Abstract The presence of high amounts of seed storage proteins (SSPs) improves the overall quality of soybean seeds. However, these SSPs pose a major limitation due to their high abundance in soybean seeds. Although various technical advancements including mass-spectrometry and bioinformatics resources were reported, only limited information has been derived to date on soybean seeds at proteome level. Here, we applied a tandem mass tags (TMT)-based quantitative proteomic analysis to identify the significantly modulated proteins in the seeds of two soybean cultivars showing varying protein contents. This approach led to the identification of 5,678 proteins of which 13 and 1,133 proteins showed significant changes in Daewon (low-protein content cultivar) and Saedanbaek (high-protein content cultivar) respectively. Functional annotation revealed that proteins with increased abundance in Saedanbaek were mainly associated with the amino acid and protein metabolism involved in protein synthesis, folding, targeting, and degradation. Taken together, the results presented here provide a pipeline for soybean seed proteome analysis and contribute a better understanding of proteomic changes that may lead to alteration in the protein contents in soybean seeds.

Keywords Glycine max, LC-MS/MS, Low-abundance proteins, Protamine sulfate precipitation, Seed storage proteins, Tandem mass tags

Introduction

Soybean seeds (Glycine max Merr. L.) are one of the most important global sources of vegetative proteins and oils for humans and livestock global. Moreover, other nutrients present in soybean seeds including isoflavones, phytate, soy-saponin, and others, exhibit health-promoting effects in treating metabolic disorders, cardiovascular diseases, and cancer (Badole et al. 2015). Multiple efforts have been put to elucidate the differential profile of transcriptome (Lambirth et al. 2015; Schmidt et al. 2011), metabolome (Schmidt et al. 2011), and proteome (Min et al. 2015; Pandurangan et al. 2012; Xu et al. 2015) using soybean seeds showing different protein contents.

The classical workflow of soybean seed proteomics including two-dimensional proteomic analysis (2-DGE) allowed the identification of a few hundreds of significantly modulated proteins due to the presence of high abundant proteins (HAPs) which account for 75% of total proteins in soybean seeds (Min et al. 2019). Recently, this limitation have been overcome by of the development of a variety of methods for the pre-fractionation of total soybean proteins (Gupta et al. 2016) using protamine sulfate (PS) (Kim et al. 2013, 2015), calcium (Krishnan et al. 2009), and polyethylene glycol (PEG) (Kim et al. 2001). Moreover, advancements in the
liquid chromatography-tandem mass spectrometry (LC-MS/MS)-
based methodologies and analytical software for downstream
data processing such as MaxQuant (Tyanova et al. 2016a),
Perseus (Tyanova et al. 2016b), Proteome discoverer (Thermo
Fisher Scientific, Waltham, MA, USA), and Skyline (Maclean
et al. 2010), have led to the improvement of sensitivity,
reliability, and coverage in proteome analysis.

Although the methodological developments have con-
tributed to the identification of thousands of proteins per
MS run with complete proteome coverage (Boersema et al.
2015; Niu et al. 2018), soybean seed proteomics is still
poorly conducted because of the presence of SSPs and
utilization of majorly 2-DGE based proteomics approaches
(Min et al. 2019). Previously, Kim’s group carried out high-
throughput proteome analysis using soybean seeds by shot-gun
proteomic approaches including label-free (Min et al. 2017)
and tandem mass tag (TMT) labeling quantitative analysis
(Min et al. 2020b). In particular, a TMT-based quantitative
analysis of filling stages of soybean seeds identified 5,918
proteins, the highest number of proteins reported to date
in soybean seeds (Min et al. 2020b). Moreover, the utilization
of the PS precipitation method with shot-gun proteome
pipeline, especially the TMT labeling approach, were carried
out and comparison of the total, PS-supernatant (PS-S), and
PS-pellet (PS-P) proteins, revealed enrichment of various
low-abundance proteins (LAPs) related to diverse seed me-
tabolism (Min et al. 2019).

