arabidopsis.

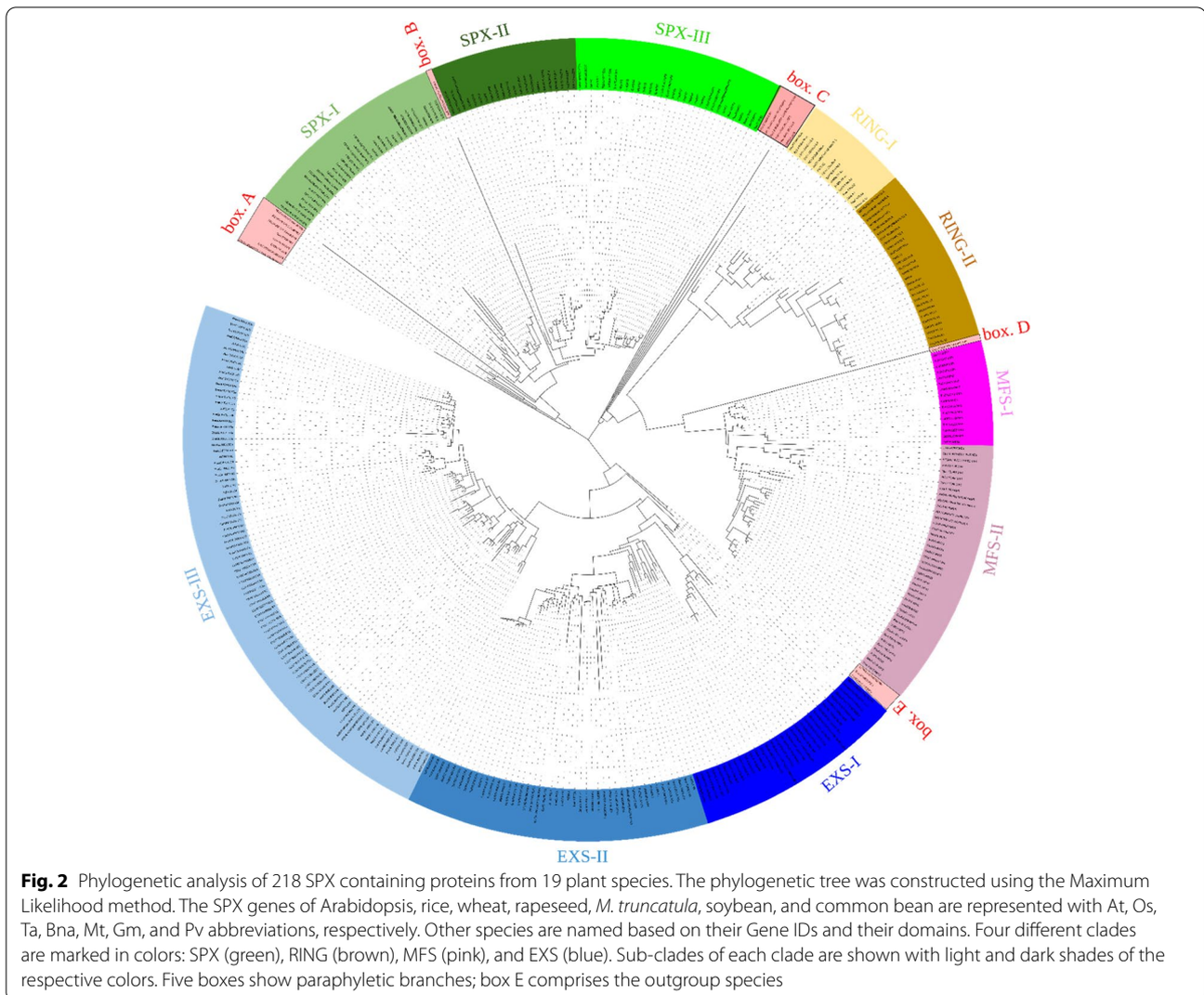
Phylogenetic tree

Multiple alignment of the SPX protein sequences from soybean, *M. truncatula*, common bean, Arabidopsis, rice,



wheat, rapeseed, *A. trichopoda*, *C. braunii*, *C. reinhardtii*, *C. crispus*, *C. melorae*, *G. sulphuraria*, *M. polymorpha*, *N. colorata*, *O. lucimarinus*, *P. somniferum*, *P. patens*, and *S. moellendorffii*, as well as proteins from mouse, human, and *Caenorhabditis elegans* as an out-group, followed by phylogenetic analysis revealed four distinct clades of SPX proteins, SPX, EXS, MFS, and RING (Fig. 2). This topology and distinct separation of four classes are consistent with previous studies on SPX gene family [3, 12, 13, 27, 34]. SPX and EXS sequences formed two distinct clades, while MFS and RING along with box C (*OSTLU26654.EXS*, *CHC T00007225001.SPX.CitMHS*, *CHLRE 09g251650V5.SPX.Na_Sulph_symp*, *C5167 020395.NLA.BET*, *Gsu16460.SPX.NLA*, and *CMP022C.SPX*) and box D (*Gsu35240.SPX.Na_sulph_symp*) have diverged from a

common ancestor and form the third major clade. SPX clade was divided into three sub-clades; SPX-I, SPX-II, and SPX-III. SPX-II and SPX-III are specific to the basal and current angiosperms and the proteins in these two sub-clades are homologs of AtSPX3 and ATSPX1/2, respectively. On the other hand, SPX-I is comprised from homologs of the basal plants (lycophytes, liverwort, moss) and algae and few proteins from the basal and current angiosperms, all being homologs of AtSPX4. Proteins in box A and in box B could be ancient homologs for SPX-I and SPX-II/III, respectively. Likewise, EXS clade was divided into three sub-clades; EXS-I is specific to lower plants (*S. moellendorffii*, *M. polymorpha*, and *P. patens*), EXS-II is a mixed group from monocots, eudicots, and basal angiosperms, all homologs of AtPHO1



and AtPHO1;H1, and EXS-III contain eudicots and the basal angiosperms without any genes of monocots. The outgroup genes used in this study were grouped in box E clustered with EXS clade. Overall, topology of EXS class is consistent with He et al., (2013), in that basal plants (lycophytes and moss) EXS homologs were grouped separately from the angiosperms, and also with the previous reports on EXS genes that monocots only possess homologs for AtPHO1 and AtPHO1;H1 [6, 24, 35].

Box C with ancient genes for both MFS and RING and box D as sister for MFS class together with MFS and RING clades seem to have evolved from a common ancestor. MFS homologs in monocots specifically grouped in MFS-I, while MFS-II contained all MFS orthologs from the other species. This could suggest that differentiation among MFS proteins has occurred after the divergence of monocot and dicots from a common ancestor. Similarly, RING clade was divided into two

sub-clades, but both contained RING orthologs from all of species; RING-I was grouped with the ancestor from *P. patens*, while RING-II included *S. moellendorffii* orthologs as its sister. The overall tree topology is very similar to results of Wang et al (2021), who investigated SPX gene family in chlorophytes and streptophytes, with focus on algae.

Protein motifs gain and loss in SPX family throughout evolution

Conserved protein motifs were predicted using MEME program for each SPX protein class and all species (Additional file 1: Figs. S1 to S5). This analysis may explain when different classes of SPX proteins have appeared and how motifs were gained or lost in each class during the evolution. The ancestral motifs in SPX domains such as motifs 3, 4, 2, and 1 seem to originate from red algae (Additional file 1: Fig. S1). There is a high fluctuation of

motif composition during the evolution. Some motifs are species specific like motifs 13, 14, and 19 that are present only in legumes, probably arising after legume whole-genome duplication event. The most variability in the motif composition was observed in *S. moellendorffii* with some specific motifs like 8, 15, and 18. The lengths of proteins in angiosperms were very similar but shorter than in the basal plants. The EXS domain was detected only in *O. lucimarinus* with 9 motifs - 9, 6, 5, 2, 3, 11, 4, 10, and 1 (Additional file 1: Fig. S2). Almost all these motifs have been retained during the evolution as ancestral motifs. In addition, some other motifs appeared in *C. braunii* such as 15, 7, 16, 20, 12, and 8, suggesting they were present in the common ancestor of Chlorophyta and Streptophyta. Although Wang et al. [24] reported one SPX-MFS in *M. polymorpha* genome, we could not find an intact SPX-MFS domain, but SPX-Sugar_tr domain with a highly similar motif composition with other MFSs was identified (Additional file 1: Fig. S3). As it has previously been reported, *PHT5* genes in *B. napus* have SPX domain connected to overlapping MFS and Sugar_tr domains [36], however, we only found SPX and Sugar-tr domains in *M. polymorpha* genome. The first SPX-MFS protein was observed in *C. braunii* with 18 common motifs with other species. Two newly observed motifs in *P. patens*, motifs 16 and 13, probably have evolved by dispersed duplication in *P. patens* and have been retained in all basal and current angiosperms. Interestingly, other five MFSs in *P. patens*, without the motifs 16 and 13, have been no longer found in angiosperms.

The evolutionary oldest NLA has been detected in *G. sulphuraria* and it was retained during the course of evolution of current angiosperms, but was not found in other Rhodophytes or Chlorophytes. In fact, the only NLA identified in *G. sulphuraria* just showed two motifs in common with other species, motifs 2 and 3 (Additional file 1: Fig. S4). Therefore, these motifs could be considered as ancestral motifs of NLA class which then further evolved by dispersed duplication in *M. polymorpha*, adding motifs 8, 7, 1, and 6 into the ancestral domains. One NLA in *P. somniferum* underwent dispersed duplication and gained motif 10 that has only been retained in the core eudicots, while two NLAs in *S. moellendorffii* segmentally duplicated and gained two specific motifs 13 and 19. Motif 16 was just observed in legume genomes that might evolved after legume whole-genome duplication (WGD) event. The most variability in motif composition of NLA class was observed in *P. somniferum*. Motif compositions in the new identified classes showed a high variation and it was impossible to find their ancestral motif (Additional file 1: Fig. S5). However, it could be concluded that SPX-Na_Sulph_sym and SPX-CitMHS with high similarity in the motif composition, probably

have similar origin and function. In summary, during the evolution different duplication events added new motifs to the ancestral motifs and other motifs specifically appeared in individual species to acquire new functions.

