

De novo RNA sequencing and analysis of the transcriptome of signalgrass (*Urochloa decumbens*) roots exposed to aluminum

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Abstract Acidic soils occupy a vast area in the world, and the aluminum (Al) in these soils can directly interact with plant cells and tissues to inhibit their growth and reduce yields. The signalgrass *Urochloa decumbens* cv. Basilisk (syn. *Brachiaria decumbens* cv. Basilisk), a widely sown tropical forage grass, is recognized for its high productivity under intensive use, vigorous growth, ease of establishment, and good forage value throughout the year, as well as its exceptional adaptation to infertile acid soils. We sequenced the transcriptome from roots of *U. decumbens* cv. Basilisk under two conditions, with and without Al, using Illumina paired-end sequencing technology and performed de novo assembly of those reads, which yielded 164,920 transcripts. Of these transcripts, 113,918 were assigned a putative protein function through comparisons with different gene set databases. Additionally, 13,375 simple sequence repeat (SSR) markers were identified. Digital gene expression analyses were conducted to identify 6698 differentially expressed genes between treatments, revealing a great differences in the root transcriptional landscape

when exposed to aluminum. An extensive annotation of the differentially expressed genes (DEGs), made possible to identify several transcripts with putative functions correlated to aluminum exposure, most belonging to vesicle transportation, cell wall modifications and metal handling ontologies. In this work, abundant, high-quality transcripts were obtained, providing a reference platform for future biotechnological studies and breeding programs for this species and its close relatives.

Keywords Aluminum tolerance · RNA-seq · *Brachiaria decumbens* · Molecular markers

Introduction

Tropical and subtropical countries account for approximately half the global beef cattle production (Jank et al. 2011). In Brazil 169 million hectares of pastures are used for feeding 208 million cattle (ABIEC 2013). The main grasses that are used as forage belong to species of the *Brachiaria* genus (syn. *Urochloa* P. Beauv.).

Although a large amount of genetic information has been deposited in public databases involving Poaceae family (28 genome sequencing projects, and 7,727,397 DNA and RNA sequences) (NCBI 2016a), the majority of the sequences are derived from a few species of high agricultural importance (rice, maize, wheat, and sorghum). In contrast with well-investigated crops, the Panicodae subfamily is of high economical importance (Toledo-Silva et al. 2013), but is genomically poorly explored.

Nevertheless, the production of genomic information is high demanding in terms of time and money; therefore, comparative genomics becomes a convenient method for studying the genetics of different species, providing

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resources in terms of molecular tools for genetic and genomic studies. As demonstrated by Surget-Groba and Montoya-Burgos (2010) and Liu et al. (2014), transcriptomic studies can be greatly improved with scaffolding steps that use closely genetically related protein sets as a proxy to assemble transcripts.

The species of the genus *Urochloa* were brought to America from Africa and were widely used in pasture for livestock grazing, especially *Brachiaria decumbens* and *Brachiaria brizantha*, which have shown wide adaptation and are extensively used as pasture grasses in this ecosystem (do Valle et al. 1996). *Urochloa decumbens* (Stapf) R.D. Webster, known as signalgrass, has gained special attention due to its high productivity under intensive use, vigorous growth, ease of establishment, and good forage value throughout the year, as well as its exceptional adaptation to infertile acid soils (Valle et al. 2010), which is mainly due to its high Al tolerance (Arroyave et al. 2013; Ramos et al. 2012; Rao et al. 1993).

Under acidic soil conditions, Al is ionized to trivalent aluminum (Al^{3+}), which is readily available and highly reactive in the soil. Al can interact with a number of extracellular and intracellular structures, inhibiting plant growth and reducing productivity and yield. Al is one of the major limiting factors for growing crops in acidic soils (Delhaize and Ryan 1995).

Given the importance of this abiotic stress, the phytotoxicity of Al^{3+} has been studied extensively (Foy 1974). The root apex is the first site of Al interaction. There, Al immediately starts to damage cells, inhibiting cell elongation and division. In vegetal cells, Al can modify the cellular wall and membrane structure, harming water and nutrient absorption and compromising plant development (Arroyave et al. 2011).

In Brazil, only one cultivar of *U. decumbens* has been commercially available since the 1960s, cv. Basilisk, which is apomictic and tetraploid ($2n = 4x = 36$), with high aluminum tolerance (Oliveira et al. 2015). To develop new cultivars of this species, diploid accessions were artificially tetraploidized using colchicine (Simioni and Valle 2009) in crosses with *U. decumbens* cv. Basilisk.

Despite the good adaptation to savanna regions of Brazil and the high economic importance of *U. decumbens* for animal production in many tropical countries, the genomic data on the species are limited; expanding the current knowledge on this species could lead to a more efficient breeding program. Although some genomic resources are available for the genus *Urochloa* (mainly 2237 EST sequences available) (NCBI 2016b), almost no genetic information is available on molecular markers or gene expression profiles under biological stresses for the genus or this particular species.

SSRs markers are one of the most abundant data correlated to genetic variability obtained by transcriptome sequencing and highly transferable to close related species, typically co-dominant markers are widely used in studies of the Poaceae family given their high abundance and stability (Wang et al. 2015; Santos et al. 2015), and also can be applied to germplasm characterization (Kumari et al. 2013). Although EST (expressed sequence tags) derived SSR markers are less informative, they represent functional sequences and are a cost-effective biotechnological tool for breeders (Salgado et al. 2014; Wang et al. 2015).

Recently, we obtained the transcriptome from roots of *U. decumbens* cv. Basilisk for study under stressed conditions using aluminum. Extensive information was generated by performing de novo assembly of this transcriptome. In this paper, we describe the analysis of this transcriptome and provide a reference platform for future biotechnological studies and breeding programs for this species and its close relatives.

This study will provide a better understanding of the molecular events underlying Al tolerance and how they vary among plant species and cultivars from same species, considering that these issues are not clearly understood (Ma et al. 2001).

Materials and methods

Plant material

Approximately 600 *U. decumbens* cv. Basilisk seeds were germinated on seed trays containing sand and vermiculite in a 1:1 proportion. Subsequently, approximately 200 12-day-old seedlings (past germination) were transferred into two hydroponic solutions [(A) 200 CaCl_2 , pH 4.2; and (B) 200 CaCl_2 + 200 AlCl_3 , pH 4.2, prepared as described by Wenz et al. (2006)] in three experimental replicates with 30 seedlings each. All of the seedlings remained in the solution for 8 h. Then, the root tips were removed and cut approximately one centimeter from the tip, yielding six bundles of root tips that were utilized for RNA extraction.

