A Glucose-6-Phosphate/Phosphate Translocator Is Involved in Carbohydrate Metabolism and Starch Biosynthesis in Sweet Potato (Ipomoea Batatas (L.) Lam.)

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Abstract

Sweet potato (*Ipomoea batatas* (L.) Lam.) is a good source of carbohydrates, an excellent raw material for starch-based industries, and a strong candidate for biofuel production due to its high starch content. However, the molecular basis of starch biosynthesis and accumulation in sweet potato is still insufficiently understood. Glucose-6-phosphate/phosphate translocators (GPTs) mediate the import of glucose-6-phosphate (Glc6P) into plastids for starch synthesis. Here, we report the isolation of a GPT-encoding gene, *IbG6PPT1*, from sweet potato and the identification of two additional *IbG6PPT1* gene copies in the sweet potato genome. *IbG6PPT1* encodes a chloroplast membrane–localized GPT belonging to the GPT1 group and highly expressed in storage root of sweet potato. Heterologous expression of *IbG6PPT1* resulted in increased starch content in the leaves, root tips, and seeds and soluble sugar in seeds of *Arabidopsis thaliana*, but a reduction in soluble sugar in the leaves. These findings suggested that IbG6PPT1 might play a critical role in carbohydrate accumulation in storage tissues and would be a good candidate gene for controlling critical starch properties in sweet potato.

Background

Sweet potato (*Ipomoea batatas* (L.) Lam.) is an important food crop that is cultivated in over 100 countries due to its stable yield, rich nutrient content, low input requirements, multiple uses, high yield potential, and adaptability under a range of environmental conditions (Burri, 2011; Hu et al., 2003; Mitra, 2012; Yang et al., 2020; Zhang et al., 2017). Sweet potato is grown mainly for its edible, starchy storage root, which is 50–80% starch by dry matter (Zhou et al., 2015). This high starch content renders sweet potato a good source of carbohydrates, an excellent raw material for starch-based industries, and a strong candidate as an inexpensive raw material for biofuel production (Koçar and Civaş, 2013; Nedunchezhiyan et al., 2012; Srichuwong et al., 2012). Although sweet potato is one of the top starch-rich root crops worldwide (Wang et al., 2019), the molecular basis of starch biosynthesis and accumulation in this species is poorly understood.

Starch is synthesized in plants through a complex pathway involving multiple enzymes and transporters (Lai et al., 2016; Schreiber et al., 2014; Zhang et al., 2017). Starch biosynthesis begins with the synthesis of sucrose, the major product of photosynthesis, in source tissues. During this process, sucrose can be converted to glucose-6-phosphate (Glc6P) and then imported into the plastid by glucose-6-phosphate/phosphate translocators (GPTs), proteins belonging to the transporter subfamily of phosphate translocators (PTs).

Two functional GPT genes have been identified in Arabidopsis (*Arabidopsis thaliana*). GPT1 is localized on the inner envelope of the plastid membrane and transports Glc6P into plastids for use in starch biosynthesis, the oxidative pentose phosphate pathway (OPPP), and fatty acid biosynthesis (Kunz et al., 2010). GPT1 is also localized to peroxisomes, where it preferentially exchanges Glc6P for ribulose-5-phosphate (Ru5P) (Baune et al., 2020).
GPTs play important roles in several physiological processes. Arabidopsis \textit{GPT1} is essential for the development of male and female gametophytes, embryos, and seeds (Chen et al., 2019; Zhang et al., 2020). GPT1 is highly expressed at the late stages of pollen development, where it drives Glc6P from the cytosol and into plastids for fatty acid biosynthesis, and thus plays an important role in lipid body biogenesis during pollen maturation (Zheng et al., 2018). By contrast, Arabidopsis \textit{GPT2} is expressed when photosynthesis is increased by light, which allows increased net import of Glc6P from the cytosol to chloroplasts, thus facilitating starch synthesis during stochastic high-light conditions (Dyson et al., 2015; Weise et al., 2019). \textit{GPT2} responds rapidly by glucose and sucrose induced and plays an essential role in interpreting environmental signals (Chen et al., 2019; Weise et al., 2019). GPT1 plays a major role in the regulation of starch synthesis in other plants. GPT1 in Narbonne vetch (\textit{Vicia narbonensis}) is critical for starch synthesis and storage in developing seeds. In \textit{Vicia} transgenic plants expressing antisense GPT1 via \textit{Agrobacterium}-mediated transformation, amyloplasts developed later and were smaller in size, starch biosynthesis was reduced, and storage protein biosynthesis increased (Rolletschek et al., 2007). In rice, pollen grains from homozygous \textit{osgpt1} mutant plants fail to accumulate starch granules, resulting in pollen sterility (Qu et al., 2021). However, the role of GPTs in sweet potato has not been investigated.