Consensus sequences of SPX domains from algae to eudicots

We then predicted conserved motifs among all identified SPXs (Additional file 1: Fig. S6). There are four conserved motifs in SPX class members, among them two motifs, 2 and 4, are common in the almost whole span of SPXs. Therefore, we can hypothesize that these two motifs have an important role for all SPXs. Afterwards, consensus sequences of these two motifs were constructed across all phyla (algae, charophytes, liverwort, bryophytes, lycophytes, basal angiosperms, and current angiosperms) and also across each class (SPX, EXS, MFS, RING, and new identified classes) (Additional file 1: Figs. S7-S10). Motif 4 is 29 aa in length and was present in all SPX proteins except the following ten: C5167_005902.EXS, C5167_032842.EXS, C5167_043562.EXS, C5167_043565.EXS, C5167_003186.NLA, C5167_046257.NLA, SELMODRAFT_419593.SPX, SELMODRAFT_419593.SPX, OsSPX4 and PvPHO1. Five amino acid residues, number 5, 9, 15, 19, and 24, were almost 100% conserved, except the fifth residue in *C. braunii* (Additional file 1: Fig. S7). Regarding conservation in different classes (Additional file 1: Fig. S8), the leucine (residue 9) was completely conserved in the EXS, MFS, RING, and new identified classes, then the phenylalanine (residue 19) was completely conserved in EXS and MFS classes, but SPX class had some members with different residues in these five positions with a very high overall conservation in this class. In addition, each class had other conserved residues, suggesting special functions.

Motif 2 is 21 aa long and was absent in CHLRE_02g111650v5.SPX, AMTR_s00106p00066860.SPX, NC1G0101580.SPX, C5167_011965.SPX, Gasu_57230.SPX, C5167_043539.EXS, SELMODRAFT_450458.EXS, SELMODRAFT_431864.SPX, SELMODRAFT_419593.SPX, and only one protein from the current angiosperm, PvPHO1;5, which is a partial protein. This motif exhibited more conserved residues at positions 1, 7, 8, 14, 15, 16, 17, 18, 20, and 21. Residue 17 was completely conserved in the all proteins containing motif 2 and residues 14, 18, and 21 were conserved in the all proteins except a few in *S. moellendorffii* and *P. somniferum* showing different residues instead of lysine (Additional file 1: Fig. S9). The lysine residues 14, 17, and 21 form a Lysine Surface Cluster (LSC), and were found to interact with sulfate in the crystal structure of human phosphate transporter XPR1, and to be a part of a larger binding site for PP-InsP [9]. Consequently, in the

different classes of the SPX proteins (Additional file 1: Fig. S10), some of the 10 conserved positions were completely conserved such as K1, N8, KILKK (14 to 18) in RING and MFS, K18 in SPX, K21 in RING, and N8, I15, K18, as well as K21 were completely conserved across the new identified classes. Overall, these two motifs were conserved in all but a few proteins from *S. moellendorffii* and *P. somniferum*, PvPHO1;5, and OsSPX4, implying that they might possibly interact with InsP/PP-InsP in a different manner, as previously reported for OsSPX4 [9]. In addition, different conserved residues in different classes could suggest that they may have different phosphate-containing ligand or different levels of Pi in cells.

Expansion pattern of SPX genes and collinearity analysis

To pinpoint the expansion modes in the land plants, we investigated duplication types in basal and current angiosperms, liverwort, hornwort, and *S. moellendorffii* (Fig. 3 and Supplemental Table S2). Taken together, WGD, segmental, and dispersed duplications contributed most to the SPX gene family expansion. The expansion patterns in soybean, *P. somniferum*, *N. colorata*, and *S. moellendorffii* mostly arose from WGD/segmental duplication type. However, *S. moellendorffii* did not have any WGD events, therefore, its expansion and unique SPX classes must have arisen through local or segmental gene duplication [37]. WGD/segmental duplication type did not participate in the SPX expansion in *A. trichopoda* and *M. polymorpha* genomes and it only resulted in one duplicated block in *P. patens* genome. In these three species,

SPX expansion were affected mostly by dispersed duplication type. The high number of WGD/segmental types of duplication in *S. moellendorffii*, soybean, and *P. somniferum* can shed light on the reason of high variation of gene family sizes in the closely related plants.

To get more information about evolutionary process of genes, collinearity analysis can provide information about conserved genomic regions of genes in different species [38]. Synteny relationship among two or a set of genes from two species means that they located in the same chromosome [39], but collinearity is a specific form of synteny with conserved gene order [40]. Collinearity analysis was conducted in three steps; 1. across *P. somniferum*, *N. colorata*, rice, Arabidopsis, and three legumes 2. Among *P. somniferum* and *N. colorata*, *P. patens*, and *S. moellendorffii* and 3. Among legumes.

Collinearity analysis among legume crops, Arabidopsis, rice, and two basal angiosperms; *P. somniferum* and *N. colorata* discovered 121 collinear blocks (Fig. 4, Supplemental Table S3); 30 blocks in Gm/Pv, 23 blocks in Gm/Mt, 15 blocks in Gm/Gm, 14 blocks in Ps/Ps, 10 blocks in Gm/At, 6 blocks in Nc/Nc and Pv/Mt, 3 blocks in Ps/Nc, Mt/At, Pv/At, and Os/Os, 2 blocks in Ps/Gm, and 1 block in At/At, Pv/Pv, Ps/Mt, and Mt/Mt. Rice as the only monocot in this analysis did not show any collinearity relationship for SPX gene family with other species.

Collinear SPX genes among *P. somniferum*, *S. moellendorffii*, *N. colorata*, and *A. trichopoda* were predicted (Supplemental Table S3). *S. moellendorffii* did not show any collinearity relationship with other species, while

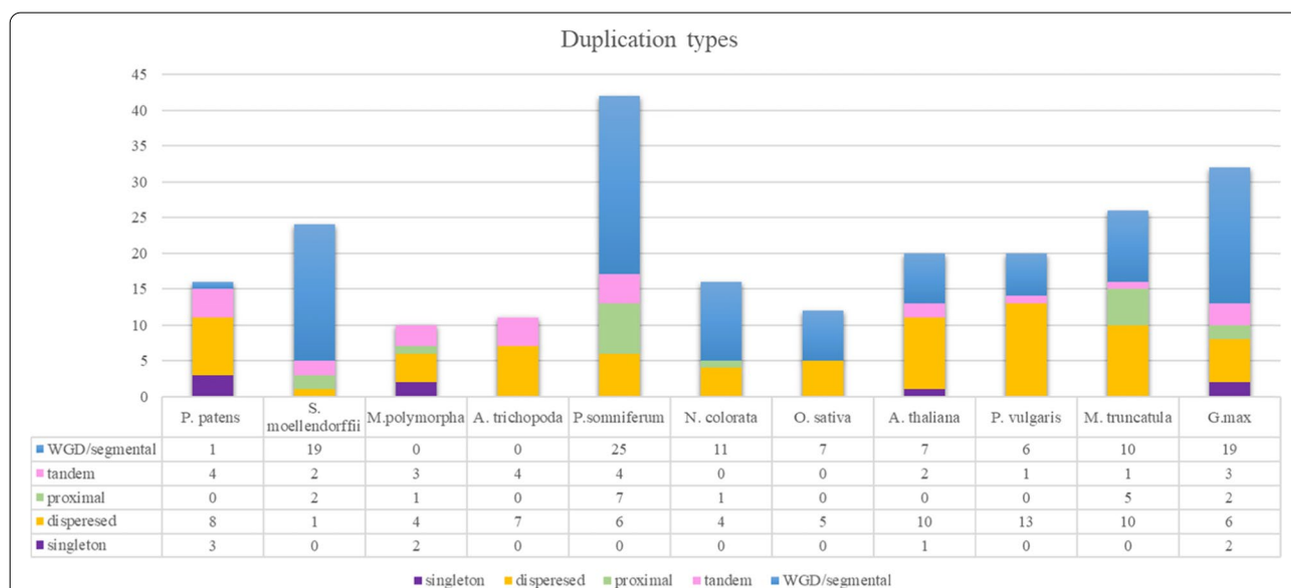
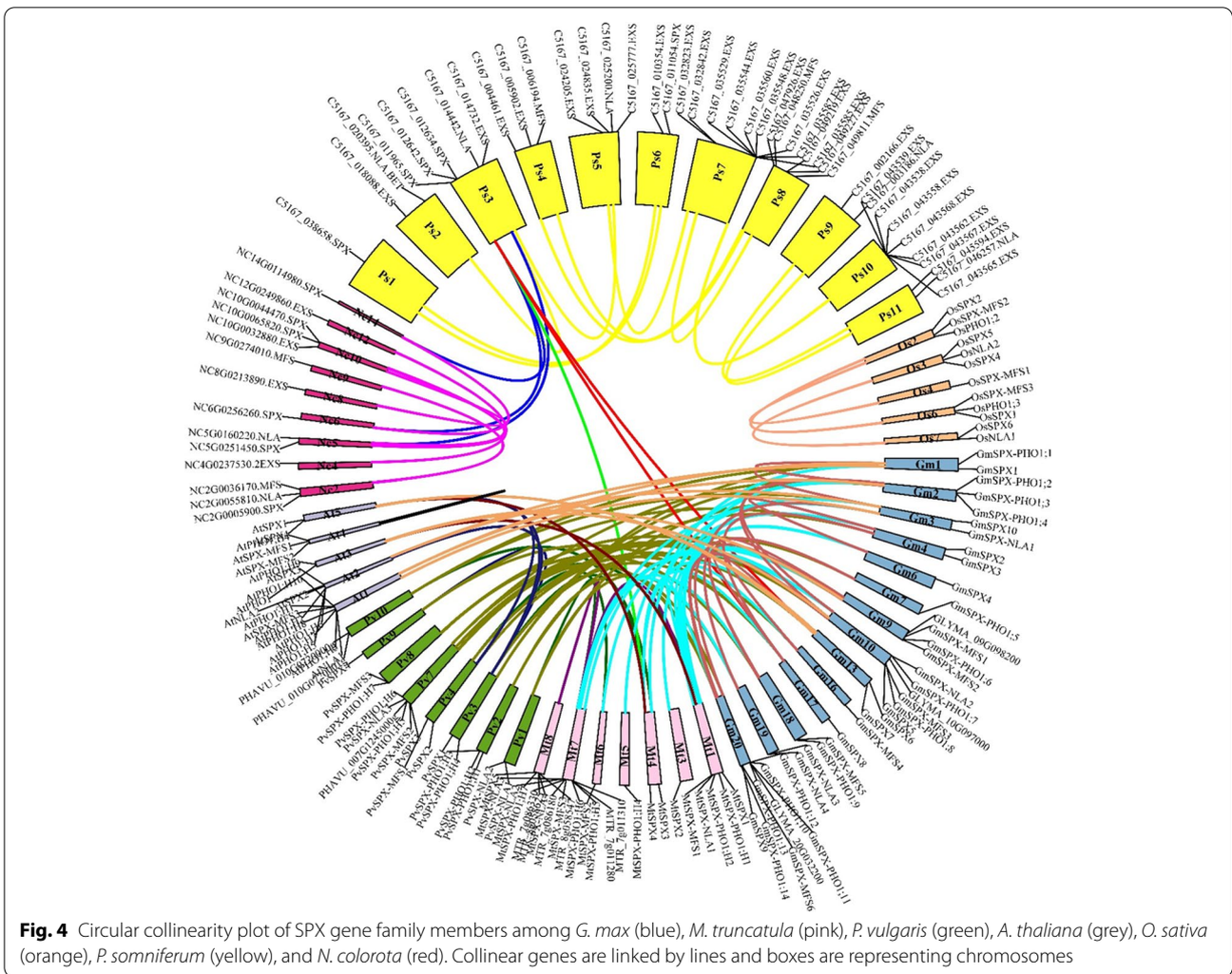


