The prohibitins (PHB) gene family in tomato: Bioinformatic identification and expression analysis under abiotic and phytohormone stresses

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1. Introduction

The prohibitins (PHB) genes concede highly conserved stomatin/prohibitin/flotillin/HflK/C (SPFH) domain in their protein sequence also recognized as band_7 domain proteins. PHBs proteins are ubiquitous proteins and are associated with a variety of biological processes including cell cycle, apoptosis, and respiration. PHBs have been identified from eukaryotes, fungi, plants, and animals. In humans, the PHB proteins act as transcriptional regulators interacting with PSF3, retinoblastoma proteins (Rb), and E2F. PHB genes were observed to be linked with the breast cancer phenotype, where they localize in the nucleus of breast cancer cell lines as a transcriptional regulator via interaction with P53, RB and E2F to regulate the expression of downstream genes. PHBs were also identified in lipid raft, a key constituent of cell membrane. Similarly, PHBs found in plasma membrane were considered to act as a target for small molecules in the inflammatory responses as well as to regulate the membrane receptor and iron channels. In short, PHB genes play crucial roles in different biological processes and are associated with various disease phenotypes. However, less is known about the role of PHB proteins in the plant kingdom.

PHB proteins are classified into type I and type II and both are complimentary for stability and...
functioning of PHB protein. In mammals, PHB1 and PHB2 have been well characterized and shown to form a 1–2 KDa protein complex on the inner mitochondrial membrane. In addition, the absence of any of these two proteins failed to produce this protein complex in Caenorhabditis elegans, resulted in decreased PHB proteins. PHB complex have been physically and functionally linked with the matrix-ATPase related to diverse cellular activities (m-AA) to regulate the degradation of respiratory chain proteins in mitochondria. PHB and PHB2/REA were found to be involved in maintaining cellular survival via Ras–Raf–MEK–Erk pathway. These findings suggest that both types of PHB are required for stable complex formation and proper functioning. Recently, various studies reported the role of PHB in plants. These proteins play a pivotal role not only in plant development and senescence but also in responses to abiotic and biotic stresses. PHB3 and PHB4 are the most broadly studied PHB genes from Arabidopsis thaliana, where they primarily expressed both in root and shoot proliferative tissues. Arabidopsis mutant, atphb3 exhibited severely retarded growth phenotypes, decreased stem, root proliferation, and declined cell division in root and stem apices. Overexpression of Arabidopsis PHB (AtPHB3/AtPHB4) exhibited irregular leaf shape and extensive branching phenotype. Notably, atphb3/4 double knockout mutants were not viable, suggesting that PHBs play important role in plant development. Similar results were obtained in petunia and tobacco, where PHB-silenced genes showed decreased cell production and prolonged senescence. In tobacco, suppression of NbPHB2 delays growth and promotes leaf senescence and apoposis. Moreover, the cells in silenced flowers were larger as compared to control flowers, suggesting a significant decrease in the number of cell division that occurs during corolla development. PHB proteins directly or indirectly interact with mitochondrial DNA (mtDNA) to regulate the reactive oxygen species (ROS) formation and oxidative phosphorylation (OXPHOS), which potentially lead to senescence phenotype both in C. elegans and plants. Furthermore, PHB protein might also involve in maintaining crista morphology to employ proteins into the inner membrane. The abovementioned finding indicates that PHB play key functioning in cell proliferation. Several studies have shown that PHB proteins play key roles not only in plant development and senescence but also in response to salinity, defense and plant hormones. For instance, Arabidopsis eer-3-1(atphb3) mutant showed an etiolated seedling phenotype upon constitutive exposure to ethylene with suppressed the expression of various ethylene inducible genes (Arabidopsis ethylene-responsive element binding protein [AtEBP]), plant defensin [PDF 1.2]), indicating the dual role of AtPHB3 in Arabidopsis. Additionally, AtPHB3 acted downstream of ethylene insensitive 2 (EIN2) and EIN3. A loss of function mutant atphb3-3 failed to affect diverse biological processes such as nitric oxide (NO) signaling, ABA (abscisic acid) induced stomatal closure, IAA (auxin) induced root formation. This mutant resulted from the substitution of Gly at position 165 with Asp of AtPHB3 protein. However, another Arabidopsis PHB (At5g64870) induced under cold, salinity, and drought but suppressed in response to hormones such as gibberellin (GA), methyl jasmonate (MeJA), and ABA. PHB proteins have been identified in various plant species including 17 in Arabidopsis, 19 in rice, 24 in Glycine max, and Zea mays with 16. The knowledge about PHB genes in tomato is insufficient. In this study, a total of 16 PHB genes were identified in the tomato genome. Phylogenetic analysis, gene structure, in silico subcellular location prediction, cis-regulatory elements, MEME motif scan, and protein chromosome location were also conducted. In addition, tissues/organ-specific expression profiling under normal conditions was evaluated. Moreover, differential expression patterns under salt, drought, and hormone-induced expression were analyzed. This study enables us to provide a foundation for the functional characterization of PHB genes in tomato.