RNA extraction was performed using Trizol[®] (Invitrogen, Camarillo, CA, USA), according to the manufacturer's manual. At the end of the extraction process, the total RNA from the six samples was suspended in 20 μL of MiliQ sterile water and treated with DNase (Invitrogen) according to the manufacturer's manual.

Quantification of the RNA samples was performed on a NanoDrop[®] 1000 (Thermo, Waltham, MA, USA) at a wavelength of 260 nm. RNA sample quality was evaluated using the $A_{260\text{nm}}/A_{280\text{nm}}$ wavelength ratio and by direct observation on a 1% agarose gel. After quantification, the samples were suspended in 70% ethanol and sent to the

Beijing Genomics Institute (BGI) for de novo transcriptome sequencing.

cDNA library construction and sequencing

Illumina sequencing was performed at the Beijing Genomics Institute (BGI)-Shenzhen, Shenzhen, China (<http://www.genomics.cn/index.php>), using a HiSeq™ 2000 platform according to the manufacturer's instructions (Illumina, San Diego, CA, USA). Briefly, poly(A) RNA was isolated from 20 µg of total RNA using Sera-mag Magnetic Oligo (dT) Beads (Illumina). To avoid priming bias when synthesizing the cDNA, the purified mRNA was first fragmented into small pieces. Then, the double-stranded cDNA was synthesized using the SuperScript Double-Stranded cDNA Synthesis kit (Invitrogen) with random hexamer (N6) primers (Illumina). The synthesized cDNA was subjected to end repair and phosphorylation using T4 DNA polymerase, Klenow DNA polymerase and T4 polynucleotide kinase (PNK). The cDNA fragments repaired using these enzymes were 3' adenylated using Klenow Exo-(3–5' exo minus, Illumina). Illumina paired-end adapters were ligated to the ends of the 3'-adenylated cDNA fragments. To select a template size range for downstream enrichment, the products of the ligation reaction were purified on a 2% TAE agarose gel (Certified Low-Range Ultra Agarose, Bio-Rad). A range of cDNA fragments (200 ± 25 bp) was excised from the gel. Fifteen rounds of polymerase chain reaction (PCR) amplification were performed to enrich the purified cDNA template using PCR Primer PE 1.0 and PE 2.0 (Illumina) with Phusion DNA Polymerase. The cDNA library was constructed with a fragment length range of 200 bp (±25 bp). Finally, after validation on an Agilent Technologies 2100 Bioanalyzer using the Agilent DNA 1000 chip kit, the cDNA library was labeled and sequenced on a PE flow cell using an Illumina Genome Analyzer HiSeq 2000.

The workflow was as follows: template hybridization, isothermal amplification, linearization, blocking, sequencing primer hybridization, and sequencing on the sequencer for Read 1. After the completion of the first read, the templates were regenerated in situ to enable a second 90-bp read from the opposite end of the fragments, i.e., the newly sequenced strands were stripped off, and the complementary strands were bridge-amplified to form clusters. Once the original templates were cleaved and removed, the reverse strands underwent sequencing by synthesis.

Pre-processing, de novo assembly, clustering, and scaffolding

After trimming adaptors, reads with unknown nucleotides longer than 5% and low-quality reads (more than 20% of

bases $Q \leq 10$) were removed. De novo assembly was performed by the BGI using Trinity (Haas et al. 2013). The database was sent to Embrapa for bioinformatic analyses. All reads were deposited in the National Center for Biotechnology Information Short Read Archive (NCBI-SRA) under accession number SRP071168, BioProject number PRJNA314352.

Because several samples were available from the same species, a clustering step was performed by TIGR Gene Indices clustering tools (TGICL) (Pertea et al. 2003) to assemble all the unigenes from different samples to form a single set of non-redundant unigenes. Orientation and CDS (protein coding sequence) predictions were obtained using Blastx (Altschul et al. 1990) against the NCBI non-redundant database (Pruitt et al. 2005). The CDSs from sequences with no hits were predicted using ESTScan (Iseli et al. 1999).

To obtain a comprehensive set of genes, a scaffolding set was assembled using TransPS [Transcriptome Post Scaffold (Liu et al. 2014)] and *Setaria italica* version 2.1 (Benetzen et al. 2012) as the reference proteome.

Annotation and screening for plant resistance genes

In an attempt to identify putative functions from the sequences through protein similarity and to identify protein families and metabolic functions, all the sequences were searched against multiple protein databases NCBI-Nr (non redundant database) (Pruitt et al. 2005), Swiss-Prot (European Molecular Biology Laboratory protein database) (Bairoch and Apweiler 2000), KEGG (Kyoto Encyclopedia of Genes and Genomes) (Kanehisa and Goto 2000), and COG (Clusters of Orthologous Groups) (Tatusov et al. 2003) using a Blastx e-value cutoff of 10^{-5} . To retrieve Gene Ontology (GO) (Harris et al. 2004) functional annotation, the Blast2GO (Conesa et al. 2005) program was applied to the sequences. The WEGO (Web Gene Ontology Annotation Plot) (Ye et al. 2006) software was used to perform GO annotation analysis and to plot the GO annotations. For a more detailed annotation of the DEGs (Differentially Expressed Genes), the scaffold proxies of those unigenes as well as the unigenes themselves were annotated with agriGO [GO Analysis Toolkit and Database for the Agricultural Community (Du et al. 2010)] and REVIGO (Reduce and Visualization of Gene Ontology) (Supek et al. 2011).

The similarity between the 164,920 de novo transcripts and 104,447 plant resistance genes (R gene) from the Plant Resistance Gene database (PRGdb) (Sanseverino et al. 2010) was analyzed using standalone BLAST tool (Altschul et al. 1990). Only sequences with e-values equal to 0.0 were accepted as result.

Gene expression

By mapping the reads back to the assembled transcriptome, the relative transcript abundance was obtained using the FPKM method (fragments per kb that were allowed to have zero or one gene in each family per million fragments) (Anders and Huber 2010). Whose formula is:

$$\text{FPKM} = \frac{10^6 \times C}{N \times L/10^3},$$

where FPKM denotes the expression of Unigene A, C denotes the number of fragments that uniquely aligned to Unigene A, N denotes the total number of fragments that uniquely aligned to all Unigenes, and L denotes the number of bases in the CDS of Unigene A. By eliminating the influence of different gene length and sequencing level on the calculation of gene expression, the FPKM values can be directly used for comparing the differences in gene expression among samples.