Fig. 3 SPX gene family expansion from algae to the current Angiosperms. Duplication event types were predicted in the *P. patens*, *S. moellendorffii*, *M. polymorpha*, *A. trichopoda*, *P. somniferum*, *N. colorata*, *O. sativa*, *A. thaliana*, *P. vulgaris*, *M. truncatula*, *G. max*

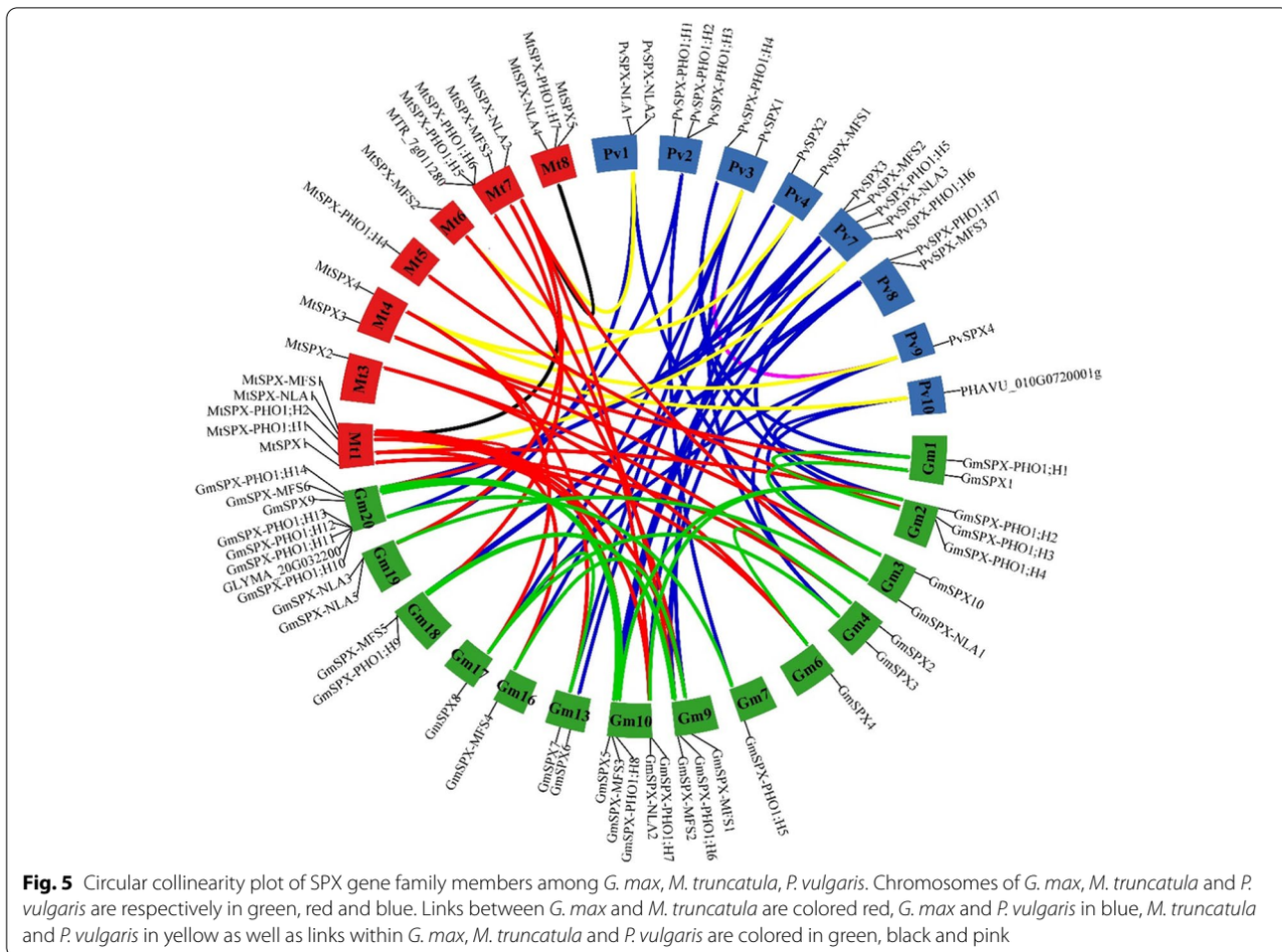


N. colorata and *P. somniferum* had the most inter species collinear relationships [14]. The most intra-genome collinear relationships were found in *P. somniferum* [14] and *S. moellendorffii* [10]. The collinear analysis was performed also for the three legume crops (Fig. 5, Supplemental Table S3). Of the 34, 19, and 17 SPX genes in soybean, *M. truncatula*, and common bean 32, 14, and 15 genes participated in collinear blocks. In total, 78 collinearity blocks between these plant species were discovered. A high level of collinearity relationships was found at 27/30 SPX genes in soybean/common bean and 19/23 SPX genes in soybean/*M. truncatula*, while the corresponding figure for *M. truncatula*/common bean was 6/7. However, just 15, 7, and 2 collinearity blocks were found in soybean/soybean, *M. truncatula* /*M. truncatula*, and common bean/common bean groups. All in all, after these three collinearity analyses, we concluded that inter-species collinearity patterns among basal angiosperms and among current angiosperms have changed. Across

basal angiosperms, SPX class had the least inter-species collinearity, while among Arabidopsis and legumes, SPX showed the most inter-collinearity relationships. It can be concluded that except in SPX class, collinearity in the other classes has been lost.

Evolution of Cis-acting elements from algae to eudicots

Transcription factors bind to the cis-acting elements (CREs) in the promoter and regulate the transcription of corresponding genes [41]. Therefore, genes with similar expression patterns may contain the same regulatory elements in their promoters [27]. To explore whether transcription factor binding sites have evolved together with the coding regions of SPX genes, 1.5 kb upstream of the transcriptional start sites of all identified SPXs were downloaded and analyzed using PlantCARE database. In total, 124 CREs were detected (Supplemental Table S4) that can be classified in three major groups: responsive to abiotic stresses (drought,



low temperature, hypoxia, wounding, defense, and stress), hormones (gibberellin, abscisic acid (ABA), salicylic acid (SA), ethylene, methyl jasmonate (MeJA), and auxin), and development-related elements (endosperm, meristem, MYB, and zein metabolism regulation). After the essential elements in promoter like TATA-box and CAAT-box, the most highly represented cis-acting elements were those involved in response to MeJA (CGTCA-motif and TGAG-motif) and ABA (ABRE and ARE). Since SPX genes play role in P homeostasis we were interested in a presence of P1BS motif (PHR1-binding site important in P starvation response) in the respective promoters. However, the P1BS motif is not included in the PlantCare database. Therefore, additionally, we performed promoter analysis using PlantPan3 for *C. reinhardtii* and *G. max* to figure out if this motif existed in both ancient and current species or it just appeared in the new angiosperms. The P1BS motif was found in all five SPX genes in *C. reinhardtii* and in almost all *GmSPXs* except *GmSPX.PHO1;3* and *GmSPX.PHO1;13* (Supplemental Table S4).

Looking for evolutionary pattern in these cis-acting elements, we performed hierarchical clustering on principal components (HCPC) using FactMineR-package. The HCPC grouped the genes into three clusters (Additional file 1: Fig. S11). Almost all SPXs from the current angiosperms fell into cluster 1 along with 2 SPXs of *G. sulphuraria* and few SPXs from basal angiosperms (Table 1, Additional file 1: Fig. S11). Cluster 2 comprised mostly genes from basal angiosperms and few members of the current angiosperms, as well as all SPXs of *C. reinhardtii* and two SPXs from *G. sulphuraria*. Cluster 3, the smallest cluster, had 12 genes mostly from *P. patens* and just one SPX of the current angiosperms, *AtPHO1;H5*. Trying to find an evolutionary pattern across these clusters, we found out that they showed different frequencies of two MeJA responsive elements, TGACG and CGTGA motifs, that in cluster 3 all genes, in cluster 2 around 93%, and in cluster 1 only around 62% of genes possessed these two elements (Table 1). Besides, we extracted the most enriched CREs in each cluster

Table 1 Number of genes having MeJA and ABA responsiveness elements in their promoter sequence

Clusters	TGACG-motif	CGTCA-motif	ABRE	ARE	Number of SPXs of each species	Total
Cluster 1	57	57	59	72	16 At, 27 Gm, 12 Mt, 15 Pv, 11 Pp, 1 Chb, 2 Gsu, 4 Nc, 2 Pp, 1 Mp,	91
Cluster 2	55	55	32	47	5 CHLRE, 2Gsu, 7Mp, 13 Ps, 2 Chb, 7 Nc, 12 Pp, 3 Sm, 2 Mt, 3 Gm, 3 At	59
Cluster 3	12	12	11	12	3 Pp, 1 Mp, 1 Sm, 1 At, 6 Ps,	12

to visualize frequencies of these elements across clusters. As can be seen in the Additional file 1: Fig. S12, CREs involved in the developmental processes (CCG TCC motif, CCGTCC box, A-box) and stress response (DRE core, MYB recognition site, CCAT box) were significantly higher in cluster 2 than in the other clusters. Cluster 1 had higher frequency of two hormone responsive elements, TCA (salicylic acid responsive elements) and ERE (Ethylene-responsive elements) in comparison to the other clusters. Overall, it seems that during the evolution of angiosperms, SPX promoters were enriched by stress responsive elements and hormonal responsive elements, especially ERE and TCA.