2. Material and Method

2.1. The Tomato PHB Gene Discovery

To predict PHB genes in the tomato genome, the Arabidopsis, rice, Zea mays, and Glycine max PHB peptide sequences were retrieved from the TAIR genome database (https://www.arabidopsis.org/), rice genome annotation project (http://rice.plantbiology.msu.edu), phytozome database (https://
phytozone.jgi.doe.gov/), respectively. These sequences were used as a query in the SOL genome network (https://solgenomics.net). The candidates’ sequences were analyzed for SPFH Domain (PF01145) in the SMART (http://smart.embl-heidelberg.de) and NCBI conserved domain database (CDD, https://ncbi.nlm.nih.gov/Structure/bwrpsb/bwrpsb.cgi). Moreover, PHB protein features including the isoelectric point (pI), the grand average of hydropathy (GRAVY), molecular weight (kDa) of each protein were calculated in sequence manipulation suite (SMS, bioinformatics.org/sms2). The deduced PHB proteins were named in their order on the tomato chromosomes.

### 2.2. Phylogenetic Analysis and Ka/Ks Analysis of Duplications

Clustal Omega (ClustaloO, https://www.ebi.ac.uk/Tools/msa/clustalo/) program was used to generate SIPHB peptide sequences alignment. For the phylogenetic relationship, SIPHBs peptide sequences from rice, Arabidopsis, Zea mays, and soybean were retrieved from phytozone (https://phytozone.jgi.doe.gov). An unrooted neighbor-joining tree was constructed using MEGAX software with the parameters set as follows: Poisson correlation of model; pairwise deletion of gaps/missing data; random seed of phylogeny test and bootstrap was set at 1000 replicates. The non-synonymous (Ka), synonymous (Ks) nucleotide substitution rates and the Ka/Ks ratio were predicted using k-estimator (http://en.bio-soft.net/format/KES estimator.html). The divergence time (T) was calculated as follows: T = Ks/2y (y = 6.56 x 10^-9).

### 2.3. Chromosome Location, Subcellular Location Prediction, and miRNA Target Prediction

The chromosome position of each SIPHB gene was obtained from the SOL genome and visualized in the MAPGene2Chromosome program (http://mg2c.iask.in/mg2c_v2.0/). In silico subcellular location, prediction analysis was performed in the WoLFPSORT program (https://wolfsort.hgc.jp). To predict miRNAs targeted putative PHBs, the cDNA sequences of each SIPHBs were submitted to psRNATarget against all tomato miRNAs reported in miRBase.

### 2.4. Gene Structure Analysis, Conserved Motif Scan, and cis-Regulatory Motif Prediction

The retrieved tomato SLPHBs coding sequences (CDS) and genomic sequences were submitted to the Gene Structure Display Server (GSDS, http://gsds.cbi.pku.edu.cn) for intron and exon distribution in each gene. MEME suite (http://meme-suite.org) was used to predict conserved motifs in SIPHB protein sequences with a parameter set as follows: (i) a maximum number of motifs – 10, (ii) number of repetitions – any, (iii) optimum motif width set to ≥10 and ≤50. A 1000bp 5’ UTR nucleotide sequences from the start codon (ATG) of each SIPHB gene were retrieved from the SOL genome and scanned in the PlantCRAE database (http://bioinformatics.psb.ugent.be/webtools/plantcare/html) for cis-regulatory elements prediction.

