Subtractive libraries for prospecting differentially expressed genes in the soybean under water deficit

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Abstract

Soybean has a wide range of applications in the industry and, due to its crop potential, its improvement is widely desirable. During drought conditions, soybean crops suffer significant losses in productivity. Therefore, understanding the responses of the soybean under this stress is an effective way of targeting crop improvement techniques. In this study, we employed the Suppressive Subtractive Hybridization (SSH) technique to investigate differentially expressed genes under water deficit conditions. Embrapa 48 and BR 16 soybean lines, known as drought-tolerant and -sensitive, respectively, were grown hydroponically and subjected to different short-term periods of stress by withholding the nutrient solution. Using this approach, we have identified genes expressed during the early response to water deficit in roots and leaves. These genes were compared among the lines to assess probable differences in the plant transcriptomes. In general, similar biochemical processes were predominant in both cultivars; however, there were more considerable differences between roots and leaves of Embrapa 48. Moreover, we present here a fast, clean and straightforward method to obtain drought-stressed root tissues and a large enriched collection of transcripts expressed by soybean plants under water deficit that can be useful for further studies towards the understanding of plant responses to stress.

Key words: gene expression, drought, transcriptome.

Introduction

Soybean is one of the most important crops in the world, and Brazil is its second largest producer. Due to its use in animal feed, vegetable oil and biodiesel, there is a sustained demand for soybeans. Nevertheless, abiotic stresses, such as water deficit, cold, wind, heat or waterlogging, reduce productivity and present a major challenge in the quest for sustainable soybean production. In order to meet the ever-increasing demand, studies to improve productivity are very important. Crop yield is highly affected by water deficit stress, particularly when it occurs during flowering and early pod expansion (Pedersen et al., 2005). However, soybean plants can acclimate to such conditions by triggering protective mechanisms that enable these plants to survive subsequent events of drought (Bray, 2004). While studying soybean plants exposed to stress at different developmental stages, Kron et al. (2008) observed that plant responses were more uniform in the V4 stage (Fehr et al., 1971). The genetic background of the cultivars BR 16 (drought sensitive) and Embrapa 48 (drought tolerant) under drought conditions was studied by Oya et al. (2004), and they found that at the vegetative stage, tolerant cultivars present higher growth rate and a larger leaf area compared to sensitive plants. However, to better understand plant physiology under stress, productivity analyses are hardly enough to distinguish significant tolerance differences among cultivars. The data indicate how cultivars behave under stress and, therefore, suggest which ones are suitable for further biochemical and molecular studies.

A stress response is initiated when plants recognize the stress at the cellular level, it triggers a complex signal transduction network which is modulated by molecular events. Gene products involved in water-deficit responses can be classified into two groups: functional or regulatory genes. Functional genes include the following: proteins, osmolytes, chaperones, heat-shock proteins (HSPs), water-channel proteins (multifunctional proteins involved in transport facilitation of solutes and water in leaves and roots), and other compounds probably conferring direct tol-
erance to abiotic stresses. Under water stress conditions, a chemical imbalance in the cell leads to production of reactive oxygen species (ROS), which are damaging to lipids and proteins (Carvalho, 2008; Molina et al., 2008). Enzymes involved in detoxification metabolism are found to protect against the stress in the tolerant common bean cultivar (*Phaseolus acutifolius*) (Türkan et al., 2005). In drought-stressed tissues, the water movement within and between cells can be facilitated by aquaporins (Coccozza et al., 2010). The second group of genes involved in the response to water deficit includes molecules involved in further regulation of signal transduction and stress-responsive gene expression. These molecules are regulatory proteins, represented by various transcription factors such as DREB1, ABA-responsive element, nitrogen assimilation control protein, ethylene-response factors, kinases, phosphatases, enzymes involved in phospholipid metabolism, and other signaling molecules, such as calmodulin-binding protein (Shinozaki and Yamaguchi-Shinozaki, 2007). In the peanut, a predicted drought tolerant legume, a set of functional and regulatory genes were detected in enriched subtractive libraries probably acting in a gradual process of drought acclimation (Govind et al., 2009).

