Assessing differential expression profiles and modeling allele-specific expression in leaves of *Saccharum* accessions contrasting in biomass production

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Thesis presented to obtain the degree of Doctor in Science. Area: Genetics and Plant Breeding
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To my parents, Claudia and Valdecir,
and my girlfriend Maria Clara
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If a thing be ordained to another as to its end, its last end cannot consist in the preservation of its being. Hence a captain does not intend as a last end, the preservation of the ship entrusted to him, since a ship is ordained to something else as its end, viz. to navigation.

*St. Thomas Aquinas*
A cana-de-açúcar é uma das mais importantes culturas agrícolas mundiais devido a seus principais produtos - açúcar e álcool -, o reuso de seus subprodutos e a capacidade de inovação de sua agroindústria. Apresenta um potencial para uma produção mais rentável e sustentável, que pode ser obtida pelo desenvolvimento de cultivares de alta produtividade. Por esse motivo, características além do teor de sacarose nos colmos devem ser exploradas. Recentemente, a chamada cana-energia fez com que os programas de melhoramento contemplassem características relacionadas à biomassa, como o conteúdo de fibra e a capacidade de perfilhamento. A variação genética associada a essas características pode ser melhorada pela inclusão de outros acessos de Saccharum, os quais ainda não foram explorados pelos melhoristas. Além disso, os estudos sobre os perfis de expressão gênica em diferentes grupos de genótipos ainda são limitados na literatura. Portanto, objetivou-se a avaliação dos transcriptomas das folhas de dois grupos de genótipos - alta e baixa biomassa - a fim de identificar genes ou alelos potencialmente envolvidos com o conteúdo de biomassa. Para esse objetivo, genótipos foram selecionados pela similaridade fenotípica, independentemente de suas classificações como cultivados ou selvagens. O estudo foi dividido em dois capítulos. No primeiro, o objetivo foi a identificação de genes diferencialmente expressos entre os grupos de biomassa e investigação dos perfis de expressão de genes co-expressos. Os resultados mostraram que a expressão gênica permitiu não só estudar a variabilidade entre os grupos, como também a variabilidade dentro de cada grupo. Apesar da similaridade fenotípica, o grupo de alta biomassa mostrou uma alta variabilidade entre seus acessos, o que resultou em número expressivo de genes diferencialmente expressos, muito maior do que a comparação intergrupo. Genes que codificam a sacarose sintase e proteínas relacionadas à síntese de sacarose foram ligeiramente mais expressas no grupo de baixa biomassa, enquanto que aqueles envolvidos com a síntese de compostos da parede celular foram significativamente menos expressos. Curiosamente, a análise de co-expressão revelou que a expressão de genes relacionados com a fotossíntese foi maior em todos os genótipos híbridos e em Saccharum officinarum. Mostrou-se, também, que diferentes níveis de quantificação possuem certa influência nas considerações biológicas desse tipo de estudo. No segundo capítulo, testou-se a expressão alelo-específica (ASE) em um subconjunto de amostras de Saccharum. Esses acessos - três híbridos, uma S. officinarum e duas S. spontaneum - foram genotipados através da técnica de genotipagem por sequenciamento, seguida das estimativas da ploidia e dosagens alélicas. Modelou-se, para cada polimorfismo, a probabilidade da expressão do alelo de referência por um modelo Beta-Binomial hierárquico, no qual as dosagens alélicas serviram de informação a priori. Os resultados revelaram que ASE afeta parte dos loci avaliados em Saccharum. Entretanto, nenhum termo funcional foi enriquecido com os genes que demonstram ASE. Este estudo foi a primeira visão geral da ocorrência de expressão alelo-específica em múltiplos genótipos de cana-de-açúcar. Ademais, o modelo hierárquico pode ser usado para avaliar ASE em outros organismos de ploidia mista.