### 2.5. Plant Material, Abiotic Stress, and Phytohormone Treatment

Tomato cv. Micro-Tom seedlings were grown in the College of Agriculture and life sciences, Kunming University, under controlled greenhouse conditions (25°C/20°C, day/night, 14 h/10 h light/dark photoperiod with relative humidity 80%). For tissue/organ-specific expression analysis of various plant parts such as root, leaves, stem, and flowers were collected from a six-week-old plant. For expression in fruit tissues, 1/2/3/cm, mature green fruit, breaker fruit, and ten days breaker fruits were harvested. For salinity, drought, and phytohormone-induced stresses, six-week-old plants were treated with 200 mM NaCl, 0.01 mM ABA, GA3, IAA, MeJA, and PEG as described previously. Roots and shoots (including stem and leaves) were harvested at 0 h, 3 h, 6 h, 12 h, and 24 h interval after treatment. All the samples were collected in triplicate and store immediately at –80°C.

### 2.6. Total RNA Extraction, cDNA Preparation, and qRT-PCR Analysis

Total RNA was extracted from selected samples using TRIZOL reagent according to the
manufacturer’s instruction. RNA was quantified using nanodrop lite (Thermo USA) and RNA integrity was assessed by running 2% agar agarose gel electrophoresis. The cDNA was synthesized with a PrimerScript Real-Time (RT) reagent kit (Takara, Japan) according to the manufacturer’s protocol as described previously.\textsuperscript{51–53} RT-qPCR was conducted in ABI 7500 Fast Real-Time system (AB, USA) using the iTaq\textsuperscript{™} Universal SYBR\textsuperscript{®} Green Supermix (BIO-RAD, USA) according to the manufacturer’s protocol. The RT-qPCR was conducted in triplicate. Tomato \textit{SlUBQ} (Solyc01g056940) gene was used as an internal control. The relative expression of tomato \textit{SlPHBs} was calculated using the 2\textsuperscript{−ΔΔCt} method\textsuperscript{54} and heat maps were generated with heat mapper program (http://www1.heatmapper.ca/ expression/).

### 2.7. Subcellular Localization of \textit{SlPHB5} and \textit{SlPHB10}

The full-length sequences of \textit{SlPHB5} and \textit{SlPHB10} excluding stop codon were fused into the vector p35S-GFP as explained previously.\textsuperscript{51,55} The Arabidopsis protoplast isolation and transformation were carried out as described by Sheen.\textsuperscript{56} After 18–20 h of transformation, the protoplast was visualized by confocal laser scanning microscope and the images were processed using photoshop.

### 3. Results

#### 3.1. Identification of \textit{SlPHB} Genes

The Arabidopsis, rice, \textit{Zea mays}, and \textit{glycine max} PHB protein sequences were used as a query in the SOL genome to identify all putative PHB protein sequences in the tomato genome. A total of 16 non-redundant genes were identified. The Pfam, SMART, and NCBI CDD searches were used to verify the SPFH domain in all \textit{SlPHB} protein sequences. The tomato PHB genes were named as \textit{SlPHB1} to \textit{SlPHB16} in order of their position in chromosomes. The peptide length to the molecular weight of \textit{SlPHBs} ranged from 261 aa (\textit{SlPHB8}) to 518 aa (\textit{SlPHB7}), and 30.08 kDa (\textit{SlPHB1}) to 57.75 kDa (\textit{SlPHB7}). The GRAVY values of all the \textit{SlPHB} proteins were negatively exhibiting indicating that these proteins are hydrophilic except \textit{SlPHB15} (Solyc11g013260) which show a positive GRAVY score. The deduced \textit{SlPHB} genes were distributed in seven chromosomes (Fig. 1(a)). A pair of genes \textit{SlPHB1} and \textit{SlPHB2}, \textit{SlPHB8} and \textit{SlPHB9}, \textit{SlPHB14}, and \textit{SlPHB15} were located on chromosomes 1, 5, and 11 each, respectively. \textit{SlPHB3}, \textit{SlPHB4}, \textit{SlPHB5}, \textit{SlPHB6}, and \textit{SlPHB7} were located on chromosome 3. Three genes (\textit{SlPHB10}, \textit{SlPHB11}, \textit{SlPHB12}) were located on chromosome 6 while a single gene was located on chromosome 10 (\textit{SlPHB13}) and chromosome 12 (\textit{SlPHB16}) each. In silico subcellular location, prediction indicated that \textit{SlPHBs} were localized in the cytoplasm, mitochondria, and chloroplast (Table 1). Tomato PHB genes displayed segmental duplication and five segmental gene duplication (eight genes) were found in tomato as shown in Fig. 1(b).