Differentially expressed genes (Audic and Claverie 1997) were identified after a multiple hypothesis test on the relative transcript abundance values, using H_0 : a gene has the same expression level in two samples; and H_1 : a gene has different expression levels in two samples. Later, a correction test using false discovery rate (FDR) (Yekutieli and Benjamini 2001) combined with the ratio of the FPKMs from control samples was used to determine significant differentially expressed genes. In our analysis, we chose genes with $\text{FDR} \leq 10^{-3}$ and expression \log_2 ratios > 1.5 . To integrate annotation and gene expression levels, MAPMAN (Thimm et al. 2004) was utilized as a visualization tool. Moreover, the top 100 up and down regulated genes with COG annotation were selected and plotted in a M (log ratios) and A (mean average) scale space, where we could relate COG or KEGG annotation codes among the transcripts with the Differential Expression and Protein Interactions Visualization Tool (DEPICTViz). DEPICTViz is an interactive JavaScript web tool for visualizing differential gene expression and protein interactions. This tool allows one to visualize genes of interest in an MA graph, including attributes such as COG Id (with category and description), Swiss-Prot, KEGG and KO (KEGG orthology) Ids, in addition to links to protein interactions and gene ontology databases, such as STRING-DB (Search Tool for the Retrieval of Interacting Genes/Proteins database) and COG. If applicable, visualization using the UCSC (University of California, Santa Cruz) Genome Browser is also allowed, which is available at <http://git.facom.ufms.br/bioinfo/depictviz.git>.

Screening for molecular markers

Searches for SSRs (Simple Sequence Repeats) from the contig dataset were performed using MISA (Microsatellite Identification Tool) (Thiel et al. 2003). The definition of microsatellites was set to 12-fold (1/12), di (2/6), tri (3/5), tetra (4/5), penta (5/5), or hexa (6/4) mononucleotide repeats.

After screening, SSR markers situated more than 150 bp from both ends of the encompassing unigene were retained and used to design primers with Primer3 (Untergasser et al. 2012).

Comparative analysis

Having as input all the protein coding genes from *B. decumbens*, *B. distachyon* (GenBank assembly accession—GCA_000005505.1), *Leersia perrieri* (Department of Energy Joint Genome Institute, DOE-JGI), *Oryza sativa* (GenBank assembly accession—GCA_000005425.2), *Panicum hallii* (DOE-JGI), *Panicum virgatum* (DOE-JGI), *S. italica* (GenBank assembly accession—GCA_000263155.1), *Sorghum bicolor* (GenBank assembly accession—GCA_000236745.2) and *Zea mays* (GenBank assembly accession—GCA_000005005.1), Orthologsorter (Farias 2013) tool was used to build the whole genome phylogenetic species tree. This package compares essentially all protein coding genes from all species in an all-against-all fashion using OrthoMCL (Li et al. 2003). Among all the families found, only those with exactly one representative gene in each genome were used. The only exception was the outgroup genome, *S. italica*, which was allowed to have zero or one gene in each family. After selecting those families, Muscle (Edgar 2004) was used to align each of them, and Gblocks (Castresana 2000) was used to remove non-informative sites. All alignments were then concatenated. To build the tree, the final complete alignment was input into RAxML (Stamatakis 2006) with the PROTCATJTT model, rapid bootstrapping and an ML (maximum likelihood) search.

Results

Pre-processing and de novo assembly

All of the 401 million clean reads (66.8 million clean reads per sample on average) generated using the Illumina platform were assembled into 164,920 unigenes, which were defined here as sequences without Ns that could not be extended on either side. The clustering step assembled unigenes from different samples to form a single set of non-redundant unigenes, resulting in 67,123 clusters and 97,797

Table 1 Assembly summary

	Sample	Total number	Length (nt)	Mean length (nt)	N50	Consensus sequences	Distinct clusters	Distinct singletons
Contig	130511256	320,333	84,479,954	264	364	–	–	–
	130511258	323,943	85,986,326	265	370	–	–	–
	130511260	313,499	82,973,249	265	369	–	–	–
	130511262	281,266	75,135,302	267	380	–	–	–
	130511264	322,648	85,081,726	264	366	–	–	–
	130511265	266,062	71,690,914	269	391	–	–	–
	Unigene	130511256	154,021	89,922,701	584	1119	154,021	47,604
130511258		156,098	92,073,465	590	1140	156,098	47,960	108,138
130511260		149,655	86,377,464	577	1097	149,655	45,078	104,577
130511262		134,088	78,331,874	584	1115	134,088	41,066	93,022
130511264		154,674	92,151,753	596	1160	154,674	48,465	106,209
130511265		126,253	74,180,821	588	1130	126,253	38,565	87,688
All		164,920	148,280,694	899	1522	164,920	67,123	97,797

singletons with lengths ranging from 200 to 18,820 nucleotides (nt) and 63.4% of all sequences with lengths between 500 and 2000+ nt (Table 1).

By retrieving the best BLAST hits from the selected databases, we extracted the CDS information from approximately 62% of the assembled sequences. The BLAST results were also used to train a gene model for ESTScan prediction. CDS prediction by ESTScan successfully resulted in 21,305 CDSs.

Scaffolding and annotation

Using the 116,644 sequences from the *Setaria italica* protein dataset, the scaffolding step successfully identified 11,634 sequences with one sequence per protein relation. A total of 4979 sequences were assembled into 2230 scaffolds that ranged from 237 to 15,200 nt; the scaffolds were formed from a maximum of 2–8 sequences.

With respect to the identification of putative protein functions, we successfully retrieved functional information from 113,918 sequences (69% of all sequences) against the NCBI-nr, NCBI-nt (NCBI nucleotide database), SwissProt, KEGG and COG databases, with low e-values (76% <1e–05) and high similarity (61% of sequences with >60% identity), mainly with proteins from *Sorghum bicolor* and *Zea mays* (Fig. 1).

COG annotation resulted in 55,451 sequences distributed into 121 categories, mainly in “general function prediction only”, “translation, ribosomal structure and biogenesis”, and “function unknown” (Fig. 2). COG annotation of the transcripts provided more insight into the protein function and was used to identify proteins related to broad functions, thus generating a more general sense from the group of transcripts. Although most of the transcript annotations

relied on poorly characterized functions, we were able to verify the significant presence of specific pathways related to AI response, toxicity and tolerance, including a mechanism for the modification of the cell wall, plasma membrane transport processes, intracellular capture, and AI³⁺ DNA binding (Brunner and Sperisen 2013). Specific to this response group, the COG analysis identified 514 transcripts belonging to the “MU” category (“cell wall/membrane/envelope biogenesis; intracellular trafficking, secretion, and vesicular transport”) and 1217 transcripts belonging to the category “V” (defense) annotated as “ABC-type multidrug transport systems”.

