Research Article

New Insights in the Sugarcane Transcriptome Responding to Drought Stress as Revealed by Supersage

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Received 31 October 2011; Accepted 2 December 2011

Academic Editor: Luigi Cattivelli

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In the scope of the present work, four SuperSAGE libraries have been generated, using bulked root tissues from four drought-tolerant accessions as compared with four bulked sensitive genotypes, aiming to generate a panel of differentially expressed stress-responsive genes. Both groups were submitted to 24 hours of water deficit stress. The SuperSAGE libraries produced 8,787,315 tags (26 bp) that, after exclusion of singlets, allowed the identification of 205,975 unitags. Most relevant BlastN matches comprised 567,420 tags, regarding 75,404 unitags with 164,860 different ESTs. To optimize the annotation efficiency, the Gene Ontology (GO) categorization was carried out for 186,191 ESTs (BlastN against Uniprot-SwissProt), permitting the categorization of 118,208 ESTs (63.5%). In an attempt to elect a group of the best tags to be validated by RTqPCR, the GO categorization of the tag-related ESTs allowed the in silico identification of 213 upregulated unitags responding basically to abiotic stresses, from which 145 presented no hits after BlastN analysis, probably concerning new genes still uncovered in previous studies. The present report analyzes the sugarcane transcriptome under drought stress, using a combination of high-throughput transcriptome profiling by SuperSAGE with the Solexa sequencing technology, allowing the identification of potential target genes during the stress response.

1. Introduction

Sugarcane (Saccharum spp.) is an outstanding crop throughout the tropical regions of the world [1]. It represents an important food and bioenergy source, being cultivated in many tropical and subtropical countries [2], and covering more than 23 million hectares worldwide, with a production of 1.6 billion metric tons of crushable stems [3]. This crop is responsible for almost two thirds of the global sugar production [1]. Brazil, the world’s largest sugarcane producer, processed and generated in 2008 about 31 million tons of sugar [4]. In contrast to most plants, sugarcane stores sucrose—rather than polymeric compounds such as starch, proteins, or lipids—as the primary carbon and energy reserve [1]. Hence, sugarcane byproducts have received greater attention, due to their multiple uses, with the ethanol generation being highlighted, as an important renewable biofuel source [5]. Moreover, the bagasse of sugarcane has been largely used for energy cogeneration at distilleries, production of animal feed and also for paper production [6]. Nevertheless, similarly to other meaningful agronomical crops, sugarcane cultivation faces considerable losses due to inappropriate or unfavorable edaphoclimatic conditions.

Abiotic stresses are among the main causes of major crops worldwide productivity losses [7], causing negative impacts on crop adaptation and productivity. In this scenario, drought figures as the most significant stress and is considered an extremely important factor when it comes of losses in the productivity of sugarcane [8]. Several plant biotechnology programs have been initiated aiming
to increase drought stress tolerance in crop plants using genetic engineering and traditional breeding [9]. Although breeding activities have provided significant progress for the understanding of the physiological and molecular responses of plants to water deficit, there is still a large gap between yields in optimal and stress conditions [10]. For this purpose, case-sensitive methods are demanded, not only to discover new genes associated to those stress conditions, but also to effectively detect differentially expressed genes on a drought tolerant variety. The identification and expression profile of such responsive genes may be helpful to unravel the basic mechanism of stress tolerance [11]. In this sense, previous works uncovered genes associated to important roles in stress perception, signal transduction, and transcriptional regulatory networks in cellular responses, useful for the improvement of stress tolerance in plants by gene transfer [12, 13].

Molecular approaches concerning drought and salinity performance in sugarcane were carried out using techniques based on molecular hybridization such as Suppression Subtractive Hybridization (SSH) [11] and micro-/macroarrays [14]. In general, the main limitations of these methods are their low sensibility and specificity [15]. Among the methodologies for transcriptomic analysis, the SuperSAGE [16] approach represents one of the most recent and informative methods [17], especially with its association to the high-performance sequencing platforms (pyrosequencer [454 Roche], Solexa (Illumina), and SOLID (Applied Biosystems)). SuperSAGE regards an evolution of the traditional Serial Analysis of Gene Expression [18] generating longer (26bp) tags and thus allowing most reliable annotation analysis. Since, it is an open architecture method (i.e., allowing the discovery of new genes), it presents the potential to provide a global and quantitative gene expression analysis, based on the study of the entire transcriptome produced in a given time and tissue, under a given stimulus. Additionally, SuperSAGE permits a simultaneous analysis of two interacting eukaryotic organisms, full-length cDNAs amplification using tags as primers, potential use of tags via RNA interference (RNAi) in gene function studies, identification of antisense and rare transcripts, and identification of transcripts with alternative splicing [19]. Besides, this method has been recently associated to the next generation sequencing technologies, allowing a less expensive and faster covering of the analyzed transcriptomes, permitting a deep insight of the modulated responses under different physiological conditions. The association of SuperSAGE with the rapid advances in high throughput sequencing opened the possibility of performing genome-wide transcriptome studies in non model organisms. Additionally, this technique has been successfully applied in plant species such as rice [16], banana [20], chickpea [21, 22], chili pepper [23], tobacco [24], and tropical crops (cowpea, soybean, sugarcane; [25]). In the present work, we profit from the high resolution power of SuperSAGE coupled to the Illumina sequencing to characterize the transcriptome of drought-stressed sugarcane roots after 24 hours of submission to this stress, aiming to elect a best group of tags to be validated by RTqPCR.

2. Methodology

2.1. Identification of Drought-Tolerant and Sensitive Sugarcane Accessions. For the selection of the drought-tolerant and sensitive accessions used in the present evaluation, a previous assay was carried out in order to identify contrasting genotypes for these features. For this purpose, 20 commercial sugarcane varieties (CTC 1 to 15, SP83-2847, SP83-5073, CT94-3116, SP90-1638, and SP90-3414) from CTC (Sugarcane Technology Center, Piracicaba, Brazil) were evaluated. Among these, the four above-mentioned varieties were used as a standard for the interpretation of results, including two varieties (SP83 and SP83-2847-5073) identified as drought-tolerant and other two (SP90 and SP90-1638-3414) indicated as drought-sensitive based on field empirical observations performed by specialized technicians during several years in sugarcane commercial fields.

For this assay, mini-cuttings from the 20 varieties above were planted in 50 L pods containing inert substrate (Plantmax) in order to allow the slow increase of water deficit by removing irrigation. Tests were performed with six-months-old plants under greenhouse conditions and the treatments included plant permanently irrigated (without stress), suppression of irrigation for three days (72 hours stress), suppression of irrigation for 10 days (240 hours stress), and suppression of irrigation for 20 days (480 hours stress). Physiological measurements applied in all treatments included chlorophyll content using an SPAD-507.B Chlorophyll Meter; analysis of chlorophyll fluorescence ratio between variable and maximum chlorophyll-a (Fv/Fm); estimation of chlorophyll content with a fluorometer; determining the relative water content. For the parameters of chlorophyll-a fluorescence and chlorophyll content, three measurements were taken from three plants from each treatment. Data analysis was performed by comparing the percentage change considering the parameters mentioned above. After this assay, four drought-tolerant and four sensitive accessions could be selected according to the parameters used, revealing a gradient of water stress tolerance among the varieties analyzed. Considering the classification of the standard varieties identified previously as drought-tolerant (SP83-2847 and SP83-5073) and drought-sensitive (SP90-1638 e SP90-3414) and also considering the measurements taken after stress under glasshouse conditions (these results will be presented in a separated manuscript) four varieties were considered as drought-tolerant (CTC15, CTC6, SP83-2847 and SP83-5073) and other four as drought-sensitive (CTC9, CTC13, SP90-3414, and SP90-1638).

2.2. Drought Stress Application and the SuperSAGE Libraries. Plants of each selected accession were grown under glasshouse conditions in 40 L pods, in randomized experimental design (comprising six repetitions) under daily irrigation until the age of three months. After that, part of the material was submitted to drought by interruption of irrigation during 24 hours. Roots of both, stressed and nonstressed plants, were collected and frozen in liquid N2, being maintained in a deep freezer until total RNA extraction using
Trizol (Invitrogen). The extracted samples were quantified by spectrophotometry, digested with DNase and purified with the aid of the RNasy Mini kit (Qiagen). The samples were quantified again by spectrophotometry, allowing the composition of the bulks using equimolar amounts of poli-A + messenger RNA, for all treatments. Four libraries have been generated: TD (bulk of four tolerant accessions under stress); TC (bulk of four tolerant genotypes without stress, as sensitive negative control); SD (bulk of four sensitive materials under stress); SC (bulk of nonstressed sensitive accessions, as sensitive negative control). The procedures for SuperSAGE library generation followed Matsumura et al. [26], including the attachment of library-specific adaptors carried out by GenXPro GmbH (Frankfurt am Main, Germany) allowing the identification of library-specific reads after SOLEXA sequencing.