#### 3.2. Phylogeny, Strong Purifying Selection, and Conserved Motif Analysis of \textit{SlPHB} Proteins

To unveil the phylogenetic relationship of tomato \textit{SlPHB} proteins with PHBs from other plant species such as \textit{Arabidopsis}, rice, maize, and soybean, an unrooted neighbor-joining phylogenetic tree was generated. It was observed that all PHB proteins were divided into four major clades (II, III, IV, and V). The subclade of each group contains 7–15 members from different species. \textit{SlPHBs} were found in all clades such as five \textit{SlPHBs} in group IV (2 in IV B and 3 in IV A). Similarly, three in subclade V B and single in V A subclade of major clade V. Moreover, clade III has four, and clade II contained two tomato \textit{SlPHBs}. Similar trends of PHB distribution were observed for other species (Fig. 2). Furthermore, three sister pairs of \textit{SlPHB} genes were detected in the phylogenetic tree such as \textit{SlPHB14}/\textit{SlPHB15} in clade IV A, \textit{SlPHB2}/\textit{SlPHB3} in subclade V B of major clade V, and \textit{SlPHB11}/\textit{SlPHB6} in clade III. It was observed that \textit{SlPHBs} localized in chloroplast were clustered together as shown in Fig. 3(a).

A comparison of the gene structure of each tomato PHB revealed a diverse structure. The number of intron and exon ranged from one to nine exons and zero to eight introns. The exon/intron pattern was similar in different clades and subclades. For example, five exon and four introns were found in clade III, nine exons and eight introns in clade II, and clade V. Similarly, five
exons in subclade IV B and two in IV A clade. Besides, the length and positions of exons were also highly similar in clades and subclade Fig. 3 (b). We identified ten conserved motifs in SIPHBs using the MEME server. It was observed that the motifs pattern was also similar within clades (Fig. 3 (c)). For instance, motif 1 and motif 2 found in clade V B; motif 1, motif 2, motif 3, motif 7, motif 8, and motif 9 in clade IV. SIPHBs in clade III contained all motifs except motif 9. To explore the fate of divergence of these genes in the tomato genome, the Ka/Ks values were estimated for three duplicate SIPHB gene pairs. The Ks was used in estimating the divergence time of each SIPHB gene pairs.
Table 1. The characteristic features of tomato SIPHB proteins in tomato genome.

| Gene locus ID | Gene Name | aa | MW  | pI   | GRAVY | Position | Start  | End    | Sub-cellular Location |
|---------------|-----------|----|-----|------|-------|----------|--------|--------|-----------------------|
| Solyc01g010770 | SlPHB1 | 272 | 30.08 | 4.55 | −0.118 | 1 | 5825560 | 5828696 | Cysk |
| Solyc01g089910 | SlPHB2 | 490 | 54.12 | 7.22 | −0.308 | 1 | 75383651 | 75385711 | Cyto |
| Solyc03g005420 | SlPHB3 | 489 | 54.67 | 8.99 | −0.419 | 3 | 296785 | 298965 | Cyto |
| Solyc03g007190 | SlPHB4 | 290 | 32.32 | 4.89 | −0.234 | 3 | 1764428 | 1769612 | Cyto |
| Solyc03g080050 | SlPHB5 | 424 | 46.49 | 9.48 | −0.302 | 3 | 45473113 | 45480181 | Mito |
| Solyc03g113220 | SlPHB6 | 285 | 31.36 | 5.73 | −0.087 | 3 | 57486387 | 57489392 | Cyto |
| Solyc03g117250 | SlPHB7 | 518 | 57.75 | 6.25 | −0.389 | 3 | 60489302 | 60492373 | Cyto |
| Solyc05g012340 | SlPHB8 | 261 | 41.09 | 6.73 | −0.377 | 5 | 5601824 | 5605948 | Chlo |
| Solyc05g015100 | SlPHB9 | 277 | 30.31 | 7.67 | −0.016 | 5 | 61017078 | 61019992 | Chlo |
| Solyc06g063850 | SlPHB10 | 484 | 53.94 | 5.26 | −0.402 | 6 | 37670367 | 37673877 | Mito |
| Solyc06g071050 | SlPHB11 | 289 | 31.82 | 5.22 | −0.119 | 6 | 40043522 | 40046056 | Cyto |
| Solyc06g073030 | SlPHB12 | 398 | 44.54 | 9.14 | −0.429 | 6 | 41388909 | 41390901 | Chlo |
| Solyc10g008140 | SlPHB13 | 289 | 31.83 | 10.11 | −0.154 | 10 | 2276303 | 2278610 | Chlo |
| Solyc11g010190 | SlPHB14 | 279 | 30.68 | 9.4 | −0.057 | 11 | 3269915 | 3270754 | Chlo |
| Solyc11g013260 | SlPHB15 | 301 | 32.95 | 9.55 | 0.059 | 11 | 6170897 | 6173381 | Chlo |
| Solyc12g005500 | SlPHB16 | 283 | 31.16 | 10.08 | −0.201 | 12 | 293644 | 295414 | Chlo |