As discussed, the ability to tolerate water deficit is a complex trait controlled by many genes (Molina et al., 2008; Ergen and Budak, 2009). Collections of cDNAs expressed under stressed conditions have been a useful resource to study gene expression and regulation. The cDNA libraries were associated to normalization steps and PCR amplification to obtain just the transcripts presented in a target situation (Diatchenko et al., 1996). A suppressive subtractive library approach makes it possible to search for genes expressed in treated samples but not in the control samples, if the forward subtraction is done. It can be used to assess changes in metabolism and in quantitative gene expression studies, such as those based on arrays. The method has been employed successfully in different research areas, such as the monitoring of citrus responses to fungus infection (González-Candelas et al., 2010) and in fish infected by iridovirus (Xu et al., 2010). Using the subtracted library of pigeonpea, Priyanka et al. (2010) isolated a gene encoding a hybrid-rich-proline protein, which affords tolerance to water deficit and other abiotic stresses such as salt, heat, cold and abscisic acid treatment. In addition, cDNA subtractive libraries have also allowed the identification of genes involved in the flower development of orange plants (Zhang et al., 2008) and of genes involved in the *Varroa destructor* mite tolerance to honey bees (Zhang et al., 2010).

In recent years, drought has resulted in severe losses in soybean productivity worldwide. In order to expand previous knowledge about the water stress-sensitive BR 16 and water stress-tolerant Embrapa 48 soybean cultivars, the present work aimed to study early responses in gene expression of leaves and roots under water stress.

### Material and Methods

#### Plant growth and water stress treatment

Seeds of the soybean cultivars Embrapa 48 and BR 16, classified as tolerant and sensitive to water deficit, respectively, were germinated on filter paper during four days in a growth chamber at 25 ± 1 °C of temperature and 100% relative humidity. Seedlings were placed in 36 L boxes containing 50% Hoagland’s solution (Hoagland and Arnon, 1950), which was continuously aerated and replaced on a weekly basis. These boxes were then transferred to a greenhouse under a natural photoperiod of approximately 12/12 h light/dark cycle, temperature of 30 ± 5 °C and 60 ± 10% relative humidity (RH), where the plants were allowed to grow until the V4 stage (Fehr et al., 1971). The experimental plan was a randomized complete block 2x7 factorial design with three repetitions. The factors were two cultivars (BR 16 and Embrapa 48) and seven times of water deficit (0, 25, 50, 75, 100, 125 and 150 min). The stress was applied by removing the plants out of the hydroponic solution and leaving them in boxes without nutrient solution for up to 150 min under ambient-air exposure. For each time of water deficit, roots from 10 plants were collected, pooled and frozen in liquid nitrogen before storage at -80 °C. The same procedure was performed for the leaf samples.

#### Physiological parameters measurements

Photosynthetic rate (*A*), stomatal conductance (*g*), intercellular CO2 concentration (*c*), transpiration rate (*E*) and leaf temperature (*T*) were evaluated using a LI-6400 Portable Photosynthesis System (LiCor, Inc.). Measurements were taken on the middle leaflet of the youngest soybean leaf, fully expanded, under photon flux density of 1,000 μmol m⁻² s⁻¹. The water use efficiency (WUE) of plants was determined through the ratio between photosynthetic (*A*) and transpiration (*E*) rate. Data was submitted to ANOVA analysis through the programs Sanest and SAS (Statistical Analysis System version 8.0), and the treatment means were compared by the Tukey test (*p* < 0.05).