2.3. Statistical Analysis and Tag-Gene Annotation. The 26-bp tags were extracted from each library. Singlets (reads appearing only once) were excluded from the present evaluation. Statistical tests were applied to the remaining tags (Audic Test, Claverie; P ≤ 0.05) with aid of the DiscoverySpace 4.1 software [27] regarding the four contrasting treatments [T (TD versus TC); S (SD versus SC); D (TD versus SD); C (TC versus SC)]. The tests allowed the identification of the total number of expressed unitags (or tag species) for each situation and contrast, as well as the differentially expressed tags, including up- (UR) and downregulated (DR) tags.

The tag-gene annotation was performed by independent evaluations via BlastN [28] against different EST databases: NCBI: (i) dbEST including only Saccharum ESTs; (ii) Gene Index (including Arabidopsis thaliana, AtGI 15.0, and Poaceae species: S. officinarum, SOGI 3.0; Sorghum bicolor, SBGI 9.0; Zea mays, ZMGI 19.0; Panicum virgatum, PAVIGI 1.0; Oryza sativa, OsGI 18.0; Triticum aestivum, TAGI 12.0; Hordeum vulgare, HVG 11.0; Festuca arundinacea, FAGI 3.0; Secale cereale, RYEGL 4.0); and (iii) KEGG (including A. thaliana and Fabaceae ESTs). Valid BlastN alignments were considered when the following parameters were observed: score from 42 to 52; integrity of the CAGT sequence at the 5′ end; plus/plus alignments. Inferences about the modulation of a specific tag (Fold Change; FC) were carried out considering the ratio of the observed frequencies of a given library in relation to the other.

2.4. Gene Ontology of SuperSAGE Hits. Matching ESTs to the analyzed tags were categorized via GO using the software Blast2GO [29] after BlastX alignment against the Uniprot-SwissProt protein database (ε-value ≤ e-10). ESTs related to the GO subcategories concerning abiotic stress response (to water deprivation, GO: 0009414; to heat/cold, GO: 0009408/GO: 0009409; to osmotic stress, GO: 0006970, to oxidative stress, GO: 0006979, to asbcsic acid stimulus, GO: 0009737; to jasmonic acid stimulus, GO: 0009753) were identified, as well as UR tags related to these classes. Sets of UR tags considering the different contrasting situations (T, S, D, and C) were annotated, generating Venn diagrams, aiming the visualization of specific or shared tags considering the different treatments.

3. Results and Discussion

3.1. Qualitative and Quantitative Analysis of the SuperSAGE Libraries. The four SuperSAGE libraries produced 8,787,315 tags, from which 1,862,064 (21.2%) regarded singlets (tags sequenced only once), and were excluded from this evaluation. The most representative libraries considering the number of tags were TC (drought-tolerant control; 2,516,454 tags) and SD (drought-sensitive under stress; 2,133,587 tags), while the less representative were TD (drought-tolerant under stress; 750,226 tags) and SC (drought-sensitive control; 762,492 tags). The coverage of the transcriptome by the tags was estimated considering the total number of tags per genotype (3,266,680 for the tolerant bulk and 2,896,079 for the sensitive bulk) in relation to the number of expected transcripts per cell. The total number of averaged transcripts was estimated to range from 100,000 [30] to 500,000 [31] per cell in higher plants. Considering the high value (500,000), the coverage provided by the tags in relation to the sugarcane transcriptome was 6.5 times higher for the tolerant bulk and 5.8 for the sensitive bulk, that is, the number of expected single copy transcripts per cell should be represented by their tags in the absolute frequencies of around six in each library. Taking the less represented libraries (TD and SC) in account, the coverage of the transcriptome regarded 1.5 times higher for both, tolerant and sensitive bulks. Considering this value, we established the n < 2 frequency as cutoff threshold, allowing the exclusion of singlet tags. Coverage of this magnitude allowed a comprehensive evaluation of a given transcriptome, also including rare transcripts expressed during the response to the evaluated stress.

Taking all valid tags (n ≥ 2) into account, a total of 205,975 unitags remained for evaluation. In a recent approach, Yamaguchi et al. [32] observed similar amounts (∼190,000 unitags) in the roots of Solanum torvum under heavy metal stress (CdCl2; 0.1 μM). The high number of unitags, here observed, shows the diversity of transcripts (and expressed genes), possibly also reflecting the allopolyploid nature of sugarcane, since tags diverging in a single nucleotide were considered to be distinct unitags. It has been speculated that, in some cases, unitags could be the result of artifacts generated by the amplification process during library construction [33] or incomplete digestion of the synthesized cDNA by the NlaIII enzyme [34], and also by PCR amplifications associated to innate features of the sequencing technology [32]. In order to minimize error sources, some precautions were taken during library development in this study, including double digestion of the total RNA extracted with DNase, double digestion with the NlaIII enzyme, and exclusion of singlet tags. An additional way to minimize potential errors would be the exclusion of unitags related to other similar sister-tags, grouping them to other most frequent, so called mother-tags. On the other hand, this procedure would eliminate transcripts bearing important single nucleotide polymorphisms (SNPs).

Still, another possibility would be to establish a minimum frequency (n) of a given tag to be considered valid. In the present work, only canonical tags were accepted, with
A more stringent value ($n > 2$) was adopted by Yamaguchi et al. [32], to reduce the number of unitags per library (from 300,000 to 450,000) for each 33 thousand tags, in an attempt to reach the number of expected genes for model species as rice (32,000 genes) and A. thaliana (26,000 genes). However, this procedure impairs the identification of rare and alternative transcripts that possibly play important roles in the cell metabolism.

Statistical analysis considering $P \leq 0.05$ (Audic-Claverie test) among libraries permitted the identification of differentially expressed tags including up- (UR) or downregulated (DR) tags for the four contrasting situations (T; S; D; C), as shown in Table 1.

### 3.2. Primary Annotation of SuperSAGE Tags

Relevant BlastN alignments comprised 567,420 tags (75,404 unitags with 164,860 different ESTs). Details about the results obtained after alignment to different databases are not itemized here, since this is not the aim of the present evaluation. Despite that the choice of the databases and the adopted criteria allowed the following: (a) the identification of ESTs related to most tags, preferentially concerning sequences from sugarcane or taxonomic the related species sequences; (b) annotation of a considerable number of tags considering a minimum alignment of 21 bp (similar to a LongSAGE tag); (c) identification of tags with perfect alignments (100% identity) or with a maximum of a single mismatch among tag and EST, important for future development of primers; (d) avoidance of plus/minus alignments, minimizing false NATs (natural antisense transcripts).

The strategy of considering the alignments without the election of a best hit allowed the maximization of annotation chances, since no alignment was disregarded in the acceptable score range. Thus, alignments with annotated ESTs could be more informative than similar alignments with a slightly superior score in relation to nonannotated ESTs. Moreover, tags aligned with distinct ESTs could be analyzed, minimizing the chance of a wrong choice that could compromise the validation of the expression results, especially considering that they are used as targets for RTqPCR primer design. In this context, seeking the maximization of the annotation procedures, the use of the Gene Index database for tag identity annotation was carried out trying to circumvent at least two limitations, when compared with the partial dbEST bank additionally used: (a) no need of clusterization concerning ESTs deposited at dbEST, since the Gene Index project provides tentative clusters (TC); (b) best functional annotation, with the Uniref100 (Uniprot) bank as reference. Thus, in view of the posterior need of primer design for RTqPCR and data validation of SuperSAGE tags, alignments with tolerance of a maximum of a single mismatch (TSM) tag-hit represented up to one third (186,191 or 32.8%) of the data, indicating high identity among 26 bp tags and similar ESTs, since a minimum of 21 bp alignment (size of a LongSAGE tag) was considered relevant. Almost all valid alignments (471,672 or 83.12%) regarded Saccharum spp. (partial dbEST) and S. officinarum (Gene Index SOGI), as expected. TSM alignments restricted to these databases comprised 163,742 tags. Considering TSM alignments with sequences of the SOGI only, from 26,884 ESTs, 73.0% presented informative gene descriptions and/or their functions, allowing the identification of molecular targets and gene-feature association. Despite of the higher number of TSM matches concerning alignments with dbEST sequences (136,858), the EST annotation was not informative for most contemplated ESTs (97.0%). To overcome this deficiency, the Gene Ontology categorization proved to bring a valuable contribution.