In our previous work, we identified \textit{GPT} genes in sweet potato using transcriptome analysis and showed that they have different expression patterns during storage root development and among sweet potato genotypes with different starch properties. Therefore, these GPTs are probably involved in starch accumulation and sucrose metabolism in sweet potato (Zhang et al., 2017). Here, we cloned two \textit{GPT} genes and analyzed their protein localizations, sequence features, and functions. Our results provide important insights into the mechanisms underlying the starch properties of sweet potato.

**Results**

*Two GPT-encoding genes were cloned from sweet potato*

To ensure that the full-length mRNA sequence of sweet potato \textit{GPT} genes could be obtained, the RACE method was used for cloning. Two cDNA sequences encoding the sweet potato \textit{GPT} genes were obtained, named \textit{IbG6PPT1} and \textit{IbG6PPT1-2}. The obtained sequences contained 1763 and 1767 bp of mRNA, corresponding to 1191 and 1200 bp of ORFs and encoding 399-aa and 396-aa protein sequences, respectively. The two genes shared 96.627\%, 98.083\%, and 98.747\% identity at the mRNA, CDS, and putative amino acid levels, respectively. The two proteins differed in only five amino acids (Fig. 1), including a deletion of the L\textsubscript{37}P\textsubscript{38}A\textsubscript{39} sequence in the shorter GPT.

The sweet potato genome is annotated with three \textit{GPT1} gene members: \textit{IbG6PPT1} located on chromosome 3 (chr3), \textit{IbG6PPT1-2} located on chr2, and another \textit{IbG6PPT1}-like gene also expected to be located on chr2. However, the sequence of this \textit{IbG6PPT1}-like gene was not cloned from our cDNA library. Amino acid differences between \textit{IbG6PPT1} and \textit{IbG6PPT1-2} were not located at conserved domains or important transmembrane domains, indicating that these proteins are likely functional.
The cloned GPT genes belong to the GPT1 group

The GPT subfamily is relatively well characterized at the molecular level. The GPT subfamily includes three groups: GPT1, GPT2, and GPT3 (Chadee and Vanlerberghe, 2020). The sweet potato GPT genes showed 98.75% and 97.99% identity with Ipomoea nil and Ipomoea triloba GPT1, respectively. A GPT phylogenetic tree showed that the GPT1 group consisted of two sweet potato GPT proteins as well as Ipomoea triloba, morning glory (Ipomoea nil), tobacco (Nicotiana tabacum), potato (Solanum tuberosum), tomato (Solanum lycopersicum), China rose (Rosa chinensis), Arabidopsis thaliana, rice (Oryza sativa), and maize (Zea mays) GPT1 proteins. The GPT2 group consisted of AtGPT2, whereas the GPT3 group mainly consisted of two N. tabacum GTP3 proteins, XP_016451801.1 and XP_016454155.1 (Chadee and Vanlerberghe, 2020) (Fig. 2). Therefore, the obtained sweet potato GPTs belong to the GPT1 group.