Here, we are reporting a comparative seed proteome
profiling of two soybean cultivars differing into protein
content. Altogether, this study resulted in the identification
of 1,146 differentially modulated proteins (13 and 1,133
protein showed different abundance profiles), providing a
list of potential protein candidates using two soybean seed
cultivars differing in protein and oil contents.

Materials and Methods

Plant materials

Soybean seeds (Daewon, and Saedanbaek) were sown in
the experimental fields of the National Institute of Crop
Science (NICS), Rural Development Administration (RDA),
in Miryang, South Korea, in June. The soil was supplemented
with a standard RDA N-P-K fertilizer (N-P-K=3-3-3.3 kg/10
acre). Seeds were harvested in October 2018 (average tem-
perature, 23.5±3.5°C; average day length, 12 hours 17 min)
(Min et al. 2016).

Protein extraction, protein digestion, and TMT labeling

Total proteins from two different cultivars of soybean seeds
were isolated using the PS precipitation method with tri-
chloroacetic acid (TCA)/acetone precipitation method (Gupta
et al. 2015; Kim et al. 2015). Briefly, for PS precipitation
method, one gram of each seed powder was homogenized
with 10 mL of ice-cold Tris-Mg/NP-40 extraction buffer (0.5
M Tris-HCl, pH 8.3, 2% (v/v) NP-40, 20 mM MgCl2) and
centrifuged at 15,922 g for 10 min at 4°C. The clear ho-
memate was incubated on ice for 30 min with 0.15% (w/v)
PS stock solution. The extract was centrifuged at 15,922
 g for 10 min at 4°C to divide the PS-S and PS-P fractions,
respectively, as described previously (Kim et al. 2015).
Finally, the PS-S fraction was dissolved in 80% acetone
containing 0.07% β-mercaptoethanol and stored -20°C until
further analysis. Trypsin digestion by filter-aided sample prep-
paration (FASP), TMT labeling and peptide pre-fractionation
by basic pH reverse phase chromatography were carried
out as described previously (Gupta et al. 2020; Kim et al.
2018; Min et al. 2019a; Wiśniewski et al. 2009). A total
of 12 peptide fractions were collected, lyophilized in a
vacuum centrifuge and stored at -80°C until further LC-
MS/MS analysis.

LC-MS/MS analysis

Obtained peptides were dissolved in solvent-A (water/
Acetonitrile (ACN), 98:2 v/v; 0.1% formic acid) and separated
by reversed-phase chromatography using a UHPLC Dionex
UltiMate ® 3000 (Thermo Fisher Scientific, USA) instrument
(Pajarillo et al. 2015). For trapping the sample, the UHPLC
was equipped with Acclaim PepMap 100 trap column (100 μm
× 2 cm, nanoViper C18, 5 μm, 100 Å) and subsequently
washed with 98% solvent A for 6 min at a flow rate of
6 μL/min. The sample was continuously separated on an
Acclaim PepMap 100 capillary column (75 μm × 15 cm,
nanoViper C18, 3 μm, 100 Å) at a flow rate of 400
nL/min. The LC analytical gradient was run at 2% to 35%
solvent B (100% ACN and 0.1% formic acid) over 90
min, then 35% to 95% over 10 minutes, followed by 90%
solvent B for 5 minutes, and finally 5% solvent B for 15
minutes. Liquid chromatography-tandem mass spectrometry
(LC-MS/MS) was coupled with an electrospray ionization source
to the quadrupole-based mass spectrometer QExactive™
Orbitrap High-Resolution Mass Spectrometer (Thermo Fisher
Scientific, MA, Waltham, USA). The resulting peptides were
electro-sprayed through a coated silica emitted tip (Scientific
Instrument Service, NJ, Amwell Township, USA) at an ion
spray voltage of 2000 eV. The MS spectra were acquired at a resolution of 70,000 (200 m/z) in a mass range of 350-1650 m/z. The automatic gain control (AGC) target value was 3 x 10^6 and the isolation window for MS/MS was 1.2 m/z. Eluted samples were used for MS/MS events (resolution of 35,000), measured in a data-dependent mode for the 15 most abundant peaks (Top15 method), in the high mass accuracy Orbitrap after ion activation/dissociation with Higher Energy C-trap Dissociation (HCD) at 32 collision energy in a 100-1650 m/z mass range (Pajarillo et al. 2015). The AGC target value for MS/MS was 2 x 10^5. The maximum ion injection time for the survey scan and MS/MS scan was 30 ms and 120 ms, respectively.