aa; amino acid, MW; molecular weight, pI; isoelectric point, GRAVY; the grand average of hydrophathy, Cysk; cytoskeleton, Cyto; cytoplasm, Chlo; Chloroplast, Mito; mitochondria, Extr; extracellular cytoplasm.

Figure 2. The phylogeny of the PHB proteins. An unrooted neighbor-joining phylogenetic tree of PHB proteins from Arabidopsis, rice, maize, soybean, and tomato was generated in the MEGA program with a bootstrap value set as 1000 replicates. The tree was clustered into various clades and subclades. The black dots represent tomato SIPHB proteins.
Figure 3. Phylogeny, gene exon/intron distribution, and conserved motif analysis of 16 tomato SIPHB genes. (a) An unrooted neighbor-joining phylogenetic tree of PHB proteins with bootstrap set at 1000 replicates and clustered into different clades and subclades. (b) Tomato SIPHB gene intron and exon distribution. The scale at the bottom is corresponding to gene size in kb. (c) The putative conserved motifs in 16 tomato PHB proteins identified using the MEME suite. A total of ten motifs (1 to 10) were identified and each color of the box is corresponding to a motif. The scale at the bottom represents the protein size in kb.

Table 2. The Ka/Ks of tomato SIPHB paralogs.

| Gene1       | Gene2       | Ka         | Ks         | Ka/Ks      | Time (Mya*) | Purify Selection |
|-------------|-------------|------------|------------|------------|-------------|-----------------|
| Solyc01g089910 | Solyc03g005420 | 0.206309991 | 1.247723962 | 0.165349065 | 95.1009117    | Yes             |
| Solyc03g113220 | Solyc06g071050 | 0.042588715 | 0.483257784 | 0.088128358 | 36.83367252   | Yes             |
| Solyc11g010190 | Solyc11g013260 | 0.067198744 | 1.332405685 | 0.050434147 | 101.553114    | Yes             |

*millions year ago

(Table 2). Our results showed that the Ka/Ks ratio of duplicated genes pairs was more than 0.04. This suggesting that the purifying selection pressure was a major factor that occurs during the evolution, function divergence was limited after duplication and was estimated to occur between 36.8 and 101.55 million years ago (Mya).

3.3. Bioinformatics Analysis of SIPHB Promoter Sequences

The cis-acting elements of potential tomato SIPHB genes were predicted by searching a 1000 bp region from the transcriptional activation site (ATG) of each gene against the PlantCARE database. As shown in Fig. 4, several putative cis-regulatory sequences were identified in SIPHB genes. For an instance, four different kinds of development-related cis-regulatory elements such as circadian control (circadian), meristem development (CAT-box), endosperm development (GCN4_motif), and zein metabolism regulation (O2-site) were predicted in the promoter region of some of the SIPHBs, suggesting that these genes may play roles in organ/tissue-specific development and growth. Moreover, a various stress-responsive element such as the MYB binding site involved in drought-inducibility (MBS), WRKY binding site involved in abiotic stress and defense response (W-box), anaerobic induction element (ARE), defense- and stress-responsive element (TC-rich repeats), low-temperature-responsive element (LTR), wound-responsive element (WUN-motif), and element for maximal elicitor-mediated activation (AT-rich sequence) were also detected. The promoters of tomato SIPHB genes possessed cis-regulatory sequences related to ethylene (ERE), suggesting that these genes may involve in ethylene responses (Fig. 4). In addition, various hormone-related responsive elements related to gibberellin (GARE-motif), methyl jasmonate (MeJA, CGTCA-motif), abscisic acid (ABRE), and salicylic acid (TCA-element) were
also detected, implying that these genes may respond to phytohormone as well (Fig. 4). The promoters of tomato *SlPHB* genes possessed *cis*-regulatory sequences related to ethylene (ERE), suggesting that these genes may involve in ethylene responses.