Palavras-chave: Saccharum, Transcriptomas, Biomassa, Desbalanço alélico
ABSTRACT

Assessing differential expression profiles and modeling allele-specific expression in leaves of *Saccharum* accessions contrasting in biomass production

Sugarcane is one of the most important crops worldwide due to its main products - sugar and ethanol -, the reuse of byproducts and the innovation capability of the agroindustry. It offers the potential for a more profitable and sustainable production, which can be accomplished by developing high-yielding cultivars. For that reason, traits other than the sucrose content in culms should be explored. The so-called energy cane has recently moved the attention of breeding programs towards biomass-related traits such as fiber content and tillering capacity. The genetic variation associated with these traits can be enhanced with other *Saccharum* accessions that have not yet been explored by breeders. In addition, studies regarding gene expression profiles in diverse groups of genotypes still limited in the literature. Therefore, we aimed to assess the transcriptomes from leaves of two groups of genotypes - high and low biomass - to identify genes or alleles potentially involved with the biomass content. To achieve such goal, genotypes were selected based on their similar phenotypes, regardless of their classification as cultivated or wild. We divided this study into two chapters. In the first chapter, the aim was to identify differentially expressed genes between the biomass groups and to investigate the expression profiles of coexpressed genes. Our results showed that gene expression allowed to study beyond the variability between the contrasting groups, the variability within each group. Despite the phenotypic similarity, the high biomass group showed an impressive variability among its accessions, resulting in many differentially expressed genes (DEGs), much more than the intergroup comparison. Genes coding for sucrose synthase and proteins related to sucrose synthesis were slightly more expressed in the low biomass group, whereas genes involved with the synthesis of cell wall compounds were significantly less expressed. Interestingly, the coexpression analysis revealed that the expression of genes related to photosynthesis was higher in all hybrids and *Saccharum officinarum* genotypes. We also showed that different quantification levels have certain influence on the biological insights provided by this kind of study. In the second chapter, we tested for allele-specific expression (ASE) in a subset of the *Saccharum* samples. These accessions - three hybrids, a *S. officinarum* and two *S. spontaneum* - were genotyped via genotyping-by-sequencing, followed by the estimation of ploidy and allelic dosages. We then modeled, for each polymorphism, the probability of expressing the reference allele using a hierarchical Beta-Binomial model, where allelic dosages served as prior information. Results revealed that ASE affects part of the loci assessed in *Saccharum*. However, any functional term was enriched with genes showing ASE. This study was the first global view of allele-specific expression in multiple genotypes of sugarcane. Furthermore, the hierarchical model can be used to evaluate ASE in other mixed-ploidy organisms.

Keywords: *Saccharum*, Transcriptome, Biomass, Allelic imbalance
1 INTRODUCTION

Sugarcane is an important crop in Brazil since the Portuguese colonization to produce sugar and, for approximately 50 years, to produce ethanol. Recently, data from the Brazilian Sugarcane Industry Association (UNICA) shows that sugarcane is planted in more than 5.5 million hectares in the State of São Paulo (http://www.unicadata.com.br/ - year 2018). According to the National Supply Company (CONAB), the Brazilian sugarcane production in the 2020/21 harvest is expected to increase in comparison with the previous year, reaching roughly 665.1 million tonnes [5]. While the total ethanol production will be reduced by 7.9%, sugar production is estimated to increase by 40.4% (41.8 million tonnes). Progress in the sugarcane industry was partially achieved through breeding high-performance cultivars. Briefly, the sugarcane breeding process relies on crossing parental genotypes, selecting superior genotypes for traits with high variability, then evaluating clones in proper experimental designs for lower heritability traits and, finally, assessing the genotype-environment interactions in competition trials [20]. Breeders have focused on increasing plant productivity to supply the industrial needs of raw material. At the same time, a more effective production in the same cultivable area is desired for a more sustainable agriculture. Scortecci and colleagues [49] stress the importance of leveraging the genetic potential of cultivars to achieve high yields and reduce the natural resources consumed by the plant. Moreover, we should explore not only the variability of sugarcane cultivars, but also from other *Saccharum* species.

Sugarcane is taxonomically classified as belonging to the genus *Saccharum*, subtribe *Saccharinace*, of the Poaceae family. Six species have been studied for understanding the evolution in the genus. Among them, four can be classified as cultivable: *Saccharum officinarum* L., *S. barberi* Jeswiet, *S. sinense* Roxb. and *S. edule* Hassk [40, 55, 39]. The same authors classify the two remaining species as wild: *S. spontaneum* L. and *S. robustum* Brandes & Jeswiet ex Grassl. Due to their proximity and the possibility of intergeneric crossings, *Erianthus*, *Miscanthus*, *Narenga*, *Saccharum* and *Sclerostachya* form the *Saccharum* complex [40, 55, 39]. Historically, the main objective of sugarcane breeding was sucrose accumulation in culms using mostly *S. officinarum* accessions. Later, crossings with *S. spontaneum* were performed to introgress traits related to stress tolerance [53]. The recent development of a group of high-productivity cultivars - energy canes - directed the breeders’ attention to biomass [22, 13, 21]. As stated in studies dating from the 80s [8, 31], energy canes should achieve high yields of both sugar and biomass. The development of such new genotypes demands genetic resources in terms of biomass-related traits, such as fiber content in culms and tillering capacity. Breeding programs can thus benefit from enhanced knowledge about the molecular basis of desired traits, obtained via molecular markers and genomic sequences [7].