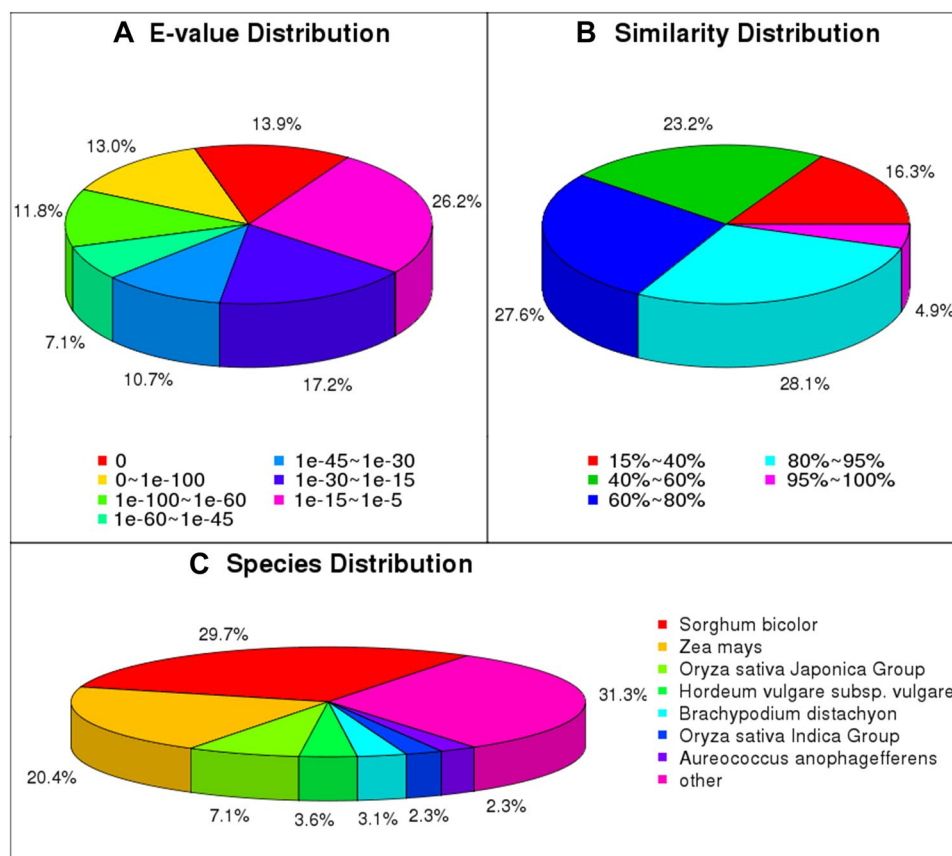
In an attempt to describe the properties of the identified transcripts and their products, a GO annotation step was performed. We successfully annotated 70,008 sequences into 5370 GO terms. Of these terms, the “cellular component” category under the class “cell” is the most representative, followed by “biological processing”, “single-organism process”, “molecular function”, and “catalytic activity” (Fig. 3).

Additionally, to gain more understanding of our set of transcripts, a KEGG annotation step was performed, and we connected 72,749 sequences to 3290 KO’s and 1065 enzyme commission terms. The most representative hits belonged to glycerophospholipid metabolism (KEGG enzyme 3.1.4.4 and KEGG orthology K01115); in addition, all the main metabolic pathways were identified. A list of biochemical pathways predicted from unigenes by KEGG is provided in electronic supplementary material

Gene expression

From the 164,920 de novo transcripts, we identified 4534 DEGs with FDRs $\leq 10^{-3}$ between treatments (Fig. 4) and

Fig. 1 Annotation statistics. **a** E-value distribution; **b** similarity distribution; **c** species distribution



log₂-fold-change values ≥ 1.5 or ≤ -1.5 . From these highly significant DEGs, 1589 were identified as belonging to 1059 scaffolds with 1726 successfully retrieved GO terms. A gene set enrichment analysis (GSEA) annotated 2240 entries into 131 significant GO terms.

When annotation and gene expression were integrated, our results suggested the increased activity of key cell functions. Transcripts annotated under the “vesicle transport” and “transport” categories were highly up-regulated (Fig. 5). A list of DEGs and the integrated annotation is provided in electronic supplementary material.

Screening for molecular markers

A total of 14,932 SSRs (excluding mononucleotide repetitions) were identified in 14,822 transcripts, ranging from 12 to 240 nucleotides in length and possessing four to eleven motif repeats (Table 2). Three-nucleotide motifs were the most common SSRs, accounting for 61% of all SSRs (Fig. 6). The trinucleotides (GCG)*5 and (CGC)*5 were the most frequent.

Additionally, 338 SSRs were found in 292 DEGs. From the scaffolding analysis, the 14,932 markers were placed into 7350 genes. By GSEA analysis, 14,030 transcripts

were annotated into 42 significant GO terms. Under the “biological process” category, the most significant terms were “cellular component organization”, “regulation of biological quality”, “transmembrane transport” and “catabolic process”.

Comparative analysis

A complete genome tree generated from the concatenation of orthologous families (see “Materials and methods”) is presented in Fig. 7. Among 48,844 protein families, 561 with exactly one representative gene from each genome were chosen. After aligning each one of these families, all alignments were concatenated, initially resulting in 166,109 columns and then 386 columns after removing non-representative sites. The topology is entirely compatible with that obtained using the online NCBI Taxonomy Browser tree builder, using the same set of species (<http://www.ncbi.nlm.nih.gov/Taxonomy/CommonTree/wwcmt.cgi>).