#### cDNA subtracted libraries synthesis

Total RNA was extracted from roots and leaves using Trizol reagent (Invitrogen) according to the manufacturer’s instructions. Samples from each biological replicate were extracted separately, and equimolar amounts of total RNA were pooled before the mRNA purification. The FastTrack MAG mRNA Isolation (Invitrogen) was applied to isolate the mRNA Poly A⁺ molecules by using oligo-dT conjugated magnetic beads. The mRNA samples were pooled as illustrated in Figure 1 to perform the cDNA reverse transcription. Libraries of roots and leaves from Embrapa 48 and BR 16 plants were constructed based on the method of Diatchenko *et al.* (1996) with the PCR Select cDNA Subtraction (Clontech) Kit. First strand cDNA was synthesized...
from 2 μg of mRNA Poly A+ pooled samples (tester and driver). After the second strand synthesis, cDNAs (tester and driver) were digested with the restriction enzyme Rsa I to produce blunt end fragments. Once digested, the cDNAs (only tester samples) were divided into two samples and each sample was ligated with two different adapters (adapter 1 and 2R, respectively). As our aim was to detect transcripts expressed only in water-deficient stressed plants, the sample T0 (control plant) was used as the driver in the hybridization step. The cDNA driver (with no adaptor) was added to tester samples (1 and 2R) and they were submitted to the first hybridization step to eliminate sequences common in both samples or present in the control (driver). For the second hybridization, tester samples 1 and 2R from the first hybridization were mixed together, and the denatured cDNA driver was added again to enrich the expressed sequences in the tester. The PCR amplification was performed using the double-strand cDNAs with different adapters as template. Sequences were exponentially amplified due to the specificity of primers to adapters. The second PCR was carried out using the nested primers to enrich the differentially expressed sequences. The PCR products were directly applied to next generation sequencing.

Data analysis

The PCR products were sequenced using the Sequencing by Synthesis Technology (Illumina). The six libraries from BR 16 cultivar (three from roots and three from leaves) were sequenced through 36 bp (base pairs) single reads in a Genome Analyzer Ille machine. The Embrapa 48 libraries were sequenced in a Genome Analyzer Ilz using single reads of 76 bp. Libraries tagged with a barcode produced thousands of reads from each library. After base-calling, reads were mapped on a reference genome (Schmutz et al., 2010) using the SOAP (Li et al., 2008), and non-mapped reads were assembled into contigs through the Edena program following the default parameters (Hernandez et al., 2008).

Contig sequences were submitted to the non-redundant protein database NCBI through Blast X (Altschul et al., 1997), to search for similarity to known proteins. In addition, sequences were analyzed by the software AutoFact (Koski et al., 2005), which is an automated annotation tool that assigns biological information for a given sequence by comparing different databases. We used the UniRef90 e UniRef100, KEGG (Kanehisa and Goto, 2000), Pfam (Finn et al., 2010), Smart (Schultz et al., 1998) databases. In order to establish the GO (Gene Ontology) terms (Ashburner et al., 2000) we employed the Blast2Go program to classify the sequences according to the molecular function and biological process described to the respective protein (Götz et al., 2008, Carbon et al., 2009).

Results

Plants grown hydroponically were exposed to a short-term water deficit and the physiological parameters \( A, g_s, c_i, T_l, T_{roo} \) and WUE are presented in Figure 2. We employed the SubtractiveSuppressive Hybridization (SSH) technique to obtain an enriched collection of ESTs expressed early in response to water deficit stress. cDNA libraries were constructed to detect genes up-regulated when compared to control plants. After sequencing and base calling, reads were mapped on the soybean genome and assembled into contigs. Approximately 65% of the total reads produced for each library was mapped on the genome. Contigs assembled were searched in the NCBI database in order to access the information about proteins. The data from each sequencing run are shown in Table 1. After the AutoFact analysis, the unknown/hypothetical genes decreased significantly. All genes identified by using the subtractive libraries are stored in the Soybean Database.