Finding cis-elements responsive to stresses and hormones triggered two major questions, how SPX genes are regulated by phytohormones and whether SPX genes and hormone and stress responsive genes show overlaps in response to phosphate deficiency. To address these questions we reanalyzed three datasets from publicly available databases related to these conditions. The first data was for Arabidopsis response to ABA, salicylic acid (SA), jasmonic acid (JA) and their combination at 3 and 24h after treatment (GSE28600). Among these three phytohormones, ABA caused changes in of almost all SPX expressions, especially for members of SPX class (Fig. 6A). However, ABA in

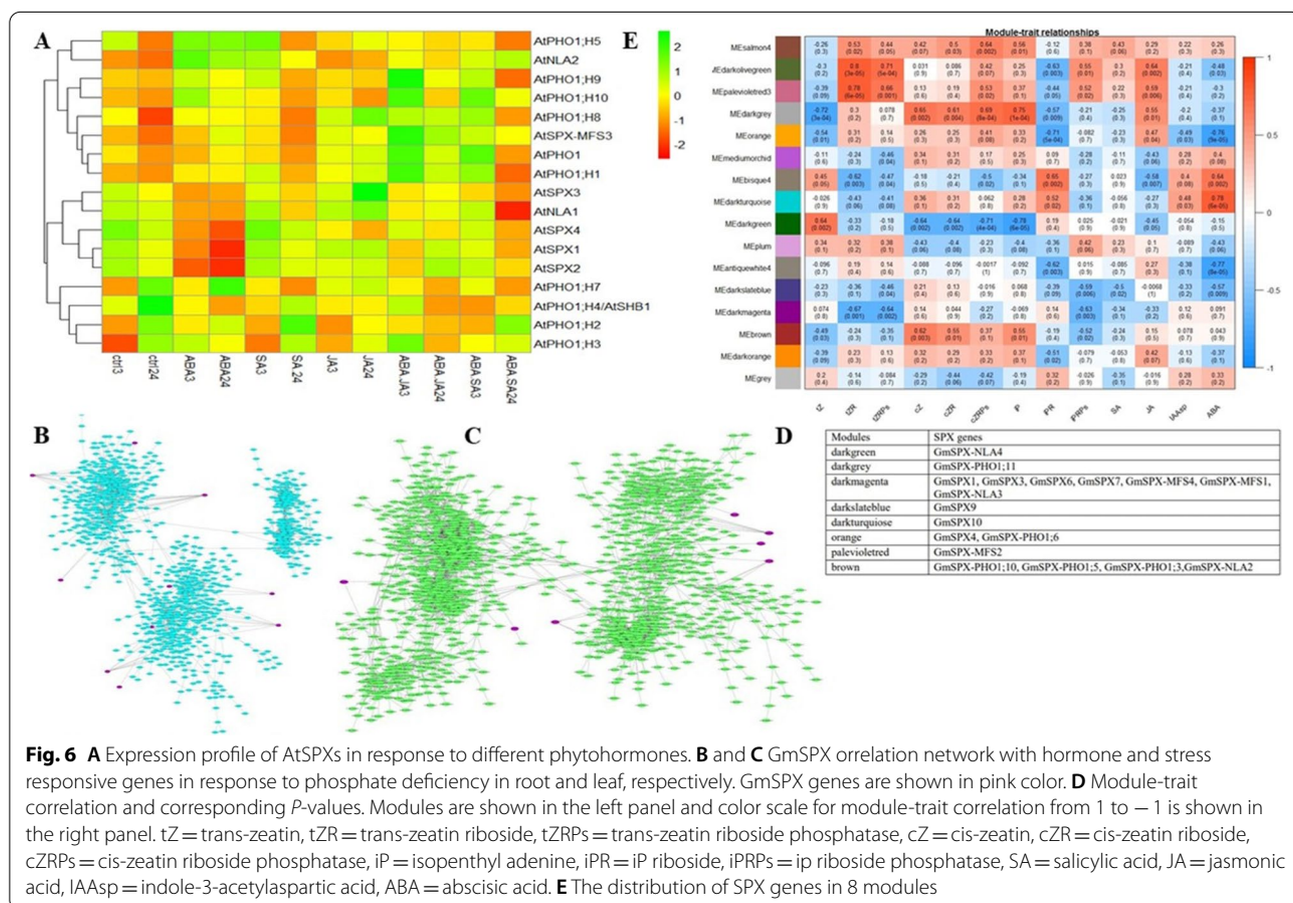


Fig. 6 **A** Expression profile of AtSPXs in response to different phytohormones. **B** and **C** GmSPX orrelation network with hormone and stress responsive genes in response to phosphate deficiency in root and leaf, respectively. GmSPX genes are shown in pink color. **D** Module-trait correlation and corresponding P-values. Modules are shown in the left panel and color scale for module-trait correlation from 1 to -1 is shown in the right panel. tZ = trans-zeatin, tZR = trans-zeatin riboside, tZRPs = trans-zeatin riboside phosphatase, cZ = cis-zeatin, cZR = cis-zeatin riboside, cZRPs = cis-zeatin riboside phosphatase, iP = isopenthyll adenine, iPR = iP riboside, iPRPs = ip riboside phosphatase, SA = salicylic acid, JA = jasmonic acid, IAAasp = indole-3-acetylaspatic acid, ABA = abscisic acid. **E** The distribution of SPX genes in 8 modules

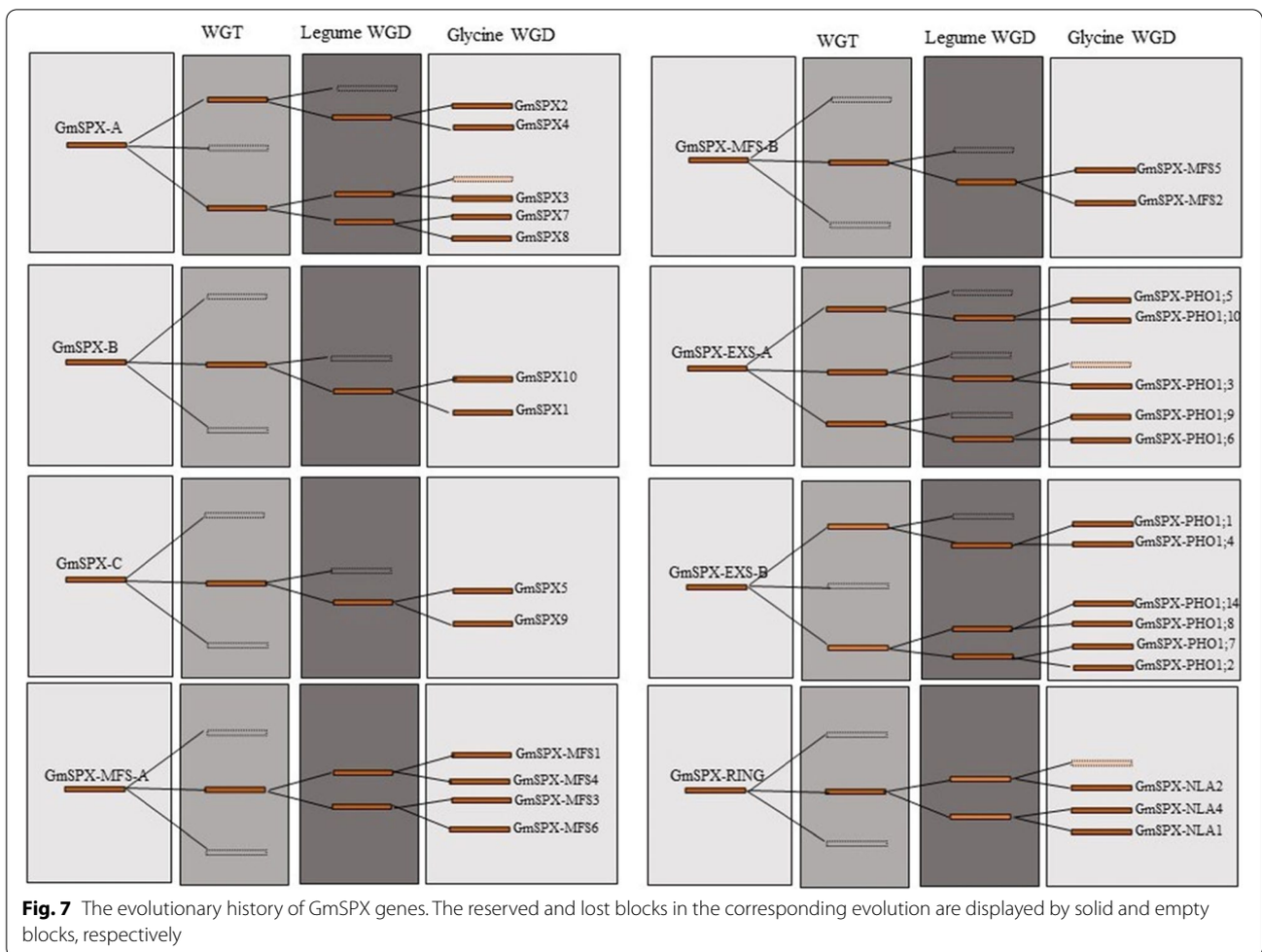
combination with (SA) and (JA) showed less induction. The second dataset analyzed how *GmSPX* genes are correlated with hormonal and stress responsive genes in response to phosphate deficiency. For this purpose, we performed correlation network analysis using TPMs calculated from 40 samples (root and leaf) of a RNA-seq dataset (PRJNA544698). Correlation network was separately constructed for root and leaf using stress and hormonal responsive genes (Fig. 6B and C). This analysis showed that in root and leaf, different SPX genes are correlated with different stress and hormone responsive genes. The third dataset was aimed to determine association of SPX genes with phytohormones in response to dehydration in different soybean organs (leaves, stems, and root). We conducted weighted co-expression network analysis (WGCNA) using microarray and phytohormone data published by Maruyama et al. (2020). WGCNA clustered 12,106 genes into 16 modules and SPXs were distributed in 8 modules (Fig. 6D). Afterwards, the module eigengenes were calculated and then used to create module-trait relationship (Fig. 6E). Among 8 modules that contained SPXs, dark grey, brown, and darkgreen modules showed significant correlation with four phytohormones; trans-zeatin (tZ), cis-zeatin (cZ), cis-zeatin riboside (cZR), cis-zeatin riboside phosphate (cZRPs), and isopentyl adenine (iP). It has been shown that cytokinin can repress PSR genes by modulating nitrate-triggered phosphate signaling [11, 42]. Module dark magenta contained the main players of phosphate starvation responses, including *GmSPX1/3/6/7*, *GmMFS1/4*, and *GmNLA3* that consistently showed negative correlations with trans-zeatin riboside (tZR), trans-zeatin riboside phosphatase (tZRPs), and iP riboside phosphate (iPRPs) forms of cytokinin. Moreover, orange, darkslateblue, and darkturquoise modules, contain *GmSPX4*, *GmPHO1/6*, *GmSPX9*, *GmSPX10* that showed a significant correlation with ABA. Jasmonic acid (JA) showed positive correlations with modules orange, palevioletred, and brown, but salicylic acid (SA) accumulation did not show any correlation with any 16 obtained modules. All in all, results from these analyses showed that while many SPX genes are indeed regulated by phytohormones or part of the same transcriptional networks like hormone regulated genes, as expected from the presence of corresponding cis elements, there is not a clear one-to-one relationship between cis-elements in promoters and the expression of corresponding genes. Previous studies also reported only weak associations between the presence of specific cis-elements and the observed expression profile [43]. Thus, other environmental and developmental factors might modulate