Screening for plant resistance gene

A comparison of the 164,920 de novo transcripts and the PRGdb (Sanseverino et al. 2010) yielded 2445 genes with high similarity, 1221 of which were different from each

COG Function Classification of All-Unigene.fa Sequence

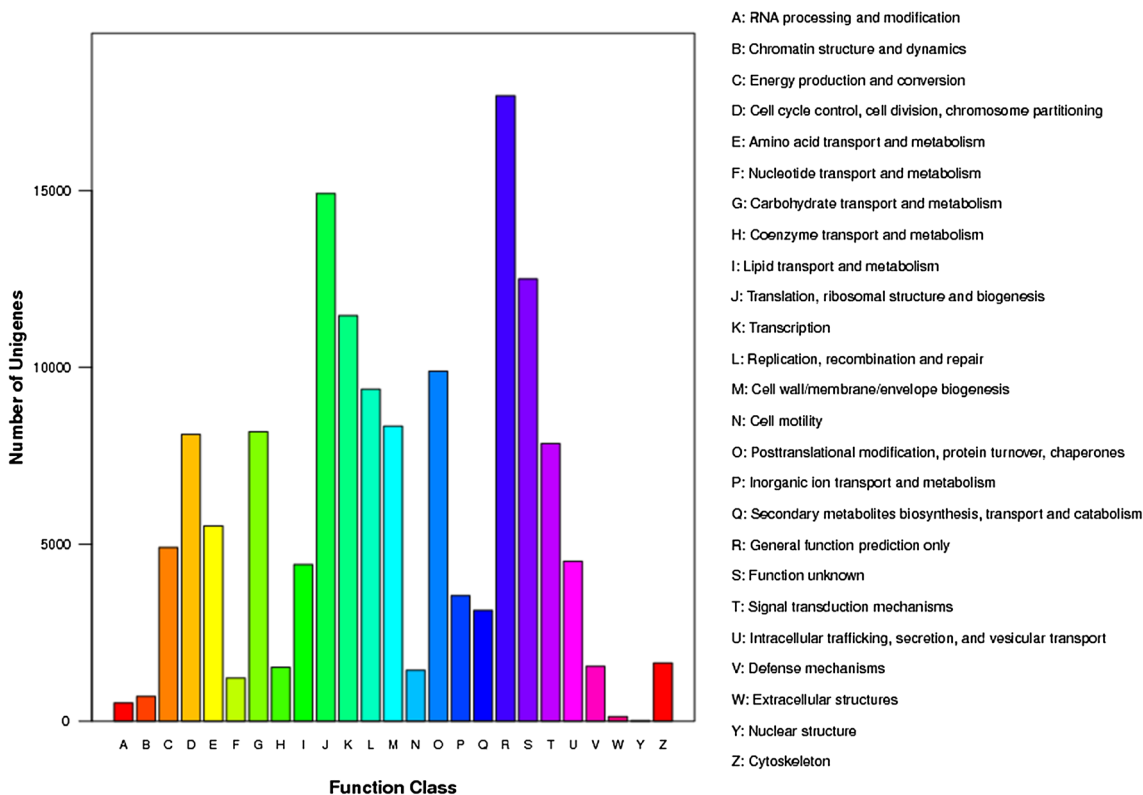


Fig. 2 Cluster of orthologous groups classification. Overall, 47% of all sequences were successfully annotated into 121 categories mainly under more general terms such as “general function prediction only” and “function unknown”, which accounted for 54% of all annotated transcripts

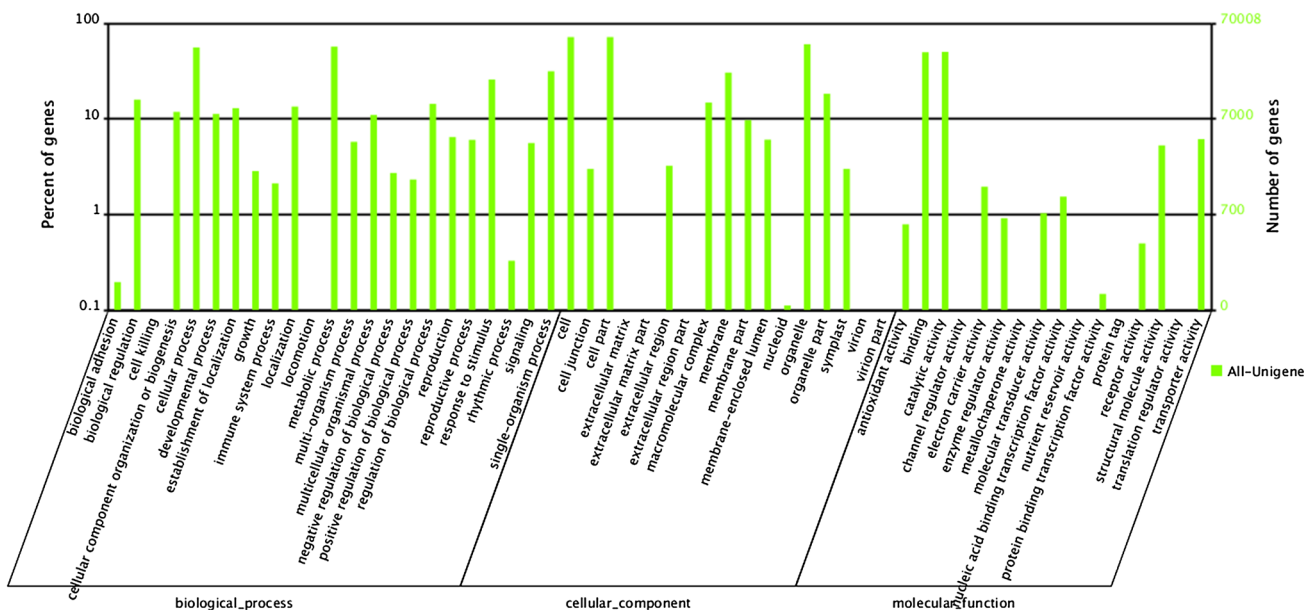


Fig. 3 GO terms histogram. The distribution of GO annotations under “biological process”, “cellular component”, and “molecular function” terms for the 5370 GO entries from 70,008 annotated transcripts

other. The number of sequences similar to the R genes found in *U. decumbens* may be attributed to the fact that the plant was exposed to Al stress in our study. Pathogen attack is a common stressor for plants. According to Atkinson (2012), conditions such as high salinity and acidity are expected to increase the expression of R genes to protect the plant host from enemy attack and environmental stress, thus maintaining its vital functions.

Under the physiological conditions of this study, *U. decumbens* expressed 12 domains and multidomains of proteins related to resistance, which were present either alone within a gene or combined with other domains within the same gene. The most expressed domains were serine-threonine kinases (Ser-Thr Kinase), nucleotide binding sites (NBS) and leucine rich repeats (LRR), as indicated in Table 3.