Biochemical roles of the proteins were determined by using the Biological Process level 3 from Gene Ontology (Figure 3). Approximately 71% of the sequences in BR 16 and 61% of Embrapa 48 showed an associated putative biological or molecular function. Consequently, the number of no match and unknown genes was larger in Embrapa 48 than in BR 16 plants. In BR 16, we observed 121 and 452 different roles for the genes expressed in the roots and leaves, respectively. In Embrapa 48, there were 251 and 128 biological processes, respectively. Categories employed in functional classification include the following: Biological regulation (gene expression regulation, protein modification regulation); Signaling (kinases, phosphatases, secondary messengers); Response to stress (water deficit, osmotic and salt stress, heat, cold); Response to stimulus (chemical stimulus, endogenous stimulus); Response to biotic stimulus (pathogen, fungus, insect); Cell organization (cytoskeleton components, cell wall organization, cellular component assembly); Cell cycle (actin filament-based process, cell division); Senescence (aging, cell death); Other biological process (cellular component organization, cell recognition, developmental process); Biosynthetic process (carbohydrate, fatty acid, hormone, small and macromolecules, pigment, nucleotide); Catabolic process (pro-
tein, fatty acid, cell wall, polysaccharide); Nitrogen compound metabolism (glutamate and glutamine biosynthetic process, cellular nitrogen compound metabolic process); Oxidation reduction (electron transport, lipid oxidation) and Secondary metabolism (terpenoid, phytochelatin, phytoalexin and phenylpropanoid metabolic process). Each category was normalized by the total number of sequences observed in the respective library, which enables comparison of the results during the time course of the water deficit (Figure 3).

Applying the Venn-Master Diagram, we also searched for genes expressed in more than one library as described in Figure 4. BR 16 root tissue genes were categorized as involved in the electron transport process (R1 and R2 libraries) or as encoding proteins related to the stress response class (R2-R3 library) (Figure 4A). In BR 16 leaves (L1-L2 libraries), we also detected genes involved in signaling reactions or involved in the transport of metabolites and electrons (oxidation reduction process) (Figure 4B). In addition, the genes in the L2-L3 libraries were also related to stress response (Figure 4B). A general view of L1-L2-L3

Figure 2 - Physiological parameters measured on drought-stressed plants. Photosynthetic rate (μmol CO₂ m⁻² s⁻¹), stomatal conductance (mol H₂O m⁻² s⁻¹), intercellular CO₂ concentration (μmol CO₂ mol⁻¹), transpiration rate (mmol H₂O m⁻² s⁻¹), and leaf temperature (°C) were measured with a LiCor Portable Photosynthesis System. The water use efficiency (mmol C/mol H₂O) was determined based on transpiration and photosynthetic rate. From each stress treatment, two plants were evaluated in each block. Results represent the mean value of all plants analyzed. The (±) standard error bars are shown.
libraries confirmed that genes functionally related to transport and stress responses were also expressed in leaves over the water deficit period (Figure 4B). It is important to emphasize that transcripts observed in R1-R2, for example, were different from the transcripts found in R1-R2-R3, although similar biological processes could be detected in both libraries. Interestingly, in the Embrapa 48 cultivar, transcription factors and genes involved in signal transduction were the most abundant transcripts in both R1 and R2 root tissues (Figure 4C). Genes encoding transcription factors were also predominant in the R2-R3 libraries in addition to the ones related to transport process (electrons and molecules). The same pattern was found in R1-R2-R3 libraries (Figure 4C). Processes related to protein metabolism (assembly, folding and modification) were very active in L2-L3 libraries as well as genes involved in transport process, response to stress and signaling. Stress response genes stand for 22% of the genes identified in L1-L2-L3 libraries (Figure 4D).

We performed a comparison between genes expressed in roots and leaves in either cultivar using the Venn-Master Diagram tool and looking for the biological processes observed in each group. When the same contig or glyma was detected in more than one library it was counted as one transcript. Then, considering only the non-repeated genes from each group (tissue or cultivar), we were able to...
observe genes expressed in root and leaves as well as genes expressed in Embrapa 48 and BR 16 plants (Figure 5A). To assess biological processes in roots and leaves of both cultivars, we highlighted the more abundant putative proteins and cellular components in each water-deficit stressed tissue (Figure 5B). According to our data, transcripts expressed by BR 16 plants in roots and leaves showed that the most frequent biological processes were similar in both tissues (Figure 5B). However, in Embrapa 48, the process of response to stress was predominant among transcripts from leaves and less common in roots (Figure 5B).