responses of SPX genes to phytohormones and stress [44].

Selective pressure and SPX history model in legumes

The Ks (number of synonymous substitutions per synonymous site) and Ka (number of nonsynonymous substitutions per nonsynonymous site) values of pairs of segmental duplicated SPX genes in soybean, *M. truncatula* and common bean were retrieved from Plant Genome Duplication Database (PGDD) (Supplemental Table S5). The Ka/Ks ratios <1 indicate purifying selection and Ka/Ks values >1 indicate positive selection [45, 46]. The Ka/Ks values for all pairs of segmental duplicated genes were <0.3 implying an intense purifying selection on these gene pairs (Supplemental Table S5). In addition, the Ka/Ks ratio of duplicated gene pairs between soybean and *M. truncatula*, soybean and common bean, and *M. truncatula* and common bean were retrieved (Supplemental Table S5). The mean Ka/Ks values of 0.18, 0.16, and 0.14, respectively, suggest that the genetic pairs between species were subjected to purifying selection.

Based on the Ks values of duplication blocks retrieved from PGDD, the divergence times were estimated. In total, 36, 7, and 3 duplication blocks were retrieved for soybean, *M. truncatula*, and common bean, respectively (Supplemental Table S5). All duplication blocks related to MFS and RING class have Ks<1.5, and the most recent duplication events belonged to MFS members in soybean. Evolutionary process of *GmSPX* genes was modeled based on Ks of duplication blocks (Fig. 7). The duplicated SPX genes in SPX, EXS, MFS, and RING were classified into 3, 2, and 1 groups, respectively. *GmSPX-A* firstly generated three copies after the Gamma WGT event, followed by loss of one copy. The two retained copies were further doubled after Legume WGD event, and after losing one copy, the rest three copies duplicated after Glycine WGD event, resulting in genes, *GmSPX8*, *GmSPX7*, *GmSPX3*, *GmSPX4*, and *GmSPX2*. *GmSPX3* lost its linked duplicated gene (Fig. 7). Unexpectedly, all three generated copies of *GmSPX-EXS-A* in Gamma WGT event were retained but their duplicated genes after Legume WGD were lost. Therefore, Glycine WGD resulted in generation of five genes (*GmSPX-PHO1/10*, *GmSPX-PHO1/5*, *GmSPX-PHO1/3*, *GmSPX-PHO1/9*, and *GmSPX-PHO1/6*) after a loss of one of the linked genes. However, *GmSPX-EXS-B* lost one copy in the first and second round of duplication events and lastly generated six genes (*GmSPX-PHO1/1*, *GmSPX-PHO1/4*, *GmSPX-PHO1/14*, *GmSPX-PHO1/8*, *GmSPX-PHO1/7*, *GmSPX-PHO1/2*). *GmSPX-B* and *-C* as well as *GmSPX-MFS-B* shared the same evolutionary trajectory and generated two duplicated genes in the same way after three rounds



of the evolution processes. In addition, *GmSPX-MFS-A* and *GmSPX-RING* were somewhat similar as both produced two duplicated blocks, although one copy was lost in *GmSPX-RING*, resulting finally in three and four genes, respectively.

Functional characterization of orthologous genes in legumes

Orthologs and orthogroups among seven current angiosperms were determined with OrthoFinder. Altogether, from 218 genes, 216 genes could be classified in seven orthogroups and just two genes of rapeseed (*BnaA6.PHO1;H3c* and *BnaA9.PHO1;H3b*) were not grouped, maybe suggesting a brassica-specific function for these proteins. All members of SPX, SPX-MFS, and SPX-RING were assigned into one group; 1, 3, and 4, respectively. On the other hand, members of EXS family were divided into four distinct groups: group 2 that was dicot-specific; group 7, brassicaceae-specific; as well as groups 5 and 6 that contained genes from all species (Table 2). All genes

in an orthogroup are descended from a single ancestral gene.

Orthologous genes across Arabidopsis and the three legume crops are presented in Table 3. Some genes showed a simple one-to-one orthology relationship, such as *GmSPX6*, *PvSPX2*, and *MtSPX5* with *AtSPX4*; *GmPHO1;3*, *PvPHO1;1*, and *MtPHO1;4* with *AtPHO1;H10*; and *GmNLA3*, *PvNLA1*, and *MtNLA2* with *AtNLA2*. Others showed one-to-many and many-to-many orthology relationships. Interestingly, the pattern of *AtSPXs* orthology relationships were the same among three legumes, and each SPX gene has the same evolutionary trajectories. To overcome the difficulty of one-to-many and many-to-many orthology inference, expressologs of *AtSPXs* with soybean and Medicago were retrieved from the Expression Tree Viewer [32]. Expression Tree Viewer allows to visualize expressologs depending on both sequence similarity and expression pattern similarity. Implementing this web tool resulted in postulating expressologs between Arabidopsis

Table 2 Ortholog groups among soybean, common bean, Medicago, Arabidopsis, rice, wheat, and brassica

Orthogroup	Pv	Gm	Ath	Bna	Os	Ta	Mt	Total	
1	4	10	4	11	5	15	5	54	SPX group
2	4	8	8	29	0	0	4	53	SPX.EXS dicot-specific group
3	3	6	3	8	4	12	3	39	SPX.MFS group
4	3	4	2	7	2	7	4	29	SPX.RING group
5	2	4	1	4	1	9	2	23	SPX.EXS group
6	1	2	1	5	2	3	1	15	SPX.EXS group
7	0	0	1	2	0	0	0	3	SPX.EXS brassicaceae-specific group

Table 3 Ortholog genes between legumes and Arabidopsis

Arabidopsis	Soybean	Common bean	Medicago
AtSPX1/2	GmSPX3/7/8	PvSPX1/5	MtSPX4
AtSPX3	GmSPX1/10	–	MtSPX3
AtSPX4	GmSPX6	PvSPX2	MtSPX5
AtPHO1	GmPHO1;2/7/8/14	PvPHO1;6/5	MtPHO1;1/2
AtPHO1;H1	GmPHO1;1/4	PvPHO1;4	MtPHO1;7
AtPHO1;H2/3/4/5/7/8	GmPHO1;5/10/11/12/13	PvPHO1;2/3	–
AtPHO1;H9	GmPHO1;6/9	PvPHO1;7	MtPHO1;3/5/6
AtPHO1;H10	GmPHO1;3	PvPHO1;1	MtPHO1;4
AtMFS1/2/3	GmMFS1/2/3/4/5/6	PvMFS1/2/3	MtMFS1/2/3
AtNLA/AtBAH1	GmNLA1/2/4	PvNLA2/3	MtNLA1/3/4
AtNLA2	GmNLA3	PvNLA1	MtNLA2

and soybean and Medicago (Supplemental Table S6). Generally, the results were in very good agreement with previous results from phylogenetic tree and OrthoFinder. Based on the Expression Tree Viewer results, we could designate *GmPHO1;2/7* and *MtPHO1;1/2* as the functional orthologs of *AtPHO1* and *AtPHO1;H1* with the function of long-distance Pi transport. However, it was difficult to find expressologs for other SPXs. Consistently, the function of *GmSPX1* [31] and *GmSPX3* [29] were characterized with negative and positive regulatory roles in phosphate deficiency that are the same for *AtSPX1/2* and *AtSPX3* [6].

Expression analysis of SPXs in Arabidopsis and soybean

SPX genes are involved in various physiological process but they are specifically known for their role in phosphate signaling and phosphate homeostasis. To get insight into the potential developmental roles and preferential tissue expression, we analyzed a raw RNA-seq dataset from different developmental stages of different soybean tissues (PRJNA238493). We profiled the *GmSPXs* expression across 17 different samples (Additional file 1: Fig. S13). Overall, we observed different expression patterns of *GmSPXs* in various developmental stages of different

tissues, indicating a functional divergence in each class of *GmSPXs* [27, 47]. For example, *GmMFS2/5* and *GmPHO1;2/7* showed the same expression in almost all samples but were preferentially expressed in leaf and root, respectively. It can be concluded that they are not involved in the developmental processes. On the other hand, duplicated gene pairs arising from Glycine-specific WGD showed very similar expression patterns across all the samples, especially the *GmMFS2/5* gene pair, but except *GmSPX5/9* and *GmPHO1;5/10* pairs. Taking together, both groups of duplicated genes with the same or different expression pattern showed the evidence of sub-functionalization during the soybean evolution [47].