Data analysis by MaxQuant, Perseus, and R software

The acquired raw data were analyzed with the MaxQuant software (version 1.5.3.30) as described previously (Tyanova et al. 2016a; Gupta et al. 2018; Min et al. 2020b). All three technical replicates were cross-referenced against the Uniprot Glycine max database (75,674 entries, UP000008827, http://www.uniprot.org). TMT data processing was performed using default precursor mass tolerances set by the Andromeda search engine, which is set to 20 ppm for the first search and 4.5 ppm for the main search. Reporter mass tolerance has to set the minimum as 0.003 Da. The product mass tolerance was set to 0.5 Da and a maximum of two missed tryptic cleavage were allowed. Carbamidomethylation of cysteine residues and acetylation of lysine residues and oxidation of methionine residues were specified as fixed and variable modifications respectively. A reverse nonsense version of the original database was generated and used to determine the FDR which was set to 1% for peptide identifications. Statistical analysis was carried out using Perseus software (ver. 1.5.8.5) and R software as described previous report (Min et al. 2020a, 2020b). For removing the batch effect within TMT-6plex, data normalization was carried out using an internal reference scaling method as described previously (Plubell et al. 2017; Gupta et al. 2019) Missing values imputation was carried out from a normal distribution (width: 0.3, downshift: 1.8) using Perseus software (Tyanova et al. 2016b). Multiple Sample test controlled by the Benjamini-Hochberg FDR threshold of 0.05, was applied to identify the significant differences in the protein abundance (> 1.5-fold change). The functional classification and pathway analysis were carried out using AgriGO v2.0 (Tian et al. 2017) web-based software for GO enrichment analysis, KEGG pathway analysis by DAVID proteome annotation web-based software (Jiao et al. 2012), and MapMan software (version 3.6.0 RC1), respectively.

Results

Quantitative proteomic analysis using soybean seeds

To investigate the differential modulation of soybean seed proteome in high- and low-protein containing cultivars, seed proteins were isolated from Daewon and Saedanbaek and subjected to protamine sulfate precipitation method for depletion of major seed storage proteins (SSPs) (Kim et al. 2015). SSPs depleted fraction, referred as PS-S fraction, from two different cultivars (marked by DS; Daewon PS-S fraction and SS; Saedanbaek PS-S fraction, respectively) were sequentially subjected to trypsin digestion by filter-aided sample preparation (FASP) method and TMT-6plex labeling in the same manner as reported previously (Min et al. 2020a, 2020b) (Fig. 1A). Sequentially, pre-fractionation by basic-pH reversed-phase (BPRP) using in-house developed stage-tip was carried out to decrease the complexity of multiplex labeling sample mixtures (Han et al. 2014). This approach led to the identification of 51,278 peptides and 22,483 unique peptides matching to 5,678 protein groups from three technical replicates of TMT labeling sample sets (Fig. 1A). Particularly, TMT labeling combined with pre-fractionation approach showed improvements of the resolution and identification of protein as observed by 4,892 (84.3%) while a previous label-free study (Min et al. 2017)
using PS-S fraction of soybean seed protein identified a comparatively lower number of protein (247 unique proteins, 0.4%) than present study (Fig. 1B).