Among genes expressed by plants in response to water deficit, we observed that transcripts classified in the response to stress category presented similarity to proteins with responsiveness to biotic as well as to abiotic stress. These transcripts represent proteins with antimicrobial activity or proteins involved in defense and immune responses. Genes related to the response to virus, pathogenic bacterium, nematodes and fungi were also detected. Under abiotic stress, we identified transcripts in response to osmotic and oxidative stress, desiccation, heat, cold, water deprivation, hyperosmotic salinity and high light intensity. To assess which transcripts are expressed in response to other adverse conditions, the genes placed in the response to stress category (n = 458) were compared to transcripts from plants infected by *Phakopsora pachyrhizi*, *Uromyces appendiculatus*, virus and treated with *Bradirhizobium japonicum* (Figure 6).

### Discussion

Plant water deficit responses and tolerance are complex biological processes that need to be analyzed at system-level using genomics and physiological approaches (Harb et al., 2010). Studies aiming to detect differences among plants in response to stress using contrasting cultivars have been performed in some legumes (Türkan et al., 2005). Empirc observations during growing seasons in the field are the first step to explore plant variability based on its yield under water deficit conditions. Oya et al. (2004) examined the physiological characteristics of the Embrapa 48 and BR 16 cultivars and determined that they were moderately tolerant and highly sensitive to drought, respectively. Here, we studied these cultivars at the molecular level, and our data demonstrated that the water deficit applied was effective as it triggered changes in the plant physiological behavior (Figure 2). As the stress increased in the plants, photosynthesis of both cultivars was affected and was strongly inhibited after 100 min. This inhibition was related to a drastic decrease in the stomatal conductance (gs) and transpiration rate (E) of both cultivars. We also observed an increase in the leaf temperature (from 100 min) and internal concentration of CO2 (from 125 min). The water deficit did not produce significant physiological differences between Embrapa 48 and BR 16 cultivars. This was expected as the short-term procedure to induce stress used here is usually not enough to produce effects on plant morphological and physiological characteristics needed to distinguish the two cultivars. The experimental treatment described here aimed to study the effects of the stress on the early plant responses at a molecular level. Additionally, the hydroponic system allowed the obtainment of a clean root sample, as demonstrated by Martins et al. (2008).

As described in the literature, subtractive libraries have been employed in a wide range of studies such as gene expression studies (Kojima et al., 2009; González-Candelas et al., 2010; Jain and Chattopadhyay, 2010; Soria-Guerra et al., 2010). Most of these studies were based on the construction of a physical cDNA collection by cloning fragments into a vector and on the traditional big dye sequencing technology. However, in vivo cloning can be considered a laborious and expensive procedure and also has technical limitations. In order to overcome these limitations, both cultivar cDNAs were sequenced using two different chemistries based on Sequencing by Synthesis Technology (Illumina). As shown in Table 1, we identified many genes being regulated under water deficit, more than is often detected through subtractive libraries. Larger sets of genes were also detected in the characterization of stress-responsive genes to heat in wheat (Chauhan et al., 2011) and by studying salt-induced genes of grapevine (Daldoul et al., 2010). Both suppressive subtractive libraries were done through in vivo cloning and by the capillary sequencing method. Interestingly, while investigating invasive species, Prentis et al. (2010) found 12442 contigs in...
the transcriptome by combining subtractive libraries and massively parallel sequencing (pyrosequencing). One of the advantages of the next generation sequencing (NGS) over the traditional Sanger methodology is a more precise measurement of the transcript levels and isoforms, in addition to other revolutionary benefits (Marioni et al., 2009; Wang et al., 2009). The advantages and limitations found in the NGS and capillary sequencing have led to comparative studies of both methods (Cheung et al., 2008; Hert et al., 2008; Tedersoo et al., 2010). Lower costs and quickness have made the NGS a powerful tool for functional genomics studies (Shendure and Ji, 2008; Harismendy et al., 2009).