Discussion

Toledo-Silva et al. (2013) sequenced the tropical grass *Panicum maximum* and presented N50 metric and mean transcript length values (1272 and 942 bp, respectively) similar to ours (Table 1), which were also similar to those of other transcriptomes sequenced using next-generation sequencing from the Poaceae family (Meyer et al. 2012; Xu et al. 2013). Moreover, with respect to reads per library, ploidy levels, and estimated genome size, our results are consistent with other studies that measured differential expression under stress conditions (Li et al. 2015; Xu et al. 2013; Wu et al. 2014).

The difference between the numbers of transcripts obtained by sequencing and sequences actually used for further analysis is due to the adoption of redundancy

Fig. 5 Gene set enrichment analysis. **a** Response to stimulus cluster: the enriched terms from this category are related to processes that occur as a result of the stimulus within or outside the cell or organism, suggesting a change in state or activity due to the presence of aluminum or derived from the chemical response to aluminum toxicity. **b** Biological regulation cluster: any process that modulates a measurable attribute of any biological process, quality or function. The relationship between the terms points to the negative regulation of cell growth, which is an expected outcome of aluminum exposure. **c** Cellular component cluster: any constituent part of a cell, the basic structural and functional unit of all organisms. Both internal and external environment terms were enriched. **d** Legend for arrows and box colors

elimination and sequence complementation steps between and among libraries (Table 1). By systematically comparing different strategies and resources, Duan et al. (2012) optimized the transcriptome assembly of common wheat and found that a scaffolding step significantly reduces redundancy. After optimization, the assembly consisted of four steps: pre-assembly, merging different samples, removal of redundancy, and scaffolding. The resulting assembled transcripts were comparable to Sanger-derived ESTs in terms of both continuity and accuracy.

Sequenced genes identities can be predicted on the basis of the most similar protein deposited and annotated in a protein database (Conesa et al. 2005). The number of annotated sequences in this work for the different databases are summarized in Table 4. Functional annotation provides valuable information, allowing to connect information between gene expression levels and the putative biological role of a transcript, and it can be used to pre-select candidates genes that show linkage to a physiological response for further investigation. With respect to the identification of putative protein functions, we successfully retrieved functional information from 113,918

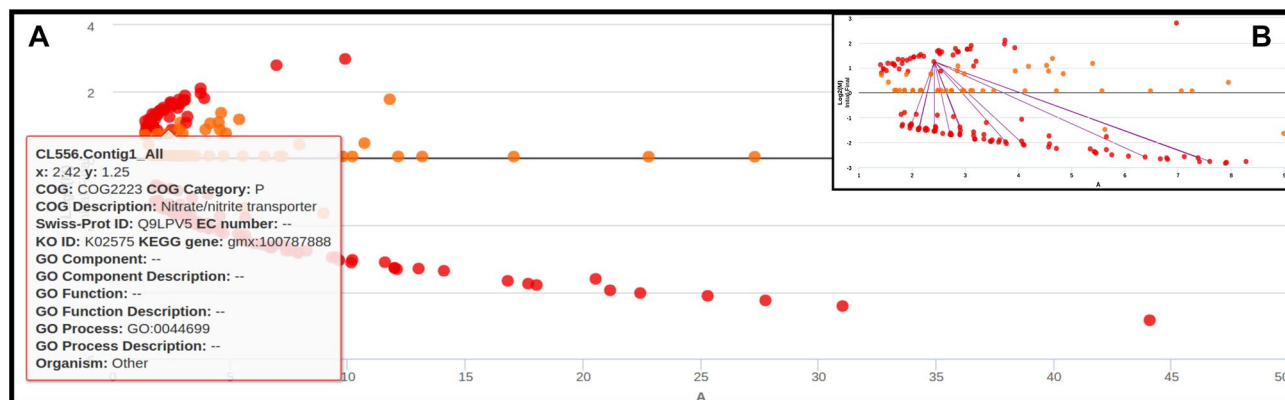


Fig. 4 MA plot. Distribution of top 100 annotated genes up and down regulated **a** in a panoramic view and **b** zoomed out view. Red points represent genes that were classified as differentially expressed, and orange points represent genes that were expressed in just one of

the conditions. X-axis represents the average of the normalized count values, dividing by size factors, taken over all samples, and the y-axis the \log_2 fold change. Genes with same COG category are linked by edges