In order to gain insight how individual SPX genes are regulated by Pi deficiency, we analysed publicly available RNAseq dataset (PRJNA544698) [48] and used DPGP software to cluster genes with similar response patterns. DPGP clustering revealed 6 and 4 clusters for root (Additional file 1: Fig. S14) and leaf (Additional file 1: Fig. S15), respectively. We designated names for each cluster based on their patterns; up-reg-fast (cluster 3 in root and cluster 1 in leaf), down-reg-fast (cluster 2 in root and cluster 3 in leaf), the lowest-peak-T1 (cluster 6 in root), the lowest-peak-T2 (cluster 5 in root), the

highest-peak-T1 (cluster 1 in root), up-reg-slow (cluster 4 in leaf), and the highest-peak-T2 (cluster 4 in root and cluster 2 in leaf). As can be seen in the Table 4, some genes have opposite pattern of regulation in different tissues. To exemplify, *GmSPX1* was placed in down-reg-fast in root and up-reg-fast in leaf, *GmSPX-PHO1;10* is found in the highest-peak-T1 in root and the highest-peak-T2 in the leaf, while *GmSPX6*, *GmSPX-NLA1*, and *GmSPX-NLA3* were in the lowest-peak-T2 cluster in root and the highest-peak-T2 in leaf. The homologs of *AtPHO1* and *AtPHO1;H1(PHO1;2/7/14)* showed an up-reg-fast pattern of cluster 4 in root and the highest-peak-T2 in clusters 2 leaf. Supporting these patterns, He et al. (2013) reported similar expression pattern for these genes, however, there is no clear association between increasing mRNA level of these genes in leaves during phosphate deficiency and growth or shoot Pi content [15]. Overall, for the genes which show tissue-specific expression, we observed different patterns in root and shoot in response to phosphate deficiency.

Finally, after investigating developmental and dynamical expression patterns of *GmSPX*, we used another RNA-seq dataset from Arabidopsis and soybean to examine the expression of *SPXs* in three different zones of root [49]. The original data were generated in multiple species, however, we only used RPKM values from Arabidopsis and soybean. A general comparison showed that almost all *SPX* tended to group species-based rather than orthology-based, except *AtPHO1* which clustered with their orthologs, *GmPHO1;2* and *GmPHO1;7* (Additional file 1: Fig. S16). Thus, we can conclude that the tissue-specific genes pose difficulty to identify functional orthologs because of probable tissue inequivalences among species.

GmSPX gene expression in response to nitrogen, phosphorus, and combined deficiency

Cross-talk between P and N signaling has been reported in previous studies [11, 50–53], and some molecular mechanisms involved in N and P cross-talk have been identified, such as the nitrate-NRT1.1B-SPX4 cascade in rice [11] and NIGT1-SPX-PHR cascade in Arabidopsis [53]. Clearly, *SPX* gene family seems to have a vital role in this cross-talk. To assess whether *GmSPX* genes are regulated during N and P deficiency and test the validity of our in silico analyses to identify functional orthologs in soybean, qRT-PCR was performed for several *GmSPX* genes. *GmSPXs* showed different regulation in leaf and root in response to N, P, and NP deficiencies (Fig. 8). Except *GmSPX6*, transcript levels of other members of *SPX* class significantly increased in response to P and, vice versa, decreased in N deficiency. This antagonistic behavior was reported previously in maize [50, 54]. The transcript levels of *GmSPX6* were not much affected by the treatment which is consistent with post-translational regulation of it rice orthologue *OsSPX4* [11]. *GmPHO1;2* and *1;7* are involved in long-distance P transportation, however, whereas *GmPHO1;2* was upregulated in both N and P deficiency in root but not in leaf, the *GmPHO1;7* was not significantly affected. *GmNLA3* was only induced by the combined deficiency (NP) and *GmMFS1* did not show significantly different expression in root but was induced in N and NP deficiency in leaf. Thus, the expression pattern of many of the tested *GmSPX* genes is consistent with their involvement in P and/or N homeostasis.

Discussion

The role of *SPX* domain-containing proteins in Pi homeostasis in Arabidopsis, rice, rapeseed, and wheat and to some extent in soybean and common bean were studied previously [3, 13, 27, 29–31, 35]. While an evolutionary

Table 4 Different patterns of clusters in root and leaf in the time series dataset of soybean

Patterns	Root clusters	Leaf clusters
up-reg-fast	Cluster3: <i>SPX4</i> , <i>SPX.PHO1;2</i> , <i>SPX.PHO1;7</i> , <i>SPX.PHO1;8</i> , <i>SPX.PHO1;14</i> , <i>SPX.PHO1;6</i>	Cluster1: <i>SPX1</i> , <i>SPX.MFS6</i> , <i>MFS-NLA2</i> , <i>MFS-NLA4</i> , <i>SPX.PHO1;11</i>
down-reg-fast	Cluster2: <i>SPX1</i> , <i>SPX10</i> , <i>SPX5</i> , <i>SPX.MFS4</i> , <i>SPX.NLA2</i> , <i>SPX.NLA4</i> , <i>SPX.PHO1;12</i>	Cluster3: <i>SPX2</i> , <i>SPX.MFS2</i> , <i>SPX.MFS5</i> , <i>SPX.MFS4</i>
up-reg-slow		Cluster4: <i>SPX4</i> , <i>SPX5</i> , <i>SPX.PHO1;1</i> , <i>SPX.PHO1;4</i> , <i>SPX.PHO1;6</i> , <i>SPX.PHO1;9</i>
highest-peak-T1	Cluster1: <i>SPX.MFS1</i> , <i>SPX.PHO1;5</i> , <i>SPX.PHO1;10</i> , <i>SPX.PHO1;9</i> , <i>SPX.PHO1;1</i>	
highest-peak-T2	Cluster4: <i>SPX.MFS5</i> , <i>SPX.PHO1;4</i> , <i>SPX.PHO1;11</i>	Cluster2: <i>SPX3</i> , <i>SPX6</i> , <i>SPX7</i> , <i>SPX8</i> , <i>SPX10</i> , <i>SPX.NLA1</i> , <i>SPX.NLA3</i> , <i>SPX.PHO1;2</i> , <i>SPX.PHO1;7</i> , <i>SPX.PHO1;14</i> , <i>SPX.PHO1;5</i> , <i>SPX.PHO1;10</i> , <i>SPX.PHO1;12</i>
lowest-peak-T1	Cluster6: <i>SPX3</i> , <i>SPX7</i> , <i>SPX8</i> , <i>SPX.MFS2</i> , <i>SPX.MFS6</i>	
lowest-peak-T2	Cluster5: <i>SPX6</i> , <i>SPX.NLA1</i> , <i>SPX.NLA3</i>	

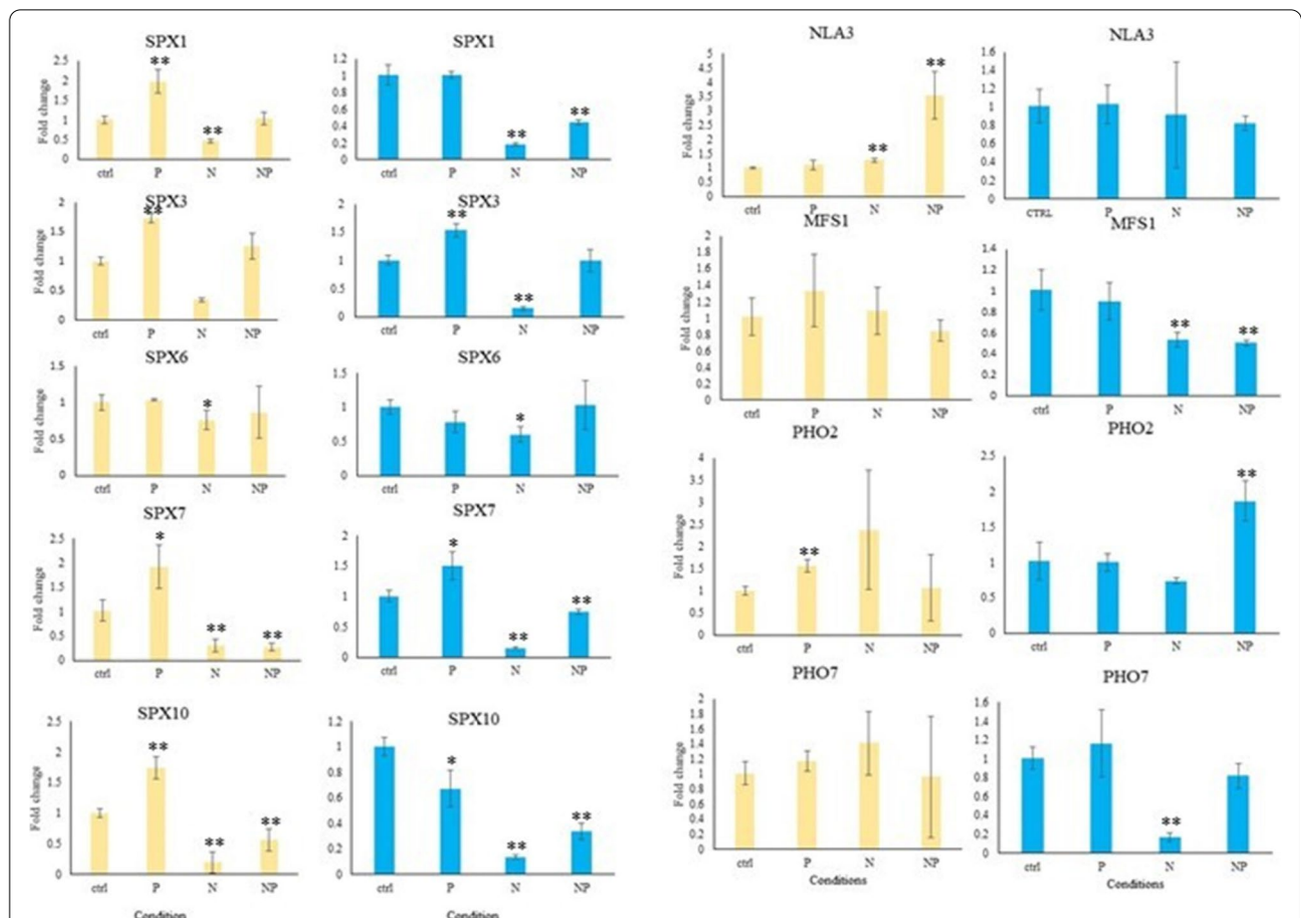


Fig. 8 Relative expression of GmSPX genes in response to nitrogen (N), phosphorus (P), and the combined deficiency (NP). Expression levels were determined by qRT-PCR using 3 biological and 2 technical replicates. The data are shown as means ± SD, the values in controls were set to 1. Yellow and blue bars are representing of root and leaf, respectively. Asterisks mark values different from controls ($P < 0.05$; T-test)

analysis of SPX-EXS [55] and SPX-MFS [24] classes has been reported, as far as we know, the evolution of all classes of SPX gene family from algae to higher plants has not been explored. In addition, despite legume crops requiring a relatively high amount of P, no systematic study of SPX gene family has been reported in legume crops. To close this knowledge gap, we performed a comprehensive search for SPX genes throughout three legume crops, including soybean, *M. truncatula*, and common bean and also algae, liverwort, hornwort, and basal angiosperms to figure out how this gene family originated and expanded during the evolution as well as to identify SPX functional orthologs in legumes.