The protein sequence alignment and phylogenetic tree showed that IbG6PPT1 was more similar than IbG6PPT1-2 to GPT1 proteins from Ipomoea nil and Ipomoea triloba (Fig. 1 and Fig. 2), indicating that IbG6PPT1 might match previous GPT1 findings better in the Ipomoea genus. Thus, we focused on IbG6PPT1 for the remainder of this work.

IbG6PPT1 is likely a chloroplast-located GPT

We created a GFP-tagged version of IbG6PPT1 and transiently expressed it in N. benthamiana. The GFP signal surrounded the chloroplast marker fluorescence, indicating that IbG6PPT1 localizes to the chloroplast membrane (Fig. 3). No signal peptide was predicted from the protein sequence, meaning that the protein wasn't secreted protein. TMPred and TMHMM predicted the presence of seven transmembrane helixes (Fig. 1), consistent with a role for IbG6PPT1 in Glc6P transport through the chloroplast membrane. Modeling of the three-dimensional (3D) structure of IbG6PPT1 predicted that two IbG6PPT1 proteins form a homodimer (Fig. 4). In addition, IbG6PPT1 contains a conserved sugar phosphate transporter domain (Jack et al., 2001). These results strongly suggest that IbG6PPT1 is a chloroplast membrane–localized, GPT family protein in sweet potato.

IbG6PPT1 is highly expressed in sweet potato storage root

We next attempted to determine whether and how strongly IbG6PPT1 is expressed in various sweet potato tissues by extracting RNA from the petiole, stem, leaf, and storage root and performing RT-qPCR to quantify gene expression. IbG6PPT1 was expressed in all tissues but showed its highest expression in storage roots, followed by the petiole, stem, and leaves (Fig. 5). Interestingly, the higher expression of IbG6PPT1 in roots than in leaves suggests that it may function in Glc6P transport in non-green tissues rather than in photosynthetic tissues.

Heterologous expression of IbG6PPT1 affects starch and sucrose content

To identify the function of IbG6PPT1, we transformed a p35S::IbG6PPT1-YFP construct into wild-type (Col-0) Arabidopsis. Analysis of IbG6PPT1-YFP expression by qPCR and western blotting showed that the
fusion protein was heterologously expressed across four independent transgenic lines, designated OV-14, OV-30, OV-76, and OV-57, that were selected from the T<sub>2</sub> progeny, as well as in the wild type (Fig. 6a and 6b). There were no differences in growth and development between the transgenic progeny and the wild-type control (Fig. 6c).

In contrast to their wild-type-like appearance, the soluble sugar content in the leaves of the transgenic <i>IbG6PPT1-YFP</i> lines was only 76.59–83.40% of control (Fig. 7a, Table S1). Meanwhile, the leaves of the 6 weeks transgenic T<sub>2</sub> plants had a measured starch content 1.65- to 2.75-fold higher than the control (Fig. 7b, Table S1), which was confirmed by iodine staining in 3 weeks seedlings (Fig. 7c). Surprisingly, the 1000 seed weights of the transgenic lines were 1.06- to 1.19-fold higher than in the control plants (Fig. 7d, Table S1). Further analyses showed that the soluble sugar content and starch content in the seeds of transgenic <i>IbG6PPT1-YFP</i> lines were 1.20- to 1.47-fold and 1.13- to 1.31-fold higher than in the control plants, respectively (Fig. 7e and 7f, Table S1). In the root tips, iodine staining showed that the starch content of transgenic <i>IbG6PPT1-YFP</i> lines was higher than that in control plants (Fig. 7g and 7h). Above all, heterologous expression of the <i>IbG6PPT1</i> gene increased both starch and sucrose content in Arabidopsis.