The association between genotypic and phenotypic data is not trivial in sugarcane. All *Saccharum* are polyploids showing a large number of chromosomes, which is variable in different accessions of the same species [55, 51]. As a consequence of the interspecific hybridization and successive backcrosses with *S. officinarum*, the modern cultivars have a very complex genome. Most of the basic chromosome architectures (x = 10) are represented by approximately eight *S. officinarum* homologs, *S. spontaneum* chromosomes and a small proportion of recombinants between the two species [51]. During sugarcane breeding, other *Saccharum* species - *S. barberi*, *S. sinense* and *S. robustum* - had a minimum contribution [38, 52]. Multiple strategies were used to unravel its genome sequence [50, 58, 33]. Recently, Garsmeur and colleagues [16] published a mosaic genome assembly of a commercial hybrid; Zhang and colleagues [26] published the sequence of a tetraploid *S. spontaneum* genome; and Souza and colleagues [4] published the gene space assembly of a Brazilian hybrid. However, analyzing the sugarcane genome is still a difficult task when different *Saccharum* accessions are being studied. Approaches using transcriptomes are useful to investigate likely cellular functions of putative genes, aiming to obtain molecular markers from functional genomic regions. Pioneering initiatives paved the way for functional genomics in sugar-
cane. First, Carson and colleagues [43] assessed gene expression in sugarcane leaf rolls using expressed sequence tags (ESTs). Two years later, after assessing the transcriptome of sugarcane leaves, they found genes functionally associated with the control and maintenance of cellular metabolism, transport and response to stresses [41]. Afterwards, researchers in the SUCEST project obtained more than 200 thousand ESTs from different samples [57]. Differentially expressed genes related to cell wall, cellulose and lignin biosynthesis were identified among different stages of culm development via transcriptome profiling [9].

These functional genomics and physiological studies in sugarcane provided evidence of important genes related to sucrose accumulation and synthesis of structural compounds. Along with advances described in the literature for other plants, efforts have also been made to connect genes in pathways to understand carbon partitioning in sugarcane. Wang and colleagues [48] showed the main steps for this process, from sucrose synthesis to its distribution to the sink cells. They showed that after photosynthesis on sugarcane leaves, sucrose is translocated in the phloem and reaches the stem parenchyma cells through both symplast and apoplast. These authors also reported key enzymes for sucrose accumulation: i) sucrose phosphate synthase (SPS) synthesizing sucrose-P from fructose-6-P and UDP-glucose; ii) sucrose phosphate phosphatase (SPP) producing sucrose from sucrose-P; iii) sucrose synthase (SuSy) being responsible for a reversible reaction converting fructose and UDP-glucose to sucrose; iv) cell wall invertase hydrolyzing sucrose into hexoses in the apoplast. There are also other classes of invertases and transporters that participate in transferring hexoses and sucrose into the cellular compartments. In addition to the transport via symplast, hexoses are transported by carriers and resynthesized into sucrose in the cytoplasm. Curiously, Saccharum species accumulate similar levels of symplastic and apoplastic solutes [2]. However, in general, high fiber species - S. robustum and S. spontaneum - show higher percentages of insoluble solids than sucrose-rich Saccharum, which in turn present a higher content of soluble solids [46, 2]. It is worth mentioning that S. spontaneum has a higher content of starch in mature culms to probably meet metabolic demands, serving as a resource for tillering and when the plant is submitted to stress [46].