Evolutionary conservation and divergence of SPX gene family from algae to legumes

Proteins harboring SPX domain has been reported to form four classes based on their domains. Meanwhile, some other classes have been revealed in the basal plants

and algae such as SPX-SLC and SPX-VTC [24]. Here we report other functional protein domains being fused to SPX domains, including EIN3, S6PP, EIN3-S6PP, and Kelch in *S. moellendorffii*, CitMHS in *C. crispus*, Na_sulph_symp in *G. sulphuraria* and *C. reinhardtii*, BET (Bromodomain extra-terminal-transcription regulation) in *P. somniferum*, EXS.rve in *C. braunii*, and Sugar_tr in *M. polymorpha*. Interestingly, some of these new domains have been lost in the land plants and all of them in angiosperms. Domains present in algae before land colonization probably had specific functions that are not required for land plants. For example, SPX-SLC and SPX-VTC were reported in algae that store polyP and are thus lost in plants with Pi vacuole storage, which in turn gained SPX-MFS [24]. Among all assayed species, *S. moellendorffii* showed the most variation of SPX genes, which could be due to its special ability of resurrection. Moreover, unlike other SPX proteins, SPX domain are located at C terminal in S6PP-SPX, EIN3-S6PP_C-SPX,

EIN3-SPX, and Kelch-SPX classes. The function of other fusion proteins is unknown so far. Particularly interesting are the fusions of SPX with EIN3 domains, because in *Arabidopsis* EIN3 is directly involved in regulation of phosphate homeostasis through binding to promoter of *PHR1* [56]. The SPX domain would then add another level of control for this interaction and allow the reciprocal regulation of ethylene signaling by phosphate. Similarly, Kelch domains are often found in regulatory proteins, for example fused to F-Box proteins [57], hence, again, the fusion with SPX may connect multiple regulatory circuits. If the SPX domain enables the activities of the additional domains to be modulated by phosphate (or InsPP), this offers an intriguing opportunity for using these domains in synthetic biology approaches to make various cellular processes controlled by phosphate. These hypotheses, however, have to be verified. On the other hand, RING and MFS classes have gradually appeared in the later-diverging plants. MFS and then RING class have the least fluctuations from 1 to 6 genes. In contrast, EXS class had high variation of gene numbers in each species and also the highest number of identified genes in comparison with the other classes. Also, presence of this domain in whole Eukarya except algae, suggest that it has been lost in some algae.

The number of whole-genome duplications is correlated with gene family size [55, 58], which is consistent with our results, since *P. somniferum* and *G. max* with two WGD events had the largest sizes of SPX family [59, 60]. The expansion of SPX family in these two plants is mostly affected by WGD duplication type, while segmental/local duplication type was the main contributor of expansion in *S. moellendorffii*, the species with third greatest SPX family, which might explain its unique classes. Algae possess 2 to 5 SPX gene family members. The expansion in *P. patens* (22 members), could suggest that duplications took place after plant terrestrialization as the SPX proteins became more important [61].

The phylogenetic analysis brought some unexpected findings. First, it showed three clades for 4 subfamilies; SPX and EXS in two different clades, but MFS and RING classes diverged from the same ancestor. Second, SPXs from algae did not group with other species in any clade, except of SPX-I. It can be concluded that genes in the SPX-I sub-clade are the most ancient genes in angiosperms that were diverged from the same ancestor with green algae. Hence, *AtSPX4*, *GmSPX6*, *MtSPX5*, *PvSPX2*, and *OsSPX4* probably have the same function with their ancestral orthologs in the green algae, but the genes in two other sub-clades, SPX-II and SPX-III have evolved after divergence of streptophytes and chlorophytes and might have

acquired additional functions. *AtSPX4* and *OsSPX4* have indeed the same function and mechanism in regulation of PSI, as in presence of phosphate both proteins interact in the cytosol with the corresponding key regulators *AtPHR1* and *OsPHR2*, and prevent them from translocating to nucleus [11, 62]. During P deficiency they are rapidly degraded, releasing thus the PHR factors to induce transcription of PSI genes. The two proteins however, also differ, as while *OsSPX4* integrates nitrate and phosphate signaling, *AtSPX4* does not seem to have this function, but on the other hand integrates phosphate signaling and anthocyanin biosynthesis [11, 63].

The MFS class as the most recently diverged class of SPX proteins was divided into two sub-clades, with MFS-I specifically containing monocots, suggesting that MFS genes in monocots diversified differently in comparison with basal angiosperms and eudicots. This may be due to different Pi storage between monocots and eudicots [6, 64, 65]. While monocots store P preferentially in the roots and their leaves have the highest P concentration in the mesophyll cells, eudicots store much more P in the leaves with the highest concentration in the epidermis [64]. It is thus possible that the different cellular localization drove a different evolution of SPX-MFS genes between monocots and dicots. Modern RING class genes have evolved two times, RING-I clade arose from a duplication of the common ancestor of mosses and angiosperms and RING-II arose from duplication of the common ancestor of lycophytes, liverwort and angiosperms. In the EXS class, EXS-III clade did not contain any orthologs from monocots but interestingly, many *AtPHO1* genes such as *AtPHO1;H2/3/4/5/6/7/8* grouped specifically with the genes from *Brassica napus*, suggesting special functions in Brassicaceae. Only *AtPHO1;H9* and *AtPHO1;H10* had two and one orthologs in the legumes, respectively.

Based on collinearity analyses, species with more WGD events showed more inter-species collinearity, but *S. moellendorffii* with locally expanded SPX and rice with mostly dispersed expanded SPX just showed intra-genome collinearity. Low collinear relationship between rice and eudicots was reported previously [66] and explained by longer evolutionary distance and more genome rearrangements [67] as well as the erosion of macrosynteny between monocots and dicots [68]. Our results are consistent with the monocot paleopolyploidy after their divergence from eudicots [66]. Having collinear relationship can arise from paleopolyploidy in the common ancestor, but *S. moellendorffii* has no evidence for WGD events and its intra-genome collinear blocks arose from segmental/local duplication [37].

Functional characterization of SPXs in legumes

Due to the functional conservation of proteins across species, determination of orthologous relationships can provide useful insights about the biological role of these proteins [69]. As plants have undergone various duplication events and had different evolutionary trajectories, relating same functions to the orthologs are difficult, especially there are one-to-many or many-to-many orthologous relationships [32]. Therefore, two different methods, phylogenetic inference of orthologs from protein sequences and expressolog identification, were conducted for prediction of functional orthologs of SPXs. This was necessary because, firstly, there are complex orthology relationships among some SPX genes that prevented Orthofinder to detect the exact functional orthologs and, secondly, some SPX genes show tissue-expression pattern that can pose problem to identify expressologs, due to difficulties in assignment of tissue equivalencies between legumes and Arabidopsis. In the dynamic *GmSPX* expression patterns, we observed tissue-specificity for most of *GmSPXs* except for homologs of *AtPHO1* and *AtPHO1;H1*. Taking together, we could assign functions of *AtSPX4*, *AtPHO1;H10* and *AtNLA2* to their predicted orthologs from Orthofinder and *AtPHO1* to their orthologs from expressolog identification results. To examine this conclusion, we analyzed two different datasets of soybean to profile *GmSPXs* expression in different tissues and developmental stages as well as their dynamic expression responses to Pi deficiency in leaf and root. Overall, we found that almost all *GmSPXs* except *GmPHO1;2/7* and *GmMFS2* have different expression patterns across the developmental samples as well as in root and leaf responses to the dynamic Pi deficiency. Also a qRT-PCR analysis of the effects of P, N, and combined P and N deficiency on *GmSPX* genes showed the induction of most by P deficiency, corroborating their role in P homeostasis, but also repression in N deficiency. In summary, these transcriptome analyses highlighted that *GmSPX* genes might be involved in different developmental processes and stresses beyond phosphate starvation response. It is probable that new or sub-functionalization in soybean and generally in legumes took place with the new functions of SPX proteins waiting to be discovered. Our analyses lay a solid foundation for the future functional studies of SPX proteins from algae to legumes.

Conclusion

In conclusion, we comprehensively analyzed SPX gene family evolution and dissected how different protein motifs and Cis-acting elements evolved, as well as identified expansion patterns, and collinear gene blocks during evolution from algae to angiosperms. Afterwards,

focusing on legumes, we tried to model evolutionary history of SPXs in soybean and identify functional orthologs. We could predict the putative SPX proteins involved in long-distance Pi transportation in soybean and Medicago. Our study not only provides a global view of the evolution and expansion of *SPX* gene family in important species but also provides the first step for more detailed investigations of the functions of individual *SPXs* in legumes.

Material and methods

Bioinformatic identification of SPX proteins

In order to identify SPX domain-containing proteins in our species; legume crops (soybean – *Glycine max*, alfalfa – *Medicago truncatula*, and common bean – *Phaseolus vulgaris*), mosses (*Physcomitrella patens*), liverwort (*Marchantia polymorpha*), Rhodophytes (*Cyanidioschyzon merolae*, *Galdieria sulphuraria*, and *Chondrus crispus*), chlorophytes (*Chlamydomonas reinhardtii* and *Ostreococcus lucimarinus*), charophytes (*Chara braunii*), basal angiosperms (*Papaver somniferum*, *Amborella trichopoda*, and *Nymphaea colorata*), and lycophytes (*Selaginella moellendorffii*), full-length protein sequences of *AtSPXs* were used for BLASTP searches across proteomes of the above mentioned species. After removing redundant sequences, the SPX proteins obtained through BLASTP search were investigated for the presence of additional domains along with SPX domain using SMART [70], Pfam [71], Conserved Domain Database (CDD) [72], and PROSITE [73] databases.