**Discussion**

<i>IbG6PPT1</i> is present in several gene copies that may have different functions

The sweet potato genome is allohexaploid (2n = 6x = 90), containing two B1 and four B2 component genomes (B1B1B2B2B2B2) (Gao et al., 2020; Yang et al., 2017; Zhang et al., 2020). Therefore, there may be up to six copies of each gene. In this study, we cloned two <i>GPT1</i> genes that share a high level of identity in both the mRNA and protein sequences (Fig. 1). However, we found three potential <i>IbG6PPT1</i> genes in the genome database, the two we cloned and another one on chr2 that might be a homolog or paralog of one of the cloned genes. During the evolution of sweet potato's polyploid genome, the duplicated genes might have developed expressional, regulatory, or functional divergence (Fan et al., 2017; Wesemael et al., 2018). Because of the very high sequence similarity between the <i>IbG6PPT1</i> genes, it is difficult to examine the expression pattern or function of a single such gene. Future work should investigate whether the three <i>IbG6PPT1</i> genes show functional divergence in Glc6 transport and thus play different roles in starch accumulation and sucrose metabolism in sweet potato. Alternately, they may not have diverged as strongly, and one gene's function may have been compensated for by the function of another gene. Future genetic engineering of the sweet potato will require gene function studies to determine the contribution of each gene copy to relevant phenotypes and identify the major gene controlling sweet potato starch properties.

<i>IbG6PPT1</i> has similar functions to other GPT1 proteins

GPT1 proteins transport Glc6P into plastids for starch and/or fatty acid biosynthesis, depending on the plant species (Zheng et al., 2018). A previous report demonstrated that starch biosynthesis is mediated
by *VnGPT1* in pea embryos (Rolletschek et al., 2007). Starch is the major carbon storage molecule of sweet potato, accounting for 50–80% of dry matter in the storage root, the organ that determines sweet potato's economic value as a crop (Zhang et al., 2017), whereas fatty acids are almost undetectable. In Arabidopsis, fatty acid biosynthesis in pollen is controlled by regulating *AtGPT1* expression through the MKK4/MKK5-MPK3/MPK6 cascade and the downstream transcription factors WRKY2 and WRKY34 (Zheng et al., 2018). In rice, pollen grains from homozygous *osgpt1* mutant plants fail to accumulate starch granules, resulting in pollen sterility (Qu et al., 2021). Like *AtGPT1*, which is expressed ubiquitously throughout Arabidopsis development (Niewiadomski et al., 2005), we found that *IbG6PPT1* is expressed in both aboveground and underground organs in sweet potato (Fig. 5), suggesting potential functions in both autotrophic and heterotrophic tissues. The localization of *IbG6PPT1* to the chloroplast membrane (Fig. 3) implied that it may function in transporting G6P from the cytosol into plastids.

To better elucidate the function of the *IbG6PPT1* gene in starch accumulation, we cloned *IbG6PPT1*, heterologously expressed it in Arabidopsis, and then measured starch accumulation in the resulting transgenic plants. *IbG6PPT1* expression increased starch accumulation in Arabidopsis leaves, seeds, and root tips, suggesting that it promotes starch biosynthesis. Lipid bodies and protein are the major storage compounds in mature Arabidopsis seeds, each accounting for up to 40% of the dry weight (Andriotis et al., 2010), whereas starch are lower. The weight of 1000 seed we observed in *IbG6PPT1*-expressing plant is greater than control plant, indicating that *IbG6PPT1* may also promote storage matter accumulation in Arabidopsis seeds.