### 3.3. Functional Categorization of SuperSAGE Tags

BlastX evaluations ($e$-value $\leq e^{-10}$) of the 186,191 ESTs (diverse databases and TSM alignments) against the peptide Uniprot-SwissProt bank allowed the characterization of 118,208 ESTs (63.5%) that presented at least one GO reference. From this categorization, the Biological Process (BP) subcategories in response to abiotic stress were considered more informative to evaluate the sugarcane response to drought conditions. The first interesting indicators were UR tags associated to EST in the BP subcategories responding to water deprivation (GO: 0009414), heat (GO: 0009408), cold (GO: 0009409), osmotic stress (GO: 0006970), oxidative stress (GO: 0006979), abscisic acid stimulus (GO: 0009737), and jasmonic acid stimulus (GO: 0009753). By the analysis of the UR tags observed in the above-mentioned subcategories (Table 2), it was possible to generate the Venn diagrams presented in Figure 1, where Figure 1(a) represents the UR tags evaluated in the contrasting situations T (TD versus TC; 20 tags) and D (TD versus SD; 25 tags), both important for future gene validation. The first case (T) related to tags from the tolerant bulk induced after water deficit when

**Table 1**: Total number of differentially expressed (DE; $P \leq 0.05$) up- or downregulated tags observed in different contrasting SuperSAGE root libraries from sugarcane under drought stress (24 hours without irrigation) as compared with negative control (irrigated materials).

| Contrasting | Upregulated | Downregulated | DE | Total |
|-------------|-------------|---------------|----|-------|
| T (TD × TC) | 12,179      | 12,482        | 24,661 | 152,049 |
| S (SD × SC) | 12,085      | 16,339        | 28,424 | 141,946 |
| D (TD × SD) | 15,591      | 12,269        | 27,860 | 148,657 |
| C (TC × SC) | 12,961      | 16,342        | 29,303 | 148,631 |

TD: bulk of tolerant genotypes under stress; TC: bulk of tolerant genotypes without stress (control); SD: bulk of sensitive genotypes under stress; SC: bulk of sensitive genotypes without stress (control).
Figure 1: Venn diagrams with numbers of differentially upregulated (UR) tags from sugarcane roots ($P \leq 0.05$) under drought stress, considering different comparisons between SuperSAGE libraries [contrasts: T (TD versus TC); S (SD versus SC); D (TD versus SD); C (TC versus SC)]. UR tags associated with gene ontology (GO) response to (a) water deprivation; (b) heat/cold; (c) osmotic stress; (d) oxidative stress; (e) abscisic acid stimulus; (f) jasmonic acid stimulus. Libraries: TD (drought tolerant bulk under stress); TC (tolerant bulk control); SD (drought sensitive bulk under stress); SC (sensitive bulk control).

compared with the bulk control; the second group refers to induced tags from both bulks submitted to drought stress (tolerant versus sensitive), with higher expression (UR) in the tolerant bulk. The first group exhibited 17 non annotated tags and only three identified genes (encoding 18S ribosomal RNA, membrane integral protein, and viviparous-14). The second group included 17 tags without annotation and other eight bearing descriptions (18S ribosomal RNA gene (two tags); ABA responsive element binding factor 2; Auxin-induced protein; DRF-like transcription factor DRFL2a; ERF/AP2 domain containing transcription factor; GST; RAPB protein) are discussed latter in this manuscript. Additionally, 11 tags are worth mentioning, since they were UR in both tolerant (T) and sensitive (S) comparisons after stress (Figure 1(a)), when compared to the expression of the respective controls. Despite of being not genotype-dependent, these tags may influence positively in the plant adaptation process under drought stress. Such results and other for similar subcategories are presented in Table 2. This table comprises the total number of UR tags induced in the tolerant bulk under stress, highlighting the exclusive (T comparison) or differentially expressed tags in comparison to the sensitive bulk (D comparison), bringing interesting candidates for validation via RTqPCR. Since the same tags may be involved in different stresses, the identified tags (exclusive in T and common in the comparisons T and D; Table 2) may not be exclusive of a given condition or response. Thus, the total number of UR tags (alone or in combination) in response to water deprivation (W), heat/cold (H), osmotic stress (Os), and oxidative stress (Ox) is presented in Figure 2(a). Likewise, the number of tags induced in response to osmotic and oxidative stress is presented in Figure 2(b), while the tags responsive to hormonal stimuli (abscisic and jasmonic acids) is shown in the Figure 2(c), and a Venn diagram showing all the categories is presented in Figure 2(d).
Table 2: Total number of sugarcane upregulated (UR) root tags observed on contrasting SuperSAGE libraries when associated with ESTs classified by Gene Ontology (GO) in the subcategories related to abiotic stress response.

| Response against GO categories | Exclusive UR tags T | Common UR tags after comparison T and D | T and S |
|--------------------------------|---------------------|----------------------------------------|--------|
| Water deprivation              | 0009414             | 20                                     | 25     | 11     |
| Heat and cold                  | 0009408; 0009409    | 19                                     | 54     | 18     |
| Osmotic stress                 | 0009408             | 32                                     | 57     | 19     |
| Oxidative stress               | 0006979             | 19                                     | 27     | 6      |
| Abscisic acid stimulus         | 0009737             | 24                                     | 37     | 12     |
| Jasmonic acid stimulus         | 0009753             | 5                                      | 19     | 6      |

EST: expressed sequence tag; contrast of libraries [T (TD versus TC); D (TD versus SD); S (SD versus SC)]; Libraries [TD: drought-tolerant bulk under stress; TC: tolerant bulk control; SD: drought-sensitive bulk under stress; SC: sensitive bulk control].

Figure 2: Venn diagrams with numbers of SuperSAGE tags overexpressed (P ≤ 0.05) in sugarcane roots under drought stress, considering different tag sets related to gene ontology (GO) subcategories associated in response to: W (water deprivation), H (heat/cold), Os (osmotic stress), Ox (oxidative stress), ABA (abscisic acid stimulus), Jas (jasmonic acid stimulus).

In relation to the 213 UR tags, including the exclusive ones from the T contrast and those presented in both T and D contrasts (Table 2), the gene-function annotation together with the GO descriptions were available for 68 of them, while 145 tags remained unknown candidates. The annotations of these 68 UR tags and respective GO subcategories, as well as the fold change (FC) for both most relevant contrasts (T and D), are listed in Table 3. Some of them will be further addressed.

3.3.1. Response to Hormone Stimulus. Response to hormonal stimulus, such as jasmonic (JA) and abscisic acid (ABA), together with other plant hormones, as salicylic acid (SA) and ethylene (ET), form a complex network that plays major roles in disease resistance and response to abiotic stresses, including drought [35, 36]. In our study, 21 potential hormone-responsive tags were identified (Table 3) and some of them are thereafter discussed.

(a) ZIM Motif Family Protein. According to the database of Arabidopsis transcription factors (DATF; http://datf.cbi.pku.edu.cn/index.php), this short motif is associated to a panel of plant transcription factors and JA signaling, which is among the most important defense-related signals in plants, acting under environmental stresses, such as UV radiation, osmotic shock, heat, and drought [37]. Examining a jasmonate-insensitive 3 (jai3-1) mutant gene, Chini et al. [38] identified a novel family of jasmonate-regulated nuclear targets of SCFCOI1, named jasmonate ZIM-domain (JAZ) proteins repressing JA signaling and targeted by the E3-ubiquitin ligase SCFCOI1 for proteasome degradation. The overexpression of this hormone activated a damping
Table 3: Upregulated SuperSAGE tags associated via Gene Ontology (GO) to abiotic stress, with fold change for T (FC_T = TD/TC) and D (FC_D = TD/SD) comparisons for tag frequencies in the sugarcane roots libraries, as well as the annotation of the best aligned EST.