Under water deficit, plants can up- or down-regulate expression of responsive genes. Different biochemical roles have been suggested for these responsive genes, and their expression patterns make an intricate network that can be related to either water deficit acclimation or the cel-
lular damage response (Shinozaki and Yamaguchi-Shinozaki, 2007). The expressed genes have been shown to belong to many classes of genes (Figure 3). However, the largest class of genes expressed under stress remains unknown, partly because there is no similarity in databases available to date, and partly due to the gene functions are not understood yet (Reddy et al., 2008; Nelson and Shoemaker, 2006). Many genes expressed under water deficit conditions are present in the contrasting cultivars Embrapa 48 and BR 16 (Figure 5A). Among others, common processes affected by the stress are those related to transport, protein folding and assembling as well as protein degradation and ubiquitination. Some proteins are synthesized in requirement of defense pathways triggered by the stress whereas other proteins are degraded because its aggregation or denaturation events are predominant in water stress conditions. The transport category involves the translocation of ions, lipids, sugar and others molecules. Also included in this category are the genes for aquaporins, which are proteins that might be directly involved in water deficit acclimation by facilitating water movement across the membranes (Zhao et al., 2008). We have identified several transcripts encoding the tonoplast and plasma membrane intrinsic proteins that were more frequently found in roots rather than in leaves, especially in the BR 16 plants (Figure 5B). Roles played by aquaporins in plants can be regulated in different complex ways, but the expression of aquaporins throughout the plant would most likely be regulated by water potential gradient (Zhao et al., 2008). However, there are differences between the mRNA changes and its respective protein level. The changes expressed here by soybean plants in response to water stress are only at the transcriptional level. BR 16 plants did not show an expressive quantitative difference in the biological processes affected by water deficit between roots and leaves (Figure 5B). Nonetheless, Embrapa 48 did show a significant contrast between both tissues (Figure 5B) regarding the genes included in the response to stress class (water deficit, osmotic and salt stress, heat and cold).

In abiotic stress responses, genes are induced to protect the plants against a disturbance in cell homeostasis. Under water stress, plants can become susceptible to other abiotic or biotic stresses because their normal metabolism is directed towards the acclimation to the first stress. Several studies have tried to make a connection between plant responses to water deficit and heat (Sakuma et al., 2006; Montero-Barrientos et al., 2010), cold (Li et al., 2010), or reactive oxygen species enhanced due to drought (Kocsy et al., 2005; Khanna-Chopra and Selote, 2007). In the response to stress category, we observed that various transcripts were similarly involved in several abiotic stresses such as osmotic stress, cold, heat and salinity as well as those involved in response to biotic stimulus (bacterium, fungus, nematode, virus) (Figure 6). From subtracted libraries of pigeonpea, Priyanka et al. (2010) identified a hybrid-proline-rich encoding gene whose multiple stress-responsive characteristics have been demonstrated under PEG, mannitol, salt and heat stress, cold and ABA-treatment. Another gene from pigeonpea, a cyclophilin gene, presented a stress-responsive nature in Arabidopsis thaliana (Sekhar et al., 2010). A bZIP transcription factor gene found in soybean plants was also associated with multiple stress responses, being its expression induced by ABA-treatment, drought, high salt and low temperature (Gao et al. 2011).

Table 1 - Summary of the total reads sequenced and number of sequences assembled and mapped in each library. (A) BR 16 was sequenced by using a 36 bp single read in Genome Analyzer lle whereas the run of (B) Embrapa 48 cultivar was carried out through a 76 bp single read in the sequencer Genome Analyzer llz.

<table>
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<th>Library</th>
<th>Reads</th>
<th>Contigs</th>
<th>No hits</th>
<th>Matched sequences</th>
<th>Known sequences</th>
<th>Unknown/hypothetical</th>
<th>GO annotation (%)</th>
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In conclusion, a hydroponic system was applied successfully to grow soybean plants and to produce clean roots. This method allowed for a large collection of cDNAs expressed by soybean plants under water deficit and their respective biochemical processes. Although the analyses presented here represent a first step in gene expression investigation, the results indicate changes which are triggered in roots and leaves of tolerant and sensitive soybean plants.

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