*IbG6PPT1* promotes carbohydrate accumulation in Arabidopsis storage tissues

Sucrose is a major end product of photosynthesis and the primary sugar transported within plants (Winter and Huber, 2000). In heterotrophic tissues, sucrose imported from photosynthetic tissues is converted to G6P, and some G6P can be transported into the plastid through GPTs for starch and/or fatty acid biosynthesis. Another portion of the G6P is metabolized in the cytosol to phosphoenolpyruvate (PEP), which is essential for the biosynthesis of lipids and other storage substances (Lee et al., 2017). In *IbG6PPT1*-expressing Arabidopsis, the starch content in the leaves increased significantly, while the soluble sugar content was reduced, compared to that in control plants (Fig. 7). Thus, heterologous expression of *IbG6PPT1* promoted starch accumulation and sucrose metabolism, probably due to the high expression of GPT, which would be expected to increase the level of G6P imported into the chloroplast or amyloplast for starch synthesis. *IbG6PPT1* expression in Arabidopsis would promote G6P transport into plastids for storage and thus contribute to the observed carbohydrate accumulation in transgenic seeds compared with controls. Therefore, it is also likely that *IbG6PPT1* plays a critical role in metabolic distribution and carbohydrate accumulation in storage tissues. This also indirectly implies that *IbG6PPT1* plays an important role in the accumulation of substances in the storage roots of sweet potatoes.

*IbG6PPT1* enhances transport activity from sink to source
In plants heterologously expressing *IbG6PPT1*, the starch contents in leaves, seeds, and root tips increased (Fig. 7), indicating that *IbG6PPT1* might have high activity in Glc6P transport and play an important role in starch biosynthesis. Sweet potato is one of the top starch-rich root crops globally (Wang et al., 2019). To ensure starch storage in the storage root, any increases in photosynthetic activity depend on corresponding development of carbon sink capacity. Thus, Glc6P transporters such as *IbG6PPT1* should exhibit a high capacity for photosynthate transport. Indeed, heterologous expression of *IbG6PPT1* increased the starch content in the leaves, seeds, and root tips in Arabidopsis. However, expression did not affect the growth and development of transgenic plants, suggesting the potential of *IbG6PPT1* to promote starch accumulation in other crops.

**Conclusion**

The changes in starch and sugar content of transgenic Arabidopsis plants showed that *IbG6PPT1* might play an important role in starch and sucrose metabolism. It is probably an important gene controlling the starch properties in sweet potato. These findings will help to elucidate the genetic basis and regulatory mechanisms underlying starch properties in sweet potato.

**Materials And Methods**

*Plant material and growth conditions*

The sweet potato varieties Xushu22 (XS22) was cultivated at temperatures of between 22 and 28°C in the experimental base of the Sweet Potato and Potato Research Institute, Southwest University, Chongqing, China. Leaf, stem, petiole, and root were sampled at 95 days after transplanting (DAP) and quickly frozen in liquid nitrogen then stored at -80°C until use for RNA extraction. All *Arabidopsis thaliana* and *Nicotiana benthamiana* plants were grown in a 22°C and 28°C climate chamber (16h light/8 dark) in Longping experimental building, Southwest University, Chongqing, China.

*Cloning sweet potato GPT genes and sequence analysis*

To obtain the full-length mRNA sequences of target sweet potato GPT-encoding genes, the cDNAs of GPT genes were cloned using the SMARTer™ RACE cDNA amplification kit (Invitrogen, USA). RNA was extracted from the leaf, stem, petiole, and storage root of sweet potato variety Xushu 22 (XS22), and residual DNA was digested using the RNAprep Pure Plant Kit with DNase I (DP432, Tiangen Biotech, Beijing, China) according to the manufacturer's instructions. A 5-mg, equally proportioned (w/w) mixture of the above RNAs was used for first-strand cDNA synthesis. The gene-specific primers 83665-5-1 (5'-GGTGTGTGCAACTGCAACTGGGAAGAGGG-3') and 83665-5-2 (5'-GCCTCACAGCCGAGATCATCATTAT-3') were designed based on *IbG6PPT* transcripts (Yang et al., 2017; Zhang et al., 2017) and used to amplify the 5' end of the GPT genes. The primers 83665-3-1 (5'-GGTGGTTGCTCGCTTGCTGCTCTTACCG-3') and 83665-3-2 (5'-TCAGTATTGGAAACACCATGAAGCGT-3') were used to amplify the 5' and 3' ends of GPT genes. PCR products were cloned into the pMD19-T vector (TaKaRa) and sequenced. Based on the
obtained 5'- and 3'-end sequences of GPT genes, the full-length cDNA sequence was amplified using a 5' primer (5'-ACACAACACACTGTACTTGTTTC-3') and 3' primer (5'-CAAAATTTGAAAGAGGTTCCCAAACG-3') that were designed to match the 5'- and 3'-end sequences. PCR products were recombined into the Gateway entry vector pENTR-D-TOPO (Thermo Fisher, Massachusetts, USA) for sequencing. Open reading frame (ORF) and sequence alignment was performed with Geneious Prime.