Data normalization and statistical analysis

For normalization and removal of batch effects within TMT data sets, we applied an internal reference scaling (IRS) method to 4,610 proteins showing more than 70% valid intensity values (Fig. 2A). As per the normalization steps, TMT data sets were normalized at the peptide spectrum match (PSM) level into the MaxQuant software (Yu et al. 2020). Sequentially, PSM-level normalized reporter ion intensities of each TMT data set were applied to the further IRS method for normalization (Plubell et al. 2017). These multiple-step normalization procedures showed the correction of batch effects that occurred by TMT-6plex reagents (Fig. 3A). IRS normalization of the data showed an improvement of the median coefficient of variation (CV) values of each sample from 19.63% to 6.06% (Fig. 3B). Besides, Pearson correlation coefficients showed a high degree of correlation among different replicates of each sample with an average $R^2$ value of 0.996 (Fig. 2B). Of these 4,610 proteins, the sequential application of fold change (FC) calculation and Student’s $t$-test controlled by a Benjamini-Hochberg FDR were applied to identify the statistically significant proteins between Daewon and Saedanbaek seeds (FDR < 0.05, FC > 1.5) (Fig. 2C). This resulting in the identification of 1,146 differential proteins, of these 1,133 and 13 proteins showed increased and decreased abundances in cluster_1 and 2 respectively (Table S1 and Fig. 4A). The PCA plot analysis revealed that PS-S proteins in Daewon vs Saedanbaek cultivars were separated at the PC1 accounting for a maximum 95.9% variation (Fig. 2D).
MapMan analysis of 1,146 differential proteins showed up- and down-regulation of various proteins in the metabolism and cell function overview categories. Proteins with increased abundance in Saedanbaek, involved in cluster_1, were mainly related to the CHO metabolism (9.3%), photosynthesis (9.3%), secondary metabolism (8.5%), lipid metabolism (15.1%), and amino acid metabolism (13.2%) (Table S2). In the cell function overview category, majority of these proteins were found to be associated with protein degradation (11.9%), stress-related protein (10.8%), signaling (9.1%), transport (8.2%), RNA regulation (7.6%), protein targeting (7.2%), protein synthesis (5.9%) (Table S2). Particularly, in the case of the protein degradation category, various types of protease including subtilases, serine, cysteine, and aspartate protease, among others showed increased abundance Saedanbaek (Fig. 4B). Furthermore, 32 proteins related to protein synthesis including various isofrom of ribosomal proteins, initiation, and elongation factors also showed increased abundance in Saedanbaek (Fig. 4B). In addition to protein synthesis, the increased abundance of 4, 9, 39, and 20 proteins related to amino acid activation, protein folding, protein targeting, and post-translational modifications respectively were observed in Saedanbaek cultivar (Fig. 4B).

GO enrichment analysis of identified proteins showed an increased abundance of the proteins associated with the major metabolic pathway. In particular, proteins involved in cluster_2 showed increased abundance of proteins associated with protein metabolic process (GO:0019538), protein localization (GO:0008104), protein transport (GO:0015031), protein folding (GO: 0006457), and protein catabolic process (GO:0030163), among others in biological process categories (Table S3). In order to get further functional insights of proteins involved in cluster_2, KEGG pathway analysis was carried out using DAVID functional annotation web-based
software (Jiao et al. 2012). KEGG pathway analysis showed that proteins with increased abundance in Saedanbaek were majorly associated with various metabolic pathways including biosynthesis of secondary metabolites, biosynthesis of amino acids, carbon metabolism, and protein processing in the endoplasmic reticulum (Table S4).