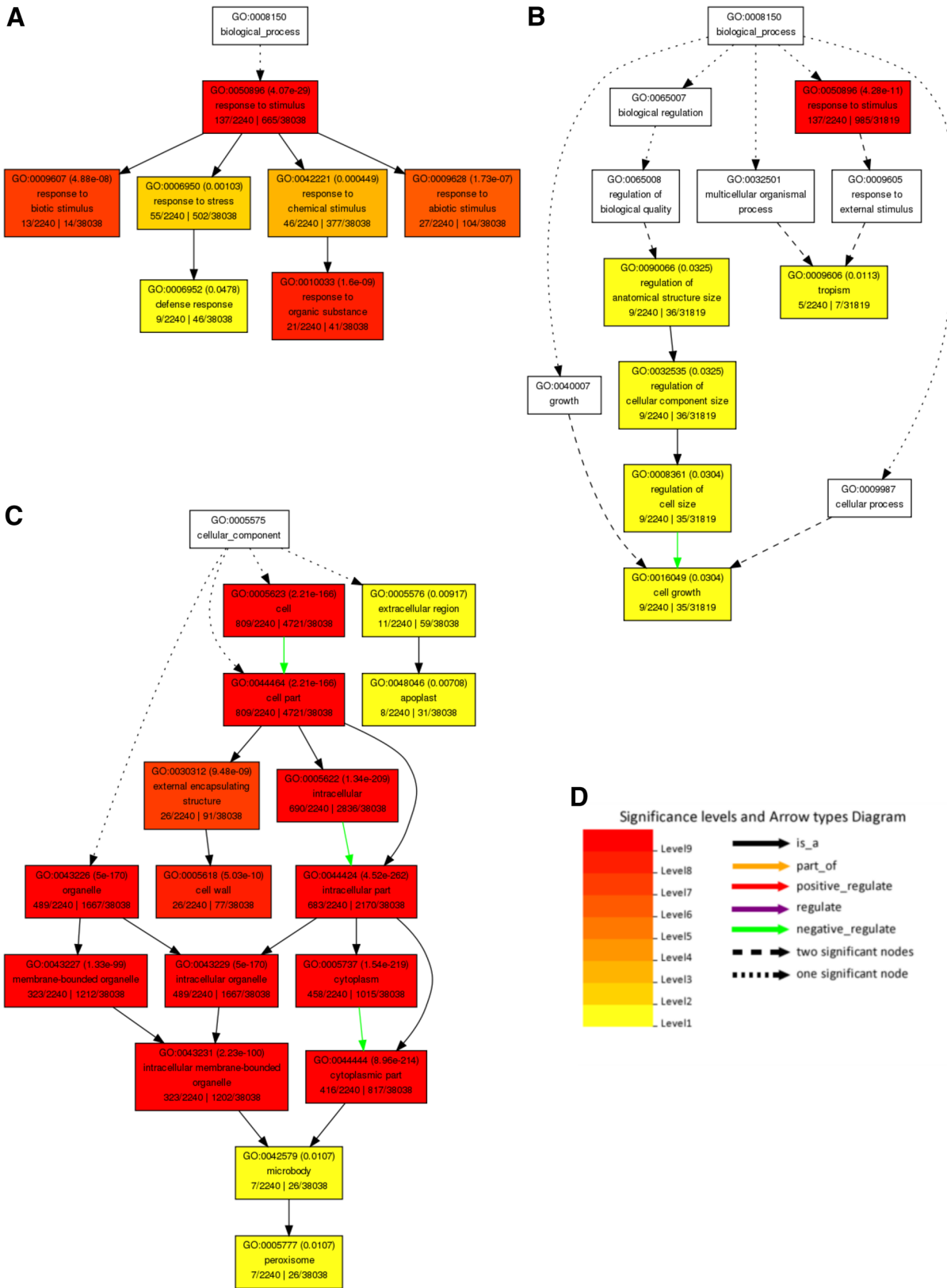
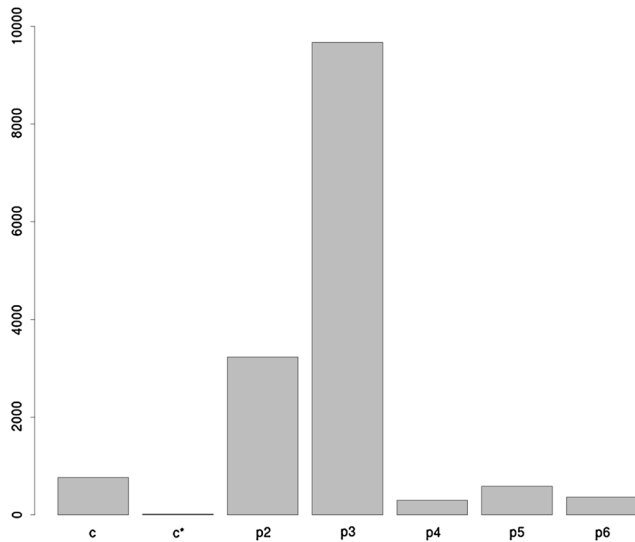
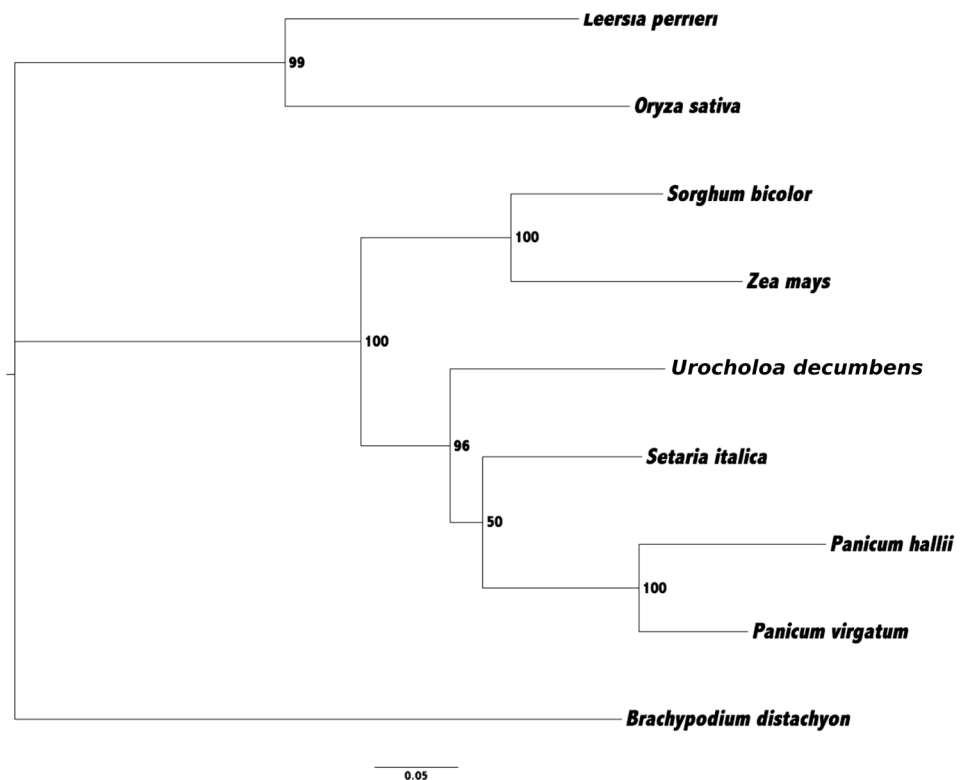


Table 2 Summary of SSR screening

Total number of sequences examined	164,920
Total size of examined sequences (bp)	148,280,694
Total number of identified SSRs	17,375
Number of SSR-containing sequences	14,822
Number of sequences containing more than 1 SSR	2056

**Fig. 6** SSR composition. A bar chart of SSR motif length. Three-nucleotide repetitions accounted for >61% of all SSRs**Fig. 7** Complete genome tree. Species tree based on the concatenation of alignments of 561 ortholog families. Node labels represent bootstrap support (100 replicates)

sequences (69% of all sequences) against NCBI nr, NCBI nt, SwissProt, KEGG (72,749 sequences to 3290 KEGG orthology and 1065 enzyme commission terms), COG (70,008 sequences into 5370 GO terms) databases, with low e-values ($76\% \leq 1e-06$) and high similarity (61% of sequences with >60% identity), mainly with proteins from *S. bicolor* and *Z. mays* (Fig. 5). The variety of distinct group functions, the number of annotated transcripts and the presence of the main metabolic pathways indicate that this study successfully captured the root transcriptional events in response to early AI exposure.