Transmembrane transport peptides were predicted by the TMPred tool in ExPASy (http://www.ch.embnet.org/software/TMPRED_form.html/, (Krogh et al., 2001) and TMHMM (http://www.cbs.dtu.dk/services/TMHMM/) (Hofmann and Stoffel, 1993) using default parameters. Signal peptides were predicted by the SignalP 4.1 (http://www.cbs.dtu.dk/services/SignalP/) server using default parameters (Petersen et al., 2011). Conserved domains in the encoded proteins were analyzed with InterPro (http://www.ebi.ac.uk/interpro/) (Southan, 2000). The three-dimensional structure of IbG6PPT1 was predicted using Swiss-Model (http://www.swissmodel.expasy.org), and the constructed model was examined and visualized with Chimera 1.2 (https://www.cgl.ucsf.edu/chimera/). Multiple sequence alignment results from ClustalW were used for phylogenetic tree construction by the neighbor-joining method with MEGAX (Kumar et al., 2018). Tree reliability was measured by bootstrap analysis with 1000 replicates.

Expression pattern assay

RNA (1 μg) was extracted from the leaf, stem, petiole, and root of 95 DAP in XS22 and was reverse transcribed in a 20 μL volume by the PrimeScript RT Master Mix (TaKaRa) according to the manufacturer's instructions. The expression pattern of GPT genes was detected using primers and RT-qPCR methods as previously described (Zhang et al., 2017). Fold changes of the GPT transcripts were calculated according to the 2^−ΔΔCt method with three samples.

Subcellular localization

The full coding sequence (CDS) of IbG6PPT1 was cloned into pCAMBIA1300, and a GFP tag was fused to the C terminus of the gene. This construct was transformed into Agrobacterium tumefaciens strain GV3101 and transiently expressed in Nicotiana benthamiana using syringe agroinfiltration (Huy et al., 2016). GFP fluorescence was observed using a Zeiss LSM780 confocal laser scanning microscope (Li et al., 2019). Signals were detected using excitation/emission wavelengths for GFP (488 nm/495–535 nm) and the chloroplast marker (633 nm/660–720 nm).

Heterologous expression of IbG6PPT1 in Arabidopsis

The full CDS of IbG6PPT1 was recombined from the Gateway entry vector pENTR-D-TOPO (see the cloning and sequence analysis method above) into the destination vector pEarleyGate101 (Earley et al., 2006), yielding the construct p35S::IbG6PPT1-YFP, which has an N-terminal YFP tag. The construct p35S::IbG6PPT1-YFP was transformed into Arabidopsis using the Agrobacterium tumefaciens–mediated floral dip method (Desfeux et al., 2000).
Positive transgenic lines were identified by PCR detection of YFP using the primers YFP-Fwd (5’-TGGTCGAGCTGGACGCGACGTAAAC-3’) and YFP-Rev (5’-TTCTCGTTGGGTTTTTGCTCAGGGC-3’) and by detection of the bar gene in the construct using the primers FBar (5’-TGGGCAGCCCGATGACAGCGACCAC-3’) and RBar (5’-ACCGAGCGCCAGGAACCGCAGGAGT-3’). \textit{IbG6PPT1} expression in the transgenic Arabidopsis plants was detected using the RT-qPCR method described in the expression pattern assay section. YFP expression was detected by western blotting using an anti-GFP antibody (Welsch et al., 2019). Thousand seed weight (g) was determined for 1000 seeds from each sample with three replicates.