| Tag       | GO     | FC_T | FC_D | Annotation                                                                 |
|-----------|--------|------|------|-----------------------------------------------------------------------------|
| SD159390  | AJ     | 5.6  | ns   | 50S ribosomal prot. L5, chloroplast                                         |
| SD191288  | AJ     | 2.9  | 2.3  | AP2 domain transcription factor EREBP                                       |
| SD122727  | AJ     | 2.2  | 2.7  | Bet v 1 allergen-like                                                       |
| SD75453   | AJ     | 2.8  | 2.8  | Chromatin-remodeling factor CHD3                                             |
| SD123546  | AJ     | 1.6  | ns   | Chromatin-remodeling factor CHD3                                             |
| SD9608    | AJ     | 2.4  | 3.3  | Initiator-binding prot.; ibp                                                |
| SD15756   | AJ     | 1.9  | 1.6  | OSK1; SNF1-related prot. Kinase                                             |
| SD108270  | AJ     | 6.4  | 2.6  | ZIM motif family prot.                                                      |
| SD258836  | AJ     | 3.1  | ns   | ZIM motif family prot.                                                      |
| SD108269  | AJ     | 2.9  | 2.1  | ZIM motif family prot.                                                      |
| SD133809  | AJ     | 2.8  | 2.6  | ZIM motif family prot.                                                      |
| SD196399  | AJ     | 3.2  | 3.2  | P18; Nucleoside diphosphate kinase l; NDK1                                  |
| SD169158  | AJ/Os  | 3.2  | ns   | Peptidyl-prolyl cis-trans isomerase                                          |
| SD252082  | AJ/WH  | 3.6  | 3.6  | Auxin-induced prot.                                                         |
| SD237930  | AJ/WH/Os | 2.4  | ns   | 18S ribosomal RNA gene                                                      |
| SD282917  | AJ/WH/Os | 1.5  | 1.6  | ABA responsive element binding factor 2                                    |
| SD237939  | AJ/WH/Os | 6.8  | 6.8  | Branched-chain-amino-acid aminotransfer.                                    |
| SD238059  | AJ/WH/Os | 2.4  | 2.4  | Branched-chain-amino-acid aminotransfer.                                    |
| SD140270  | AJ/WH/Os | 3.2  | ns   | viviparous-14                                                              |
| SD237936  | AJ/WH/Os/Ox | 2.8  | 2.8  | Ribosomal prot. L28e domain cont. prot.                                     |
| SD178862  | AJ/WH/Os/Ox | 2.8  | 2.8  | 18S ribosomal RNA gene                                                      |
| SD205616  | WH     | 1.3  | ns   | RAP2-like prot.                                                             |
| SD246714  | WH     | 2.8  | 2.8  | CoA-thioester hydrolase CHY1                                                |
| SD286424  | WH     | 1.4  | 2.6  | ERF/AP2 domain cont. transcription factor                                   |
| SD279457  | WH     | 2.8  | ns   | Mitochondrial uncoupling prot. 2                                            |
| SD107875  | WH     | 3.7  | 5.2  | Nucleic acid binding                                                        |
| SD191687  | WH     | 2.1  | 6.9  | RAPB prot.                                                                  |
| SD147607  | WH     | 5.6  | ns   | Salt tolerance prot.                                                        |
| SD109060  | WH     | 1.4  | 1.3  | Transposable element Mu1 sequence                                           |
| SD199146  | Os     | 1.1  | 1.1  | Alpha tubulin-4*                                                            |
| SD54073   | Os     | 2.3  | 6.8  | Calreticulin-like prot.                                                     |
| SD102228  | Os     | 6.8  | 6.8  | Endo-1,4-beta-glucanase Cel1                                                |
| SD80163   | Os     | 4.7  | ns   | Endo-1,4-beta-glucanase Cel1                                                |
| SD13344   | Os     | 1.9  | ns   | Eukaryotic translation if 2 alpha sub family                               |
| SD182876  | Os     | 4.4  | ns   | Phosphopantetheine adenyllyl transf. dephospho CoA kinase                   |
| SD129463  | Os     | 3.0  | ns   | Serine/threonine-prot. kinase SAPK1                                         |
| SD87319   | Os     | 2.4  | 1.6  | Serine/threonine-prot. kinase SAPK6                                         |
| SD270381  | Ox     | 6.5  | 4.5  | Allene oxide synthase                                                       |
| SD272257  | Ox     | 2.4  | ns   | Allene oxide synthase                                                       |
| SD63148   | Ox     | 2.0  | 2.5  | Allene oxide synthase                                                       |
| SD113907  | Ox     | 2.4  | 2.4  | Brassinosteroid biosynthesis-like prot.                                     |
| SD219102  | Ox     | 3.2  | ns   | Glutathione peroxidase                                                      |
| SD213044  | Ox     | 2.2  | 2.4  | Na+/H+ antiporter                                                           |
| SD54454   | Ox     | 2.0  | 3.1  | Nicotianamine aminotransferase A                                            |
Table 3: Continued.

| Tag          | GO    | FC_T | FC_D | Annotation                                      |
|--------------|-------|------|------|------------------------------------------------|
| SD125582     | Ox    | 3.8  | 1.9  | Nicotinate phosphoribosyltransferase-like       |
| SD122742     | Ox    | 3.2  | ns   | Nucleotide repair prot.                         |
| SD102844     | Ox    | 1.6  | ns   | Peroxidase precursor                            |
| SD17103      | Ox    | 6.4  | ns   | Tyrosine/nicotiamine aminotransf. family        |
| SD17107      | Ox    | 2.5  | ns   | Tyrosine/nicotiamine aminotransf. family        |
| SD17108      | Ox    | 1.8  | ns   | Tyrosine/nicotiamine aminotransf. family        |
| SD151691     | WH/Os | 1.8  | 1.6  | DRF-like transcription factor DRFL2a            |
| SD9805       | WH/Os | 25.0 | 25.0 | Glycine-rich RNA binding prot.                  |
| SD9802       | WH/Os | 14.7 | 14.7 | Glycine-rich RNA binding prot.                  |
| SD9806       | WH/Os | 13.1 | 13.1 | Glycine-rich RNA binding prot.                  |
| SD9767       | WH/Os | 2.8  | 2.8  | Glycine-rich RNA binding prot.                  |
| SD9803       | WH/Os | 2.4  | 2.4  | Glycine-rich RNA binding prot.                  |
| SD9800       | WH/Os | 2.4  | 2.4  | Glycine-rich RNA binding prot.                  |
| SD9801       | WH/Os | 1.1  | ns   | Glycine-rich RNA binding prot.                  |
| SD108120     | WH/Os | 6.0  | 6.0  | Glycine-rich RNA-binding prot.                  |
| SD108115     | WH/Os | 1.3  | 1.3  | Glycine-rich RNA-binding prot. 2; GRP2          |
| SD264077     | WH/Os | 3.6  | ns   | Membrane integral prot.                         |
| SD92627      | WH/Ox | 5.7  | 2.4  | Glutathione transferase III                     |
| SD243418     | WH/Ox | 4.0  | ns   | Serine hydroxymethyltransferase                 |
| SD179937     | WH/Ox | 3.8  | 2.5  | Serine hydroxymethyltransferase                 |
| SD21923      | WH/Ox | 1.3  | 1.7  | Whitely-induced gp91-phox                       |
| SD184083     | Os/Ox | 3.2  | 3.2  | Delta-1-pyrroline-5-carboxylate dehydrog.       |
| SD8088       | Os/Ox | 3.2  | ns   | MutT domain prot.-like                          |
| SD251703     | Os/Ox | 7.1  | ns   | P5cs; delta 1-pyrroline-5-carboxylate synth.    |

Libraries [TD: drought-tolerant bulk under stress; TC: tolerant bulk control; SD: drought-sensitive bulk under stress]; ns: fold change of tag not significant (P ≤ 0.05). WH: response to water deprivation and to heat/cold; Os: response to osmotic stress, Ox: response to oxidative stress; AJ: response to abscisic acid stimulus and to jasmonic acid stimulus.

mechanism concerning the JA signaling cascades after stress initiation [39]. In our evaluation, five UR tags were hormone related, with one candidate (SD108270) presenting expressive fold change in both contrasts (FC_T = 6.4 and FC_D = 2.6; Table 3).

(b) Chromatin-Remodeling Factor. CHD3 has been implicated in the repression of transcription [40]. Association of these proteins to drought-responsive genes was related during Arabidopsis seed germination process by regulating the ABA-dependent and gibberellic acid (GA) dependent responses, modulating the plant reaction to mild osmotic stresses and limiting the expression levels of transcription factors, preventing a maladapted growth arrest. In other words, it refines the pace of seed germination in response to ABA and maintains embryonic characters silent in response to GA [41]. Our results indicate a differential expression of CHD3 also in roots of adult sugarcane plants undergoing water deficit, with two UR tags (SD75453 and SD123546) with FC values of 1.6 (SD123546; FC_T) and 2.8 (SD75453; FC_T = FC_D; Table 3).