The Ser-Thr Kinase domain was the most abundant in the *U. decumbens* sequences. A study of *Nicotiana benthamiana* demonstrated that genes containing such receptors significantly contribute to resistance against *Phytophthora infestans* (Chaparro-Garcia et al. 2011). The LRR domain was found in 873 transcribed sequences of *U. decumbens*, alone or with other domains. Genes with this domain were observed in *Oryza sativa* L., in which they conferred resistance to the rice blast disease caused by *Magnaporthe oryzae* (Wu et al. 2015). Alone or with other domains, the NBS domain was found in 763 transcripts from *U. decumbens*. The majority of the plant R genes have an NBS and LRR site. These domains are highly conserved through generations and have evolved in different species (Pan et al. 2000).

The high-throughput sequencing of RNA (RNA-seq) provides tools that allow researchers to perform gene expression profiling to identify the transcriptional

Table 3 R gene domains and multidomains from *Urochloa decumbens*

Domain	Total sequences	Percentage (%)
LRR	74	3.02
LRR, Ser-Thr Kinase	349	14.27
MLO	14	0.57
NBS	251	10.26
NBS, LRR	449	18.36
NBS, Ser-Thr Kinase	62	2.53
Ser-Thr Kinase	1055	43.14
Unknown	107	4.37
Ser-Thr Kinase, GNK2	78	3.19
Receptor-Like Kinase	4	0.16
NBARC	1	0.04
NBS, TIR, LRR	1	0.04
TOTAL	2445	100.00

Table 4 Distribution of annotated sequences by database

Database	Annotated sequences
NCBI - NR	103,944
NCBI - NT	80,593
Swiss-Prot	72,794
KEGG	72,749
COG	55,451
GO	70,008
TOTAL	113,918

complexity of biological stress responses. As demonstrated by Delhaize and Ryan (1995), the excess Al causes a rapid (minutes to few hours) inhibition of root growth, resulting in a reduced and damaged root system which limits mineral nutrient and water uptake, extending the root tips exposure until nine hours we expect to be able to capture the early molecular events of the exposure to aluminum through the counting and relativization of the sequence data, thus elucidating how the roots of *U. decumbens* cv. Basilisk first react to Al exposure.

The enrichment of cellular component terms highlights the participation of structural components, such as the cell wall, apoplast (Fig. 5), membrane vesicle. Al binds to the cell wall, particularly to the pectic matrix, and to the apoplastic face of the plasma membrane, which leads to injured cells and the formation of reactive oxygen species (ROS). Increasing physiological, biochemical and molecular evidence suggests that the modification of apoplastic Al binding modulates Al toxicity by reducing interactions between Al and cell wall components, which emphasizes the involvement of cell wall properties in Al resistance (Horst et al. 2010).

The two main mechanisms of Al tolerance in plants include internal detoxification and exclusion from their tissues. The former is related to the ability to capture and enclose Al and form harmless complexes. The strategy involved in the latter is to inhibit aluminum uptake by releasing organic acids that chelate Al (Simões et al. 2012).

Both Al exclusion and internal tolerance mechanisms require the intense participation of plasma membrane transporters (Brunner and Sperisen 2013). Two main transporter families have been reported to play a role in Al tolerance: ABC transporters and MATE-type transporters. MATE transporters are likely responsible for citrate efflux, a strong chelator of Al, and are linked to two major aluminum tolerance quantitative trait loci (QTLs) in maize (Maron et al. 2010). ABC-transporters are involved in various plant systems and largely contribute to the membrane transport of endogenous secondary metabolites in the plant body. Using the NCBI BLAST annotations, we successfully identified 223 DEGs directly related to Al-tolerance mechanisms. Of these transcripts, 38 were annotated as multidrug and toxic compound extrusion (MATE) genes (Yazaki 2006).

COG annotation from DEGs successfully identified 190 transcripts annotated as ABC-type multidrug transport systems. ABC-type transporters are present in vacuolar membranes and have been hypothesized to act in the enclosure of Al, segregating this metal away from sensitive tissues (Huang et al. 2012). *Arabidopsis* mutants for the ALS1 and ALS3 (Al³⁺-sensitive) genes, which encode proteins that belong to the ATP-binding cassette (ABC) transporter superfamily, exhibit significantly increased sensitivity to Al and accumulate increased amounts of Al in the cytosol and nucleus. This finding suggests that these gene products are involved in Al transport across the plasma membrane and subcellular sequestration (Larsen et al. 2007). Moreover, the OsALS1 gene, which encodes a half-sized ABC transporter that is a member of the TAP (transporter associated with antigen processing) sub-group, is also related to Al sequestration in *O. sativa*, mutants of which display significantly increased Al sensitivity but not increased sensitivity toward other metals or low pH (Huang et al. 2012).

By comparing the SSR markers from nine Poaceae species (*O. sativa* L. ssp. japonica, *O. sativa* L. ssp. Indica, *Z. mays*, *S. bicolour*, *Brachypodium distachyon*, *Foxtail millet*, *Moso bamboo*, *Triticum urartu* and *Aegilops tauschii*) Wang et al. (2015) showed that the SSR density of the grass family varies greatly between species. Moreover, the authors found similar numbers of minimum-length SSRs (12 base pairs, bp), a high guanine-cytosine (GC) content (especially in trimers), and hexamers as the most abundant SSR type. In contrast, our results indicated a major percentage of trimers, although also with a high GC content. As demonstrated by Šmarda et al. (2014), Poaceae species exhibit an increased percentage of C/G bases, which

facilitates more complex gene regulation and leads to more stable genetic materials.

Conclusions

This is the first large-scale transcriptomic study of the *Urochloa* genus. The amount of sequencing data generated in this work greatly exceeds the existing resources in public databases and significantly increases the number of reads and annotated transcripts available.

The amount of reads generated, the high number of identified transcripts associated with the diversity of their locations, and the variety of predicted functions demonstrate the success of our approach in capturing the root transcriptome.

The differential gene expression analysis in response to Al stress enabled us to identify genes that are related to Al response in both intracellular and extracellular environments and provided candidates for more detailed assays concerning Al tolerance in *U. decumbens* cv. Basilisk and future assays concerning elucidate molecular mechanisms of aluminum tolerance. The data presented in this manuscript represent an important contribution for the research community interested in plant stress response at a genomic level.

Additionally, the high number of identified molecular markers provides a valuable source of genetic information that will enable downstream analysis for biotechnological purposes and breeding programs.

Availability of supporting data

All the reads resulting from the sequenced libraries used for de novo transcriptome assembly and downstream analysis are deposited on the National Center for Biotechnology Information Short Read Archive (NCBI-SRA) Number SRP071168, Bioproject Number PRJNA314352.

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Compliance with ethical standards

Conflict of interest The authors declare that they have no competing interest.

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