\textit{Starch and sugar measurement}

The starch and soluble sugar contents of leaves and seeds in transgenic and control Arabidopsis plants were determined using a previously published method (Kunz et al., 2010). Briefly, leaves and roots of 3 weeks seedlings were stained with an iodine solution (2% KI+1% I\textsubscript{2}) and examined under a light microscope (Nikon, Japan), and images were captured using NIS-Elements BR 4.30.00 software as previously described (Kunz et al., 2010).

\textbf{Declarations}

\textbf{Author Contributions}

Wu Z planned and performed the experiments and prepared the manuscript. Wang Z performed the experiments. Zhang K edited the manuscript and gave advice regarding the work. All authors have read and approved the manuscript.

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\textbf{Conflicts of Interest}

The authors declare no conflict of interest. The funders had no role in the design of the study; in the collection, analyses, or interpretation of data; in the writing of the manuscript, or in the decision to publish the results.

\textbf{Availability of data and materials}

The datasets supporting the conclusions of this article are included within the article and its additional files. About proteins database could download from NCBI by their accession number.

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Alignment of IbG6PPT1, IbG6PPT1-2, and Ipomoea genus GPT1 proteins. ItG6PPT1, Ipomoea triloba GPT1 (XP_031105621.1); InG6PPT1, Ipomoea nil G6PPT1 (XP_019193616.1). The amino acids underlined in red form transmembrane helixes based on prediction using TMHMM; black and grey highlighting indicate amino acid differences between the species.
Figure 2

Phylogenetic analysis of GPT proteins. The phylogenetic tree was constructed using the Neighbor-Joining method implemented in MEGA-X software. The numbers on the branches are bootstrap values (based on 1,000 repeats).
Figure 3

IbG6PPT1 localizes to the chloroplast membrane. (a) Nicotiana benthamiana leaf viewed in bright field, (b) IbG6PPT1-YFP viewed in the green channel, (c) the chloroplast marker viewed in the red channel, and (d) the merged images illustrating the localization of IbG6PPT1 to the chloroplast membrane.

Figure 4

Predicted three-dimensional structure models of IbG6PPT1. Two IbG6PPT1 proteins (shown in yellow and blue) form a dimer.
Figure 5

Expression of IbG6PPT1 in the petiole, storage root, stem, and leaf of the sweet potato variety Xushu 22, as determined by qRT-PCR. Each value is the mean±SD of at least three independent measurements.

Figure 6

Heterologous expression of IbG6PPT1 in Arabidopsis. (a) qRT-PCR detection of IbG6PPT1 expression; each value is the mean±SD of at least three independent measurements. (b) Western blot detection of
IbG6PPT1 expression. (c) Phenotype of 4 weeks Arabidopsis plants heterologously expressing IbG6PPT1. Col-0, control plants; OV-14, OV-30, OV-76, and OV-57, four transgenic lines.

Figure 7

Heterologous expression of IbG6PPT1 in Arabidopsis alters the starch and soluble sugar content of the plants. (a) Soluble sugar content and (b) Starch content of the leaves of 6 weeks Arabidopsis plants. (c) Iodine-stained starch in the leaves of 3 weeks seedlings. (d) Weight in 1000 seed, (e) soluble sugar content and (f) starch content in the seed in the transgenic and control plants. (g) and (h), Iodine-stained starch in
the root tips of control and transgenic plants, respectively. Col-0, control plants; OV-14, OV-30, OV-76, and OV-57, four transgenic lines. Each value is the mean±SD of at least three independent measurements.

**Supplementary Files**

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- TableS1.xlsx