### 3.4. miRNAs Targeting the PHB Family Members of the Tomato

To find out miRNAs targeting the SIPH Bs of tomato, the sequences were subjected to the miRNA database. The psRNATarget predicted that four SIPH B genes family members were targeted by conserved miRNAs belongs to different miRNAs gene families each. *SlPHB7* was targeted by the sly-miRNA869 family and sly-miRNA4239 cause the cleavage of *SlPHB3*. A single member from sly-miR396 and sly-miR397 family member target to cleavage of *SlPHB15* and *SlPHB13* gene, respectively (Table S1).

### 3.5. Expression Analysis of SIPH B Genes in Different Plant Parts

To understand the role of putative SIPH Bs in tomato plant growth and development, the expression profile analysis of SIPH Bs in various plant parts was evaluated. The SIPH Bs exhibited a diverse expression pattern among various plant parts. It was found that two SIPH Bs were expressed in leaves, and root tissues. One *SlPHB* gene had high expression levels in fully opened flower and three expressed in flower at bud condition. It was observed that the number of genes was expressed in fruit at different development stages with more and less expression levels. For example, *SlPHB1* in 3 cm fruit, *SlPHB6* in ten days fruit breaker, *SlPHB8*, and *SlPHB9* in 2 cm fruit. However, *SlPHB5*, *SlPHB14*, and *SlPHB15* exhibited increasing expression during fruit development and ripening (2 cm fruit till ten days breaker fruit) (Fig. 5). The results showed that tomato SIPH B genes play an important role in the growth and development of specific plant parts or tissues.

### 3.6. Expression Profile of Tomato SIPH B in Response to Salinity and Drought Stress

To further investigate the role of SIPH B in tomato against abiotic stresses, the expression profile of SIPH B in response to salt and drought was analyzed at various time points. It was observed that under salt stress, the transcript abundance of *SlPHB9* was
sharply increased at 3 h and peak at 6 h time point and subsequently declined at 12 h and 24 h time points. ****SIPHB7** and **SIPHB11** had maximum transcript levels at 24 h while, **SIPHB4**, **SIPHB13**, and **SIPHB14** exhibited transcript abundance at 12 h time point. **SIPHB5** and **SIPHB8** induced only at 3 h after treatment but **SIPHB10** induced at 6 h time point (Fig. 6(a)). Under drought conditions, the majority of genes were expressed at the late time point (12 h and 24 h). **SIPHB2** and **SIPHB9** induced only at 6 h after treatment (Fig. 6(b)). In comparison, **SIPHB5**, **SIPHB13**, **SIPHB14**, **SIPHB15**, **SIPHB9**, and **SIPHB7** showed similar trends of expression under both drought and salinity stresses but **SIPHB4**, **SIPHB2**, and **SIPHB8** exhibited opposite trends under both stresses (Fig. 6). These results suggest that tomato **SIPHB** genes may play a key role in regulating abiotic stress responses.

### 3.7. Phytohormone Induced Expression Profile Analysis of SIPHBs in Tomato

To check the effectiveness of exogenous phytohormone application, the expression profile of tomato **SIPHB** under various hormones such as abscisic acid, gibberellin, auxin, and methyl jasmonate was examined. For ABA treatment, **SIPHB13** and **SIPHB15** were induced at 3 h time points while **SIPHB6** and **SIPHB12** were upregulated at 6 h after application with decreased expression in later time points. **SIPHB11** expression was downregulated upon treatment with ABA but **SIPHB8**
and SlPHB16 were induced only at 12 h after treatment. Moreover, SlPHB9, SlPHB10, SlPHB4, SlPHB12, and SlPHB3 was upregulated at 24-time points (Fig. 7(a)). SlPHB5, SlPHB14, and SlPHB16 transcript levels were sharply induced at 3 h interval and reach a maximum at 6 h time point but decreased in subsequent time intervals to GA3. SlPHB3, SlPHB4, SlPHB5, SlPHB11, SlPHB12, SlPHB1, and SlPHB2 were induced with maximum transcript levels at 12 h after exposure to GA3 (Fig. 7(b)). The transcript abundance of SlPHB7 and SlPHB8 was increased temporally but SlPHB13 expression was downregulated upon treatment with GA3. For auxin, SlPHB10 and SlPHB14 genes were downregulated after application but SlPHB3 showed maximum transcript accumulation at 3 h point interval. SlPHB5, SlPHB12, SlPHB8, SlPHB13, and SlPHB16 was upregulated with time and reached maximum expression at 6 h after treatment while, SlPHB15, SlPHB4, SlPHB9, SlPHB2, and SlPHB11 expression levels were upregulated across 6 h to 24 h time points and showed maximum expression at 24 h interval (Fig. 7(c)). The SlPHB8s exhibited a unique expression profile upon exposure to MeJA. It was observed that all the genes were upregulated temporally across all time intervals and have high transcript accumulation at 24 h time point except for SlPHB4 (Fig. 7(d)). The data suggest that tomato SlPHB genes may play various important roles in cross-talk with different kinds of hormones signaling.