(c) AP2/EREBP. It is a large family of plant transcriptional regulators that plays key roles in the development and environmental stress response pathways. Transcription factors encoded by AP2/EREBP genes contain the highly conserved AP2/ERF DNA binding domain [42] constituting a plant supergene family [43] subdivided into five subfamilies according to the number of AP2/ERF motifs [44]. The AP2/EREBP subgroup induced by biotic and abiotic stresses was identified by Sharoni et al. [45]. Among the upregulated genes, 52 were induced in response to diverse abiotic stress, such as cold, drought, and salt. Lin et al. [46] working with a full-length cDNA OsEBP2 (ethylene-responsive-element binding protein2) in japonica rice leaves infected by blast fungus Magnaporthe grisea observed that OsEBP2 responded transiently to the treatments with methyl jasmonate (MeJA), ABA, and ethophen (ethylene generator). In our analysis, a UR tag was annotated as APETALA 2/ethylene response element binding protein (AP2/EREBP) showing expressive modulation (FC_T = 2.9 and FC_D = 2.3; Table 3). Additionally, one UR tag (SD286424; FC_T = 1.4 and FC_D = 2.6; Table 3) annotated as AP2/ERF domain containing transcription factor was associated to our WH group (response to water deprivation + response to heat/cold; Table 3), indicating an important candidate for validation, since the overexpression of an ERF transcription factor GmERF3 from soybean in tobacco plants raised the tolerance to salinity (up to 400 mM, NaCl) and drought [47] in transgenic plants.
3.3.2. Response to Water Deprivation, Oxidative and Osmotic Stress. In our analysis, 47 potential stress-responsive UR tags with acceptable annotation were identified (Table 3) and some of them deserve special mentioning when considering their GO categorization and the fold change data.

(a) Glycine-Rich RNA Binding Protein (GRP) Superfamily. This superfamily, characterized by the presence of a glycine-rich domain arranged in (Gly)n-X repeats, was recently reviewed by Mangeon et al. [48] that highlighted the diversity in structure, expression pattern, and subcellular localization, suggesting that these proteins perform different functions in plants, such as processing, transport, localization, stability, and translation of mRNA molecules. This supposition is consistent with literature data regarding GRPs and biotic and abiotic stresses [49, 50]. Wang et al. [50] analyzing consistent with literature data regarding GRPs and biotic plants, such as processing, transport, localization, stability, suggesting that these proteins perform different functions in plants, such as processing, transport, localization, stability, and translation of mRNA molecules.

In our analysis, 47 potential stress-responsive UR tags were identified. According to Moreno et al. [62], Arabidopsis SHMT1 functions in the photosynthetic pathway and influences resistance to biotic and abiotic stresses. The Arabidopsis SHMT1 mutant (shmt1-1) showed enhanced susceptibility to pathogens, as well as to abiotic stresses (50 mM NaCl and high light intensity). The reduced activity in shmt1-1 mutant appears to hinder the ability of the plant to cope with any kind of additional stress, compromising the cellular mechanisms during oxidative stress. In proteome analysis [63], ten out of twelve drought responsive proteins identified from rice leaf sheaths were upregulated including an SHMT. The authors suggested that SHMT was induced for protection from oxidative degradation under drought stress.

(b) CoA-Thioester Hydrolase (CHY1; Synonym: β-Hydroxy-isobutyryl-CoA Hydrolase). In our analysis, one UR tag of this class showing FC of 2.8 (FC_T = FC_D; Table 3) was identified. This peroxisomal metabolic enzyme is needed for valine catabolism and fatty acid b-oxidation. Analyzing freezing sensitive Arabidopsis mutants (chyl-10) after cold acclimation, Dong et al. [51] observed that the disruption of CHY1 function leads to an excess of methylacrylyl-CoA, causing accumulation of Reactive Oxygen Species (ROS), electrolyte leakage, impairing cold-induced gene expression. Additionally, methylacrylyl-CoA may be sequestered in the peroxisome leading to localized changes in this sub cellular region and influencing peroxisome-derived signals after cold-induction. Potential alterations in auxin response or homeostasis in the chyl mutant may contribute to the impaired cold stress tolerance of the mutant, since peroxisome-defective mutants showed resistance to the inhibitory effects of exogenous IBA, analogous to the IAA molecule (a hormone that inhibits the root elongation and promotes lateral root formation).

(c) Glutathione Transferase (GST; EC 2.5.1.18). GSTs encode an ancient, heterogeneous, and widely distributed protein group in living organisms catalyzing a variety of reactions [52], including hormonal metabolism, vacuolar sequestration of anthocyanin, tyrosine metabolism, hydroperoxide detoxification, and regulation of apoptosis [52, 53]. In our study, one UR tag (SD92627) associated to GST showed a significant expression modulation (FC_T = 5.7 and FC_D = 2.4; Table 3). GST expression is induced by a wide variety of stresses, as oxidative stress [54], xenobiotic-type of stresses [55], and dehydration [56]. Expression of TaGSTU1B (Triticum aestivum) was induced by drought stress in four genotypes investigated, but high transcript amounts were detected only in drought-tolerant genotypes [57]. George et al. [58] reported the subcellular localization and the ability of GST from Prosopis juliflora (PjGSTU1), a drought-tolerant woody Fabaceae species, to confer drought tolerance in transgenic tobacco. Ji et al. [59] working with tobacco plants overexpressing a GST gene from Glycine soja showed six-fold higher GST activity enhanced dehydration tolerance than wild-type plants.

(d) Serine Hydroxymethyltransferase (SHMT; EC 2.1.2.1). The SHMT genic family comprises five genes in A. thaliana [60] bearing both cytosolic and mitochondrial isoforms in eukaryotes [61] with activity associated to the Serine and Glycine metabolism (EMBL, 2010). In our evaluation two SHMT candidates (SD243418 (FC_T = 4.0; FC_D = ns); SD179937 (FC_T = 3.8; FC_D = 2.5); Table 3) were identified. According to Moreno et al. [62], Arabidopsis SHMT1 functions in the photosynthetic pathway and influences resistance to biotic and abiotic stresses. The Arabidopsis SHMT1 mutant (shmt1-1) showed enhanced susceptibility to pathogens, as well as to abiotic stresses (50 mM NaCl and high light intensity). The reduced activity in shmt1-1 mutant appears to hinder the ability of the plant to cope with any kind of additional stress, compromising the cellular mechanisms during oxidative stress. In proteome analysis [63], ten out of twelve drought responsive proteins identified from rice leaf sheaths were upregulated including an SHMT. The authors suggested that SHMT was induced for protection from oxidative degradation under drought stress.

(e) Peptidyl-Prolyl Cis-Trans Isomerase (PPlase). It is also known as rotamases or immunophilins (cyclophilins included), which is an enzyme superfamily with catalytic function, facilitating metabolism regulation through a chaperone or a cis-trans isomerization of proline residues during protein folding [64, 65]. A UR tag (SD169158) showing an FC_T = 3.2 (Table 3) concerns a potential PPlase. In plants PPlases have been associated with the response to adverse environmental conditions. Using contrasting genotypes of Sorghum bicolor under water deficit, Sharma and Singh [64] observed a significant increase in leaf- and root-PPlase activity in the drought-tolerant cultivar. Similarly, various rice PPlases were differentially expressed under water deficit and salinity (200 mM NaCl) stresses [65]. Also, a correlation with plant hormones was pointed out by Godoy et al. [66] working with cyclophilins (CyPs) of Solanum tuberosum. CyPs are ubiquitous proteins with an intrinsic enzymatic activity of PPlase that catalyzes the rotation of X-Pro peptide bonds. StCyP mRNA accumulation was stimulated by the application of abscisic acid (ABA) and methyl jasmonate (MeJA) in potato tubers. The accumulation of StCyP transcripts was also detected when the potato tubers were exposed to heat-shock treatment.

(f) Viviparous14. It is a key enzyme involved in the biosynthesis of the phytohormone abscisic acid [67], represented
in our analysis by the SD140270 tag with FC$_T$ of 3.2 (Table 3). Viviparous genes are encoded in the process of plant vivipary, also reported as early germination. Of the 15 genes described so far for maize, 12 control specific steps in ABA biosynthesis [68, 69], with \textit{vpi14} (viviparous-14), associated to the control of final steps of ABA synthesis, encoding a 9-cis-epoxytocarotenoid dioxygenase 1 (NCED1) enzyme that catalyzes the cleavage of the C40 neoxanthin chain into the C15 ABA skeleton xanthoxin [70]. maize mutants for the nced1 gene have strongly reduced kernel ABA content [71] while in \textit{Arabidopsis}, NCED1 overexpression conferred a significant increase in ABA accumulation in the plant and also in drought tolerance [72]. Wan and Li [73] demonstrated that the expression of \textit{AtNCED1} gene in peanut plants was significantly upregulated by dehydration and high salinity (250 mmol L$^{-1}$ NaCl).