The sequences of identified SPX proteins in the three legume crops were analyzed for their physiochemical properties; including isoelectric point (pI), molecular weight (Mw), instability index (II), grand average of hydropathicity (GRAVY), and aliphatic index (AI) using ProtParam tool of ExPASy website (<https://web.expasy.org/protparam/>). Subcellular location prediction was conducted using Wolf Psort [74].

Phylogeny analysis and identification of conserved motifs

The amino acid sequences of identified SPX proteins in our surveyed species and Arabidopsis as reviewed in [6], rice [6], wheat [3], and *Brassica napus* [27] were downloaded from EnsemblPlants (<https://plants.ensembl.org/index.html>). Three sequences to be used as outgroup, XPR1 from human and mouse, and SYG1 from *C. elegans*, were downloaded from NCBI database (<https://www.ncbi.nlm.nih.gov/>). Multiple sequence alignment of these full-length sequences was performed by ClustalX (ver. 2.1; <http://www.clustal.org/>). Then, we used Maximum Likelihood method and JTT matrix-based model in MEGA 7 software to build a phylogenetic tree from the sequence alignment using following parameters:

p-distance model, partial deletion and 1000 bootstraps. To predict conserved motifs of SPX proteins across all species, as well as Arabidopsis and rice, MEME (<http://meme-suite.org/tools/meme>) tool with the maximum number of motifs 20 was used. Logo sequences of conserved motifs were obtained by Weblogo 3 (<http://weblogo.threeplusone.com/>).

Collinearity analysis and gene expansion pattern of SPX from algae to eudicots

In order to get insight about how collinear blocks have been conserved during the evolution, we performed collinearity analysis three times with different species; 1. Among three legume crops, Arabidopsis, rice, *P. somniferum*, and *N. colorata*, 2. Among *S. moellendorffii*, *P. patens*, *N. colorata*, and *A. trichopoda*, and 3. Among three legume crops using MCScanX toolkit [75] to get collinear gene blocks and also duplication types by duplicate_gene_classifier program. To visualize the collinear blocks among the first and third runs, ttools was used [76]. Because of non-chromosomal reference genomes in *P. patens* and *S. moellendorffii* we just retrieved their collinear gene blocks without visualization.

Selective pressure and evolutionary models of SPX genes in the legume crops

Duplication blocks between each two species of soybean, common bean and *M. truncatula* were retrieved from the Plant Genome Duplication Database (PGDD, <http://chibba.agtec.uga.edu/duplication/>). SPX gene blocks were manually extracted and used for further analyses. The selective pressure on duplicated genes were estimated by retrieving synonymous (Ks) and non-synonymous (Ka) per site between the duplicated gene-pairs using from PGDD database. The Ka/Ks ratio was assessed to determine the molecular evolutionary rates of each gene pair. Generally, the $Ka/Ks < 1$ indicates purifying selection, $Ka/Ks > 1$ indicates positive selection, and $Ka/Ks = 1$ indicates neutral selection. The divergence time of the duplication blocks was evaluated to investigate the evolution of GmSPX genes. If the $Ks > 1.5$, the divergence time is after the Gamma whole-genome triplification (WGT); if the $Ks < 0.3$, the divergence time is after the Glycine whole-genome duplication (WGD) event; and when the Ks is between 0.3 and 1.5, the divergence time is after legume WGD event but before the Glycine WGD event [77, 78].

Identification of Cis-acting-elements in the promoters of SPX gene family

For finding evolutionary pattern of Cis-acting-elements from algae to eudicots, 1500bp upstream from the start codon of SPX genes in all assayed species and

Arabidopsis were downloaded from the EnsemblPlants and analyzed using the PlantCARE database (<http://bioinformatics.psb.ugent.be/webtools/plantcare/html/>). PlantPan3 database (<http://PlantPAN.itsps.ncku.edu.tw>) was used for finding PIBS motif in soybean and *C. reinhardtii* genes. Afterwards, SPX genes were clustered with hierarchical clustering on principal components (HCPC) method by FactMineR package. All detected cis-acting elements were merged into one matrix with 1 and 0 values for present or absent elements in each promoter, respectively.

Prediction of functional orthologs of AtSPXs across legumes

To identify functional orthologs in the three legumes, we used OrthFinder to compare SPX genes among 7 species (rice, wheat, rapeseed, Arabidopsis, *M. truncatula*, soybean, and common bean), resulting in orthogroups and orthologs based on sequence similarities [79]. Then, to overcome the weakness of sequence-based ortholog identification for one-to-many and many-to-many orthologs, expressolog identification among Arabidopsis, soybean, and Medicago (http://bar.utoronto.ca/expressolog_treeviewer/cgi-bin/expressolog_treeviewer.cgi), was used.

Expression analysis of SPX genes

Three different expression analyses were performed as follows:

1. To compare tissue and developmental expression pattern of GmSPXs, RNA-seq data of 17 samples from different tissues of soybean (*Glycine max* cv Williams82) (flower, root, shoot meristem, seed, and leaves) in five developmental stages (germination, trefoil, flowering, seed development, and plant senescence) (PRJNA238493) [80] were analyzed. The gene expression profiles were visualized by heatmap using R package pheatmap (<https://www.r-project.org/>).
2. To visualize changes in SPX gene expression in soybean (*Glycine max* cv Clark) in response to P deficiency we used publicly available dataset (PRJNA544698) [48]. The data were reanalyzed and TPM (Transcript Per Million) values were calculated from samples over different time points of Pi deficiency, including early stress (T, 24h), recovery (TC, 24h deficiency, 48h resupply), and repeated stress (TCT, additional 24h deficiency) in root and leaf tissues. The data were clustered using the Dirichlet process with Gaussian process mixture model (DPGP) [81].
3. To assess if the predicted functional orthologs in Arabidopsis and soybean (*Glycine max* cv Williams82) show the same expression in different root

development zones, including meristemic zone (MZ), elongation zone (EZ), and differentiation zone (DZ) data from [49] have been used. RPKM values for the SPXs were collected (GSE64665), and \log_2 (RPKM + 1) was used to construct correlation heatmap using the pheatmap package (<https://www.r-project.org/>).

Co-expression module analysis of GmSPX genes

WGCNA [82] was used to investigate co-expression modules related to phytohormones. We used microarray and phytohormone datasets which were measured in different organs of soybean (*Glycine max* cv Williams82) in response to dehydration (accession number E-MTAB-7010) [83]. Firstly, the low-quality data were removed, then the modules were detected. The power value was 0.8, with minimum 30 genes in a module, and other parameters set to default parameters. Afterwards, eigengene of module were calculated and used for construction module-trait relationship.

Soybean growth condition

Soybean seeds (Williams82) were planted in pots (containing river sand, 1-l per pot) in a completely randomized design with three biological replicates. Based on the experimental plan, pots were watered with 4 different solutions (Additional file 2, Table 1) which were nitrogen normal-phosphorus normal (N1P1), nitrogen normal-phosphorus deficiency (N1P0), nitrogen deficiency-phosphorus normal (N0P1), and nitrogen deficiency-phosphorus deficiency (N0P0) [84, 85]. To inhibit any nodulation, all solutions were prepared with distilled water, and also the sand was autoclaved. The pots were watered with the nutrient solution once every 2 days for 28 days after planting. For sampling, roots and leaves were harvested, carefully cleaned and frozen in liquid N₂ and stored at -80 °C for qRT-PCR experiment.

RNA extraction and qRT-PCR analysis

Total RNA was extracted by standard phenol/chloroform extraction and LiCl precipitation as in Koprivova et al. (2019) [86]. DNase treatment to remove of contamination of genomic DNA and first-strand cDNA synthesis was performed using QuantiTect Reverse Transcription Kit (Qiagen). Quantitative real-time RT-PCR (qPCR) was performed for nine *GmSPXs* using gene-specific primers (Additional file 2, Table 2) as in Koprivova et al. (2019) [86]. The Actin-6 (*GLYMA_18G290800*) gene was used as the internal reference gene for qRT-PCR.

Abbreviations

PHR: Phosphate starvation Response; PP-InsPs: Inositol pyrophosphates; MFS: Major Facilitator Superfamily; RING: Really Interesting New Gene; NLA: Nitrogen Limitation Adaptation; PSI: Pi starvation-induced; Ein3: Ethylene intensive 3; VTC: Vacuolar transporter chaperone; CitMHS: Citrate transporter; Na_sulph_symp: Sodium sulphate symporter; S6PP_C: Sucrose-6F-phosphate phosphohydrolase C-terminal; Kelch: Galactose oxidase; rve: Integrase core domain; GRAVY: Grand average of hydropathicity; LSC: Lysine Surface Cluster; Gm: *Glycine max*; Mt: *Medicago truncatula*; Pv: *Phaseolus vulgaris*; Ps: *Papaver sumniferum*; Nc: *Nymphaea colorata*; At: *Arabidopsis thaliana*; Os: *Oryza sativa*; CRE: Cis-acting elements; MeJA: Methyl jasmonate; tZ: Trans-zeatin; tZR: Trans-zeatin riboside; tZRPs: Trans-zeatin riboside phosphatase; cZ: Cis-zeatin; cZR: Cis-zeatin riboside; cZRPs: Cis-zeatin riboside phosphatase; iP: Isopentenyl adenine; iPR: iP riboside; iPRPs: iP riboside phosphatase; SA: Salicylic acid; JA: Jasmonic acid; IAAsp: Indole-3-acetylserine; DPGP: Dirichlet process with Gaussian process mixture model; HCPC: Hierarchical clustering on principal components.

Supplementary Information

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Additional file 1.

Additional file 2.

Additional file 3.

Additional file 4.

Additional file 5.

Additional file 6.

Additional file 7.

Additional file 8.

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Authors' contributions

MNC, AN, EE, and SK designed the study. MNC performed the analyses. SK assisted with interpretation. MNC wrote the manuscript. All authors reviewed the manuscript. The authors read and approved the final manuscript.

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Availability of data and materials

The datasets generated and/or analyzed during the current study are included in the supplemental material.

Declarations

Ethics approval and consent to participate

Not applicable.

Consent for publication

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Competing interests

The authors declare no competing interests.

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