3.8. Subcellular Localization Assay

The amino acid sequence of SlPHB5 and SlPHB10 was submitted to the WoLFPSORT (https://wolfpsort.hgc.jp/) to predict subcellular localization. The predicted results showed that both SlPHB proteins were expressed in the mitochondria. To experimentally verify, full-length sequences of candidate SlPHB5 and SlPHB10 were fused to a GFP reporter gene and transferred to Arabidopsis protoplast (Fig. 8). Subcellular localization experiment results revealed that both proteins were localized in the mitochondria as predicted. LoTPS3 protein from Lilium Siberia was used as a positive control.

Scale bar 5 μm.

4. Discussion

PHB, a highly conserved multigene family has been identified in many organisms from humans to various plant species playing essential roles in various aspects of growth and development. In plants, the PHB gene family has been reported from Arabidopsis (17), rice (19), Glycine max (24),...
and *Zea mays* with 16.\(^{17}\) However, no genome-wide identification of the PHB gene family has been reported in the tomato genome. In this study, a total of 16 PHB genes were identified in the tomato genome (Table 1). The tomato genome size (960Mb) is 7.68 folds of the *Arabidopsis* genome (125 Mb), 2.46 folds of rice (389 Mb) but 2.3 folds less of maize (2300 Mb) and 1.14 folds less than soybean (1100 Mb) genome. However, the number putative PHBs in the tomato genome even lower than *Arabidopsis* and rice\(^{31}\) but equal to reported in maize.\(^{17}\)

Gene duplication either segmental or tandem plays an important role in the expansion of the genome. The expansion of the PHB gene family in *Arabidopsis*, rice, and soybean was caused by segmental duplication while tandem duplication was another cause of an increasing number of PHB genes in *Arabidopsis* but was absent in tomato. This implying that gene duplication of the PHB gene family in tomato was

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**Figure 7.** Phytohormone induced expression profile of *SlPHB* genes. (a) abscisic acid (ABA), (b) gibberellin (GA3), (c) auxin (IAA), (d) methyl jasmonate (MeJA) induced expression profile at 0 h, 3 h, 6 h, 12 h, and 24 h time points. A log2 transformed heatmap was generated using heatmapper program. Blue, white, and red color is corresponding to low, moderate, and high expressions. The genes were clustered by applying the Euclidean method.
different from *Arabidopsis*. We have analyzed Ka/Ks values of three pairs of SIPHB gene duplication and found that tomato PHB genes undergo purifying selection (Table 2).

The PHB genes from fungi and mammals including humans were clustered in five phylogenetic clades. However, like *Arabidopsis*, rice, Glycine max, and Zea mays, tomato SIPHBs were also clustered in four clades. The genes sharing clades and subclades displayed a similar gene structure and conserved motifs patterns. PHB genes are involved in various aspects of plant growth and development. In this study, cis-regulatory sequences were predicted. It was observed that tomato SIPHB genes contained various development, abiotic stress, and phytohormone responsive elements in their promoter regions (Fig. 4). It has been well documented that PHB genes involved in leaf yellowing, hormone signal transduction pathways, and abiotic stress responses. For example, *Arabidopsis* AtPHB3/4 causes proliferation of root and shoot tissues. Similarly, *petunia* PHBs, tobacco *NbPHB1/2* promote leaf senescence. In this study, the expression profile of *SLPHBs* in various parts of tomato plant was also investigated. Tomato SIPHB genes showed diverse expression patterns in different parts such as *SIPHB4* and *SIPHB10* was expressed in flower and root tissues, respectively. Two genes (*SIPHB8, SIPHB9*) were highly expressed in 2 cm fruit while *SIPHB5, SIPHB14*, and *SIPHB15* showed increasing expression pattern with the fruit development stages (Fig. 5). These results suggest the crucial role of SIPHB genes in development of these organs in tomato plant.