\textbf{(g) Branched-Chain Amino Acid Transaminase.} BCAATs are enzymes that play a crucial role in the metabolic pathway of BCAAs (branched-chain amino acids that include leucine, isoleucine, and valine) by catalyzing the last step of synthesis and the initial step of degradation of these amino acids [74]. Plants contain a small family of \textit{bcat} genes, which have been characterized in \textit{Solanum tuberosum} (potato), \textit{Hordeum vulgare}, and \textit{A. thaliana} [75, 76]. Malatrasi et al. [77] evaluated the role of these enzymes in the drought tolerance process. In this study, the transcriptional levels of \textit{Hybcat-1}, in \textit{H. vulgaris}, increased seven folds (results obtained by double checking with RTqPCR) after progressive drought stress (up to 14 days of water deprivation). Physiologically, the authors associated the overregulation to the activation of the BCAAs catabolism, since this is the first enzyme in the branched-chain amino acid (BCAA) catabolic pathway. In high concentrations, these amino acids are toxic to the cells; therefore, activation of their catabolism may play an important role as detoxification mechanism. In our analysis, two UR tags annotated as BACTs were identified exhibiting an expressive modulation of the FC, mainly for the SD270381 tag (FC of 6.8 for both FC$_T$ and FC$_D$; Table 3), while the other tag (SD238059) showed an FC of 2.4 (FC$_T$ = FC$_D$).

\textbf{(h) Allene Oxide Synthase.} AOS is the first enzyme in the pathway leading to the biosynthesis of Jasmonic acid (JA), catalyzing the production of unstable allene epoxides that cyclize to form cyclopentenone acids, the precursors for JA [78]. Three tags of this category were identified (SD270381, SD272257, SD63148) being upregulated in most comparisons (Table 3). For example, the SD270381 tag presented high FC values in both T and D comparisons. The overexpression of AOSs has been observed also in other drought assays, as reported by Ozturk et al. [79] and Talame et al. [80] with barley (\textit{H. vulgaris}) and peanut (\textit{Arachis hypogaea}) [81].

\textbf{(i) Na$^+$/H$^+$ Antipporter.} Membrane proteins involved in the Na$^+$ and H$^+$ transport of both eukaryotes and prokaryotes act in the homeostasis maintenance of such ions [82]. In our analysis, the SD213044 tag, annotated as potential Na$^+$/H$^+$ antiporter, was overexpressed in both analyzed contrasting situations (Table 3). Assays evaluating those proteins under salinity stress showed that these salt-responsive genes may be able to activate the expression of drought-related genes in the tolerance acquisition [83]. Thus, ions are stored in vacuoles, acting as osmolytes, decreasing the hydric potential of the cell. Evaluations with transgenic plants overexpressing those genes, including \textit{Petunia hybrida} [83], \textit{A. thaliana} [84], and \textit{A. hypogaea} [85], conferred higher tolerance to dehydration under drought and salinity.

\textbf{(j) Glutathione Peroxidase (EC 1.11.1.9).} In the present approach, a UR GPX candidate (SD219102) was overexpressed FC$_T$ of 3.2 (Table 3). These enzymes are known as cell protectors against oxidative damage generated by reactive oxygen species [86]. They present a very broad distribution in the cell, occurring in several subcellular compartments [87]. Miao et al. [88] suggested that ATGPX3 might play dual and distinctive roles in H$_2$O$_2$ homeostasis, acting as a general scavenger and relaying the H$_2$O$_2$ signal, and also as an oxidative ABA signal transducer during drought stress signaling. Their differential regulation during biotic and abiotic stresses was reported by Navrot et al. [87], indicating their importance for plant breeding.

\textbf{(l) Serine-Threonine Kinase SAPK1 (Also Known as JNK).} It belongs to the MAPK family [89], including important proteins active in the osmosensory signal transduction pathways in cells exposed to osmotic stress [90]. A wheat candidate (W55a) with about 90% homology to rice SAPK1 was evaluated by Xu et al. [91]. Transgenic \textit{Arabidopsis} plants overexpressing W55a exhibited higher tolerance to drought, being also upregulated by salt, exogenous abscisic acid, salicylic acid, ethylene, and methyl jasmonate. In addition, W55a transcripts were abundant in leaves, but not in roots or stems, under environmental stresses. Expression of SAPK members analyzed by RNA gel blot hybridization with samples of leaves (blades and sheath), roots, and treatments with ABA (50 \textmu M), NaCl (150 mM), or mannitol (600 mM) showed that SAPK1 was upregulated by all three treatments in both roots and leaves, although the effect of ABA was weaker than those of the other two treatments. SAPK6 was weakly upregulated by all treatments in the blades and the sheaths, and weakly by ABA or NaCl but strongly by mannitol treatment in the roots [92]. Overexpressed candidates analyzed here (SD129463; FC$_T$ = 3.0; Table 3) included an SAPK1 as well as a second tag matching SAPK6 (SD87319; FC$_T$ = 2.4 and FC$_D$ = 1.6; Table 3), both in roots, indicating their activation also in this tissue.

\textbf{(m) Delta-1-Pyrroline-5-Carboxylate Synthetase.} P5CS is an enzyme that catalyzes the initiation of the proline biosynthesis in plants [93]. The excessive production of this amino acid would increase the osmotolerance in plants [94]. Rice plants transformed with the P5CS gene underwent 10 days of irrigation withdrawal with higher growth rates, when compared to the control group [94]. Effects of salt in transgenic tobacco transformed with P5CS gene revealed the overexpression of P5CS after 24–48 h exposure to...
NaCl (300 mM), when compared with non-transgenic plants under the same stress [95]. Transgenic lines of petunia [96] and tobacco [97] with enhanced accumulation of proline showed also high drought tolerance. Transcripts involved in amino acid metabolism, such as P5CS, OAT and AS, were also induced more than 10 folds during the identification of drought-responsive genes during sucrose accumulation and water deficit in sugarcane [98]. In our study, a UR tag (SD251703) showing an expressive induction (FC_T 7.1) was annotated as a potential P5CS candidate (Table 3).

4. Concluding Remarks

The present report is the first to analyze contrasting sugarcane accessions under drought stress with a combination of the high-throughput transcriptome profiling SuperSAGE technology coupled with a next-generation sequencing platform. This approach allowed the identification of many potential target candidates in the drought stress response. The adopted methodology of annotation and GO categorization revealed the success of the work in accessing genes from very different pathways, ranging from those controlling the perception and first reaction against the stress (as transcription factors) to those known as classic genes of the osmotic stress (as P5CS). The number of induced tags (213) with GO categorization and high modulation is surprising, especially considering the short time (24 h) after drought stress application. Besides, a high number of important gene candidates with no hits (145)—probably completely new to the research community—will demand additional efforts for the recognition of their function. Validation procedures as well as transient expression assays are planned for future works, aiming to collaborate with breeding and biotechnological approaches for the benefit of the sugarcane culture, especially facing the scenario of future climate changes.

Conflict of Interests

The authors declare that they have no conflict of interests.

Acknowledgments

The authors thank Professor Günter Kahl (Frankfurt University, Germany). This work has been funded by Brazilian institutions: Financiadora de Estudos e Projetos (FINEP), Fundação de Amparo à Ciência e Tecnologia do Estado de Pernambuco (FACEPE), and Conselho Nacional de Desenvolvimento Científico e Tecnológico (CNPq).

References

[1] A. D’Hont, G. M. Souza, M. Menossi et al., “Sugarcane: a major source of sweetness, alcohol, and bio-energy,” in Genomics of Tropical Crop Plants, P. H. Moore and R. Moore, Eds., pp. 483–513, Springer, New York, NY, USA, 2008.

[2] A. J. Waclawowsky, P. M. Sato, C. G. Lembke, P. H. Moore, and G. M. Souza, “Sugarcane for bioenergy production: an assessment of yield and regulation of sucrose content,” Plant Biotechnology Journal, vol. 8, no. 3, pp. 263–276, 2010.

[3] FAOSTAT, “Food and Agriculture Organization of the United Nations. In: FAO Statistical Databases,” 2010, http://faostat.fao.org/.

[4] UNICA, “União da indústria de Cana-de-açúcar. In: Statistics of sugarcane sector—2009,” 2009, http://www.unica.com.br/dadosCotacao/estatistica/.