Discussion

Recently, the next-generation proteomics approaches including label-free and isotope labeling-based quantitative analysis have been showing significant improvements in protein quantification and thus identification of differential proteins (Boersema et al. 2015; Min et al. 2019). However, soybean seed proteomics is still elusive due to several limitations including a narrow range of detection, low reproducibility, and difficulty to detect LAPs due to the presence of high abundant proteins (HAPs) (Gygi et al. 2000; Thompson et al. 2003). Therefore, a number of HAPs depletion methods have been developed specifically for the enrichment of LAPs from soybean seeds using protamine sulfate (Kim et al. 2015), calcium (Krishnan et al. 2009), and PEG (Kim et al. 2001). Our previous study showed a broad application of PS for the enrichment of LAPs from different plant samples including seeds and leaves of rice, soybean, pea, and peanut (Kim et al. 2015). Moreover, a previous report revealed that LAPs related to various major metabolism in filling and matured stages of soybean seeds were successfully enriched and identified in PS-S fraction using TMT-based quantitative analysis (Min et al. 2020b). Therefore, here we utilized the PS precipitation method in combination with TMT-based quantification to identify the differential proteins from the seeds of Daewon and Saedanbaek differing in total protein contents (Gupta et al. 2020; Min et al., 2020a, 2020b). This approach led to the identification of 1,146 significantly modulated proteins (FDR < 0.05, FC > 1.5) by the comparison between Daewon and Saedanbaek cultivars. Moreover, further
functional classification of the increased abundance proteins, particularly in Saedanbaek cultivar showed accumulation of various LAPs associated with major seed metabolic pathways including photosynthesis, major/minor CHO metabolism, amino acid metabolism, lipid metabolism, and secondary metabolism, among others.

For the accumulation of storage compounds such as proteins and lipids, an enormous amount of energy is required for which the diffusion of oxygen in plant tissues is prerequisite for the energy production in mitochondria (Krishnan and Coe, 2001; Galili et al. 2014). Therefore, energy production through photosynthetic activity is required and critical for the accumulation of reserved metabolites during seed desiccation (Fait et al. 2006). Here, we identified 22 proteins including psbP, psb28 subunits, ATP synthase, plastocyanin, ferredoxin, NADH-ubiquinone oxidoreductase chain 1, Ribulose bisphosphate carboxylase, Glyceraldehyde-3-phosphate dehydrogenase, and among others associated with photosynthesis showing increased abundance in Saedanbaek which is similar to that reported previously (Table S1) (Min et al. 2020b).

Besides, enrichment of LAPs led to the identification of 24 and 39 proteins related to major/minor CHO and lipid metabolism which showed increased abundance in the Saedanbaek cultivar (Table S2). Out of these, six and ten proteins related to starch synthesis/degradation and lipid degradation, respectively, showed increased abundance along with an increased abundance of two raffinose synthase proteins in the Saedanbaek cultivar (Table S1 and S2).

Moreover, during seed maturation stages, 10 to 15% of lipids are converted to raffinose family oligosaccharides (RFOs) when the supply of exogenous resources from maternal plants are limited (Kambhampati et al. 2020). These RFOs are produced by carbon remobilization from lipid along with sucrose during the development of seeds (Kambhampati et al. 2020).

In addition, we observed the accumulation of 34 proteins mainly associated with amino acid metabolism including GABA, glutamate, aspartate, branched-chain amino acids, tryptophan, serine, glycine, cysteine, and histidine synthesis (Table S2). Furthermore, MapMan functional classification of metabolism overview revealed the increased abundance of 3 proteins (more than 1.5 and 2.0-FC increase) involved in nitrogen metabolism which have an important role in determining the total amount of storage proteins. The ammonia derived from nitrogen uptake by maternal vegetative tissues is the primary source for supply the nitrogen predominantly as amino acid such as glutamine and asparagine to the seeds (Ohyama et al. 2017). In addition, amino acids participate in the synthesis of storage proteins and thereby contributing the carbon remobilization through proteolysis activity during the late seed developmental stages (Galili et al. 2014; Kambhampati et al. 2020). Here, 64 proteases showed increased abundance in Saedanbaek that might be having a crucial role in the remobilization of endogenous nitrogenous products such as amino acid or proteins to storage proteins (Gallardo et al. 2006, 2007). Taken together, our results suggest a positive correlation of various metabolism-related proteins involved...
in major/minor CHO metabolism, photosynthesis, nitrogen, amino acid metabolism, and among others with a higher protein content of soybean seeds.

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