In this study, cis-regulatory elements involved in diverse signaling pathways were identified. Most PHBs contain cis-regulatory elements involved in ABA, GA, JA, and ethylene. In addition, cis-elements involved in abiotic stresses, such as MBS (MYB binding site involved in drought-inducibility), LTR (low-temperature responsiveness element), HSE (heat stress responsiveness element), were also observed in the promoter regions of SIPHB genes (Fig. 4). In Glycine max, most of PHBs contained numerous hormone-responsive, development and stress-related cis-regulatory elements in the GmPHB promoters. It was observed that the expression of SIPHB genes was altered under these stresses. For salt treatment, *SIPHB5, SIPHB8, SIPHB9*, and *SIPHB10* were upregulated at early time points (3 h and 6 h) while, *SIPHB7, SIPHB11,*
SIPHBA, SIPHB13, SIPHB14, and SIPHB12 were induced at 12 h and 24 h after treatment (Fig. 6(a)). Similar response was observed in Arabidopsis, where PHBs were involved in abiotic stimulus and phytohormones functioning. SIPHB2 and SIPHB9 genes were induced under drought at 6 h time point but SIPHB4 was downregulated (Fig. 6(b)). SIPHB1, SIPHB14, SIPHB9, SIPHB10, SIPHB4, and SIPHB3 were upregulated after 24 h exposure to ABA but SIPHB11 and SIPHB15 were downregulated upon exposure (Fig. 7(a)). Moreover, SIPHB13, SIPHB14, SIPHB15, and SIPHB16 were suppressed in late intervals of GA3 exposure but the rest of the genes were upregulated (Fig. 7(b)). SIPHB10 and SIPHB14 were downregulated after auxin application but, SIPHB3 was induced after 3 h of treatment. SIPHB7, SIPHB1, and SIPHB6 exhibited maximum expression at a 24 h time point (Fig. 7(c)). For MeJA treatment, all the genes were induced sharply along with all the time points and peaked at 24 h after treatment except for SIPHB4, which was suppressed upon exposure to MeJA (Fig. 7(d)). Likewise, Atphb3 mutant was highly responsive to ethylene in etiolated seedlings. One Arabidopsis prohibitin (At5g64870) was down-regulated under some hormones (GA, MeJA and ABA), while highly upregulated under salt, drought and cold treatment. In Capsicum annuum, hypersensitive-induced reaction (HIR) proteins (PHB encoding proteins), such as CaHIR1, maize ZmHIR1-3, barley HvHIR1/3 and AtHIR1-3 were induced under abiotic stresses. Our findings are in line with previous studies that PHB genes showed differential expression pattern under different development stages as well as under different stimulus. The above-mentioned findings highlighted the potential diverse role of PHB genes.

5. Conclusion

In short, this study provides knowledge about the PHB gene family in the tomato genome. All the identified SIPHBs were clustered in four clades according to the phylogenetic tree. The gene structure and conserved motifs distribution patterns in each clade validated the phylogenetic classification of tomato SIPHBs. Cis-regulatory sequences prediction in combination with complex regulation of tomato PHB genes family expression against salinity, drought, and various phytohormones such as ABA, IAA, GA, and MeJA provide a foundation for further functional characterization of these genes in tomato and other plant species.

Disclosure Of Potential Conflicts Of Interest

The author(s) declare neither financial nor non-financial conflict of interest.

Author Contributions

All authors contributed to the study conception and design. Conceptualization: Yanguo Ke, Feiyang Huang and Lei Y; Methodology: Zhijiang Wang, Yan Ding and Muhammad Waseem; Formal analysis and investigation: Muhammad Waseem, Lei Yu, Xianjie Cai, Xianwen Ye, Umair Ashraf and Yanguo Ke; Writing - original draft preparation: Yanguo Ke, Farhat Abbas, Muhammad Waseem and Feiyang Huang; Writing - review and editing: Yanguo Ke, Xiaolong Chen, Umair Ashraf and Farhat Abbas; Funding acquisition: Xianjie Cai, Feiyang Huang and Yanguo Ke; Resources: Yanguo Ke, Xiaolong Chen and Feiyang Huang.

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