[5] J. Goldberg, “Ethanol for a sustainable energy future,” Science, vol. 315, no. 5813, pp. 808–810, 2007.

[6] M. Menossi, M. C. Silva-Filho, M. Vincentz, M. A. Van-Sluiys, and G. M. Souza, “Sugarcane functional genomics: gene discovery for agronomic trait development,” International Journal of Plant Genomics, vol. 2008, Article ID 458732, 11 pages, 2008.

[7] E. A. Bray, J. Bailey-Serres, and E. Wetternhly, “Responses to abiotic stresses,” in Biochemistry and Molecular Biology of Plants, W. Gruissem, B. Buchanann, and R. Jones, Eds., pp. 1158–1249, American Society of Plant Physiologists, Rockville, Md, USA, 2000.

[8] L. Taiz and E. Zeiger, Fisiologia Vegetal, S. A. Artmed Editora, Porto Alegre, Brazil, 2004.

[9] W. Wang, B. Vinocur, and A. Altman, “Plant responses to drought, salinity and extreme temperatures: towards genetic engineering for stress tolerance,” Plantas, vol. 218, no. 1, pp. 1–14, 2003.

[10] L. Cattivelli, F. Rizza, F. W. Badeck et al., “Drought tolerance improvement in crop plants: an integrated view from breeding to genomics,” Field Crops Research, vol. 105, no. 1-2, pp. 1–14, 2008.

[11] V. Y. Patade, A. N. Rai, and P. Suprasanna, “Expression analysis of sugarcane shaggy-like kinase (SuSK) gene identified through cDNA subtractive hybridization in sugarcane (Saccharum officinarum L.),” Protoplasma, vol. 248, no. 3, pp. 613–621, 2010.

[12] K. Nakashima and K. Shinozaki, “Regulons involved in osmotic stress-responsive and cold stress-responsive gene expression in plants,” Physiologia Plantarum, vol. 126, no. 1, pp. 62–71, 2006.

[13] T. Umezawa, M. Fujita, Y. Fujita, K. Yamaguchi-Shinozaki, and K. Shinozaki, “Engineering drought tolerance in plants: discovering and tailoring genes to unlock the future,” Current Opinion in Biotechnology, vol. 17, no. 2, pp. 113–122, 2006.

[14] F. A. Rodrigues, M. L. de Laia, and S. M. Zingaretti, “Analysis of gene expression profiles under water stress in tolerant and sensitive sugarcane plants,” Plant Science, vol. 176, no. 2, pp. 286–302, 2009.

[15] R. A. Shimkets, “Gene expression quantitation technology summary,” in Gene Expression Profile: Methods and Protocol, R. A. Shimkets, Ed., pp. 1–12, Humana Press, New Haven, Conn., USA, 2004.

[16] H. Matsumura, S. Reich, A. Ito et al., “Gene expression analysis of plant host-pathogen interactions by SuperSAGE,” Proceedings of the National Academy of Sciences of the United States of America, vol. 100, no. 26, pp. 15718–15723, 2003.

[17] R. Terauchi, H. Matsumura, D. H. Krüger, and G. Kahl, “SuperSAGE: the most advanced through comparative genomics,” in The Handbook of Plant Functional Genomics: Concepts and Protocols, G. Kahl and K. Meksem, Eds., pp. 37–54, Wiley-VCH, Weinheim, Germany, 2008.

[18] V. E. Velculescu, L. Zhang, B. Vogelstein, and K. W. Kinzler, “Serial analysis of gene expression,” Science, vol. 270, no. 5235, pp. 484–487, 1995.
a common motif?” Biochimica et Biophysica Acta, vol. 1492, no. 1, pp. 1–14, 2000.
[50] S. Wang, D. Liang, S. Shi, F. Ma, H. Shu, and R. Wang, “Isolation and characterization of a novel drought responsive gene encoding a glycine-rich RNA-binding protein in Malus prunifolia (Wild.) borkh,” Plant Molecular Biology Reporter, vol. 29, no. 1, pp. 125–134, 2011.
[51] C. H. Dong, B. K. Zolman, B. Bartel et al., “Disruption of Arabidopsis CHY1 reveals an important role of metabolic status in plant cold stress signaling,” Molecular Plant, vol. 2, no. 1, pp. 59–72, 2008.
[52] P. G. Sappl, A. J. Carroll, R. Clifton et al., “The Arabidopsis glutathione transferase gene family displays complex stress regulation and co-silencing multiple genes results in altered metabolic sensitivity to oxidative stress,” The Plant Journal, vol. 58, no. 1, pp. 53–68, 2009.
[53] D. P. Dixon, M. Skipsey, and R. Edwards, “Roles for glutathione transferases in plant secondary metabolism,” Phytochemistry, vol. 71, no. 4, pp. 338–350, 2010.
[54] S. S. Gill and N. Tuteja, “Reactive oxygen species and antioxidant machinery in abiotic stress tolerance in crop plants,” Plant Physiology and Biochemistry, vol. 48, no. 12, pp. 909–930, 2010.
[55] R. Edwards and D. P. Dixon, “The role of glutathione transferases in herbicide metabolism,” in Herbicides and Their Mechanisms of Action, A. H. Cobb and R. C. Kirkwood, Eds., pp. 38–71, Sheffield Academic Press, Sheffield, UK, 2000.
[56] M. W. Bianchi, C. Roux, and N. Vartanian, “Drought regulation of GST8, encoding the Arabidopsis homologue of ParC/Nt107 glutathione transferase/peroxidase,” Physiologia Plantarum, vol. 116, no. 1, pp. 96–105, 2002.
[57] A. Gallé, J. Csiszár, M. Secenji et al., “Glutathione transferase activity and expression patterns during grain filling in flag leaves of wheat genotypes differing in drought tolerance: response to water deficit,” Journal of Plant Physiology, vol. 166, no. 17, pp. 1878–1891, 2009.
[58] S. George, G. Venkataraman, and A. Parida, “A chloroplast-localized and auxin-induced glutathione S-transferase from phreatophyte Prosopis juliflora confer drought tolerance on tobacco,” Journal of Plant Physiology, vol. 167, no. 4, pp. 311–318, 2010.
[59] W. Ji, Y. Zhu, Y. Li et al., “Over-expression of a glutathione S-transferase gene, GsGST, from wild soybean (Glycine soja) enhances drought and salt tolerance in transgenic tobacco,” Biotechnology Letters, vol. 32, no. 8, pp. 1173–1179, 2010.
[60] C. R. McClung, M. Hsu, J. E. Painter, J. M. Gagne, S. D. Karlsberg, and P. A. Salomé, “Integrated temporal regulation of the photorespiratory pathway. Circadian regulation of two Arabidopsis genes encoding serine hydroxy-methyltransferase,” Plant Physiology, vol. 123, no. 1, pp. 381–391, 2000.
[61] E. A. Cossins and L. Chen, “Folates and one-carbon metabolism in plants and fungi,” Phytochemistry, vol. 45, no. 3, pp. 437–452, 1997.
[62] J. I. Moreno, R. Martin, and C. Castresana, “Arabidopsis SLIM1, a serine hydroxymethyltransferase that functions in the photosynthetic pathway influences resistance to biotic and abiotic stress,” The Plant Journal, vol. 41, no. 3, pp. 451–463, 2005.
[63] G. M. Ali and S. Komatsu, “Proteomic analysis of rice leaf sheath during drought stress,” Journal of Proteome Research, vol. 5, no. 2, pp. 396–403, 2006.
[64] A. D. Sharma and P. Singh, “Comparative studies on drought-induced changes in peptidyl prolyl cis-trans isomerase activity in drought-tolerant and susceptible cultivars of Sorghum bicolor,” Current Science, vol. 84, no. 7, pp. 911–918, 2003.
[65] J. C. Ahn, D. W. Kim, Y. N. You et al., “Classification of rice (Oryza sativa L. Japonica nippobire) immunophilins (FKBPs, CYPs) and expression patterns under water stress,” BMC Plant Biology, vol. 10, article 253, 2010.
[66] A. V. Godoy, A. S. Lazzaro, C. A. Casalongué, and B. San Segundo, “Expression of a Solanum tuberosum cyclophilin gene is regulated by fungal infection and abiotic stress conditions,” Plant Science, vol. 152, no. 2, pp. 123–134, 2000.
[67] S. A. J. Messing, S. B. Gabelli, I. Echeverria et al., “Structural insights into maize viviparous14, a key enzyme in the biosynthesis of the phytohormone abscisic acid,” The Plant Cell, vol. 22, no. 9, pp. 2970–2980, 2010.
[68] D. R. McCarty, “Genetic control and integration of maturation and germination pathways in seed development,” Annual Review of Plant Physiology and Plant Molecular Biology, vol. 46, pp. 71–93, 1995.
[69] D. Durantini, A. Giuliani, A. Malgioglio et al., “Vivipary as a tool to analyze late embryogenic events in maize,” Heredity, vol. 101, no. 5, pp. 465–470, 2008.
[70] S. H. Schwartz, B. C. Tan, D. A. Gage, J. A. D. Zeevaart, and D. R. McCarty, “Specific oxidative cleavage of carotenoids by VP14 of maize,” Science, vol. 276, no. 5320, pp. 1872–1874, 1997.
[71] B. C. Tan, S. H. Schwartz, J. A. D. Zeevaart, and D. R. McCarty, “Genetic control of abscisic acid biosynthesis in maize,” Proceedings of the National Academy of Sciences of the United States of America, vol. 94, no. 22, pp. 12235–12240, 1997.
[72] S. Iuchi, M. Kobayashi, T. Taji et al., “Regulation of drought tolerance by gene manipulation of 9-cis-epoxycarotenoid dioxygenase, a key enzyme in abscisic acid biosynthesis in Arabidopsis,” The Plant Journal, vol. 27, no. 4, pp. 325–333, 2001.
[73] X. R. Wan and L. Li, “Regulation of ABA level and water-stress tolerance of Arabidopsis by ectopic expression of a peanut 9-cis-epoxycarotenoid dioxygenase gene,” Biochemical and Biophysical Research Communications, vol. 347, no. 4, pp. 1030–1038, 2006.
[74] F. Gao, C. Wang, C. Wei, and Y. Li, “A branched-chain aminotransferase may regulate hormone levels by affecting KNOX genes in plants,” Planta, vol. 230, no. 4, pp. 611–623, 2009.
[75] M. A. Campbell, J. K. Patel, J. L. Meyers, L. C. Myrick, and J. L. Gustin, “Genes encoding for branched-chain amino acid aminotransferase are differentially expressed in plants,” Plant Physiology and Biochemistry, vol. 39, no. 10, pp. 855–860, 2001.
[76] R. Diebold, J. Schuster, K. Däschner, and S. Binder, “The branched-chain amino acid transaminase gene family in Arabidopsis encodes plastid and mitochondrial proteins,” Plant Physiology, vol. 129, no. 2, pp. 540–550, 2002.
[77] M. Malatrasi, M. Corradi, J. T. Svensson, T. J. Close, M. Gulli, and N. Marmiroli, “A branched-chain amino acid aminotransferase gene isolated from Hordeum vulgare is differentially regulated by drought stress,” Theoretical and Applied Genetics, vol. 113, no. 6, pp. 965–976, 2006.
[78] M. J. Mueller, “Enzymes involved in jasmonic acid biosynthesis,” Physiologia Plantarum, vol. 100, no. 3, pp. 653–663, 1997.
[79] Z. N. Ozturk, V. Talamé, M. Deyholos, T. A. E. Zeevaart, and D. A. Gage, “Aminotransferase gene family displays complex stress regulation and co-silencing multiple genes results in altered metabolic sensitivity to oxidative stress,” The Plant Journal, vol. 41, no. 3, pp. 437–452, 1997.
[80] M. J. Mueller, “Enzymes involved in jasmonic acid biosynthesis,” Physiologia Plantarum, vol. 100, no. 3, pp. 653–663, 1997.
[80] V. Talamé, N. Z. Ozturk, H. J. Bohnert, and R. Tuberosa, "Barley transcript profiles under dehydration shock and drought stress treatments: a comparative analysis," *Journal of Experimental Botany*, vol. 58, no. 2, pp. 229–240, 2007.

[81] K. M. Devaiah, G. Bali, T. N. Athmaram, and M. S. Basha, "Identification of two new genes from drought tolerant peanut up-regulated in response to drought," *Plant Growth Regulation*, vol. 52, no. 3, pp. 249–258, 2007.

[82] E. Padan and S. Schuldiner, "Na+/H+ antiporters, molecular devices that couple the Na+ and H+ circulation in cells," *Journal of Bioenergetics and Biomembranes*, vol. 25, no. 6, pp. 647–669, 1993.

[83] K. Xu, P. Hong, L. Luo, and T. Xia, "Overexpression of AtNHX1, a vacuolar Na+/H+ antiporter from *Arabidopsis thaliana*, in *Petunia hybrida* enhances salt and drought tolerance," *Journal of Plant Biology*, vol. 52, no. 5, pp. 453–461, 2009.

[84] F. Brini, M. Hanin, I. Mezghani, G. A. Berkowitz, and K. Masmoudi, "Overexpression of wheat Na+/H+ antiporter TNHX1 and H+-pyrophosphatase TVP1 improve salt- and drought-stress tolerance in *Arabidopsis thaliana* plants," *Journal of Experimental Botany*, vol. 58, no. 2, pp. 301–308, 2007.

[85] M. A. Asif, Y. Zafar, J. Iqbal et al., "Enhanced expression of AtNHX1, in transgenic groundnut (*Arachis hypogaea* L.) improves salt and drought tolerance," *Molecular Biotechnology*, vol. 49, no. 3, pp. 250–256, 2011.

[86] M. A. R. Milla, A. Maurer, H. A. Rodríguez, and J. P. Gustafson, "Glutathione peroxidase genes in *Arabidopsis* are ubiquitous and regulated by abiotic stresses through diverse signaling pathways," *The Plant Journal*, vol. 36, no. 5, pp. 602–615, 2003.

[87] N. Navrot, V. Collin, J. Gualberto et al., "Plant glutathione peroxidases are functional peroxiredoxins distributed in several subcellular compartments and regulated during biotic and abiotic stresses," *Plant Physiology*, vol. 142, no. 4, pp. 1364–1379, 2006.

[88] Y. Miao, D. Lv, P. Wang et al., "An Arabidopsis glutathione peroxidase functions as both a redox transducer and a scavenger in abscisic acid and drought stress responses," *The Plant Cell*, vol. 18, no. 10, pp. 2749–2766, 2006.

[89] Y. Fleming, C. G. Armstrong, N. Morrice, A. Paterson, M. Goedert, and P. Cohen, "Synergistic activation of stress-activated protein kinase 1c-JNK isoforms by mitogen-activated protein kinase kinase 4 (MKK4) and MKK7," *Biochemical Journal*, vol. 352, no. 1, pp. 145–154, 2000.

[90] S. Kültz, "Evolution of osmosensory MAP kinase signaling pathways," *Integrative and Comparative Biology*, vol. 41, no. 4, pp. 743–757, 2001.

[91] Z. S. Xu, L. Liu, Z. Y. Ni et al., "W55a encodes a novel protein kinase that is involved in multiple stress responses," *Journal of Integrative Plant Biology*, vol. 51, no. 1, pp. 58–66, 2009.

[92] Y. Kobayashi, S. Yamamoto, H. Minami, Y. Kagaya, and T. Hattori, "Differential activation of the rice sucrose nonfermenting1-related protein kinase2 family by hyperosmotic stress and abscisic acid," *The Plant Cell*, vol. 16, no. 5, pp. 1163–1177, 2004.

[93] P. B. K. Kishor, S. Sangam, R. N. Amrutha et al., "Regulation of proline biosynthesis, degradation, uptake and transport in higher plants: its implications in plant growth and abiotic stress tolerance," *Current Science*, vol. 88, no. 3, pp. 424–438, 2005.

[94] B. Zhu, J. Su, M. Chang, D. P. S. Verma, Y. L. Fan, and R. Wu, "Overexpression of a Δ1-pyrroline-5-carboxylate synthetase gene and analysis of tolerance to water- and salt-stress in transgenic rice," *Plant Science*, vol. 139, no. 1, pp. 41–48, 1998.

[95] R. Razavizadeh and A. A. Ehsanpour, "Effects of salt stress on proline content, expression of delta-1-pyrroline-5-carboxylate synthetase, and activities of catalase and ascorbate peroxidase in transgenic tobacco plants," *Biological Letters*, vol. 46, no. 2, pp. 63–75, 2009.

[96] M. Yamada, H. Morishita, K. Urano et al., "Effects of free proline accumulation in petunias under drought stress," *Journal of Experimental Botany*, vol. 56, no. 417, pp. 1975–1981, 2005.

[97] J. Gubis, R. Vaňková, V. Červená et al., "Transformed tobacco plants with increased tolerance to drought," *South African Journal Botany*, vol. 73, no. 4, pp. 505–511, 2007.