Attention has been devoted to understand the synthesis of cell wall compounds, as the fibrous part can now be used as raw material by the sugarcane industry. The cell wall can be used in diverse manners, such as a prime source of energy, as feedstock and to develop cellulose-based materials [1]. Regarding the structure, primary and secondary walls of grasses are formed mostly by cellulose, followed by hemicellulose - arabinan- and xylan-derived compounds -, phenolic compounds, pectins, proteins and silica [1, 47]. The composition varies in different developmental stages of the culm. While the hemicellulose content is higher in younger internodes, cellulose is higher in mature internodes [46]. The synthesis of these elements requires the action of enzymes coordinated in different molecular pathways. More than a hundred candidate genes were found to be significantly associated with different fiber composition traits [47]. For cellulose, UDP-glucose from SuSy reaction is used by a complex set of cellulose synthase proteins to synthesize the glucan chain [45, 48]. This is corroborated by the significant association of both SuSy and UDP-glucosyl transferase with cellulose [47]. The biosynthesis of lignin is carried out by many enzymes of the phenylpropanoid pathway. In this pathway, Jardim-Messeder and colleagues [44] defined a core set of genes involved in lignin biosynthesis from the following families: phenylalanine/tyrosine ammonia-lyase, 4-(hydroxy) cinnamoyl CoA ligase, cinnamate 4-hydroxylase, hydroxycinnamoyl CoA shikimate:quininate hydroxycinnamoyltransferase, p-coumaroyl shikimate:quininate 3´-hydroxylase, caffeoyl CoA O-methyltransferase, caffeic acid/5-hydroxyferulic acid O-methyltransferase, 5-hydroxyferulic acid:coniferaldehyde/conifer alcohol 5-hydroxylase, (hydroxy)cinnamoyl CoA reductase and (hydroxy)cinnamyl alcohol dehydrogenase. Authors reported that the expression of genes of the biosynthesis of monolignols have both genotype- and tissue-specificity [46]. High-fiber Saccharum species - S. robustum and S. spontaneum - show more diverse lignin oligomers [46]. The set of 15 phenylpropanoid core genes showed increased expression levels according to culm development [44].
These authors also analyzed the haplotypes of these genes, revealing an uneven distribution in the *S. spontaneum* genome. However, they could identify similar distribution of *cis*-elements in the upstream region of different haplotypes of a gene. Transcription factors can bind to such regions and regulate the expression of members of the phenylpropanoid pathway. In fact, biosynthesis of secondary cell wall can be regulated by myeloblastosis (MYB) and NAC transcription factors, as they are correlated to genes acting on the synthesis of lignin, tricin and hemicellulose [45].

New sequencing technologies, the possibility of assembling transcriptomes *de novo* and the development of statistical methods led to a revolution in the analysis of transcriptomes. The so-called RNA-Sequencing [30] has allowed an increase in the number of characterized sugarcane transcripts, as well as the comparison between contrasting conditions. In 2014, Cardoso-Silva and colleagues [56] assembled the transcriptomes of six cultivars, discovering 5,272 new putative genes not found in the SUCEST database. These authors found genes related to sucrose accumulation and responses to diseases. In the same year, the transcriptomes of the cultivar SP80-3280, accessions of *S. officinarum* and *S. spontaneum* were investigated [10]. These authors showed a high number of *S. spontaneum*-specific transcripts related to stress, signal transduction and transcription factors in sugarcane leaves. They also found that 78.28% of the transcripts were expressed in all genotypes and suggested that major phenotypic differences may be due to reasons other than expression variation at the gene level, such as isoforms, allelic variation and polymorphisms. Later, more than 500 transcripts associated to carbohydrate metabolism and transport were identified in the transcriptome of a high-sucrose cultivar [17].

The advance of sequencing methods has allowed the identification of isoforms, their occurrence in different tissues, development stages or growth conditions. In sugarcane, libraries from different organs were combined: i) first, second and third visible dewlap leaves; ii) immature and mature roots; and iii) the third internode from the top and the third internode from the base [59]. They generated a *de novo* transcriptome using Illumina sequencing on samples of (iii) and the isoform sequencing (Iso-Seq) from Pacific Biosciences on (i), (ii) and (iii) to identify isoforms. The *de novo* assembled transcriptome had a higher percentage of read alignment, more predicted proteins with homology to *Viridiplantae* and allowed the discovery of a larger number of KEGG pathways. Iso-Seq, on the other hand, recovered more complete transcripts, which aligned better to the *Sorghum bicolor* genome [59]. These results indicate the potential of Iso-Seq for comparative analyses.

Gene expression can be quantified after mapping reads to the transcripts from which they were originated. RNA-Sequencing has the potential to capture the dynamism of expressed genes from a population of cells, in a given experimental condition, creating the base for differential expression studies [28, 19, 30]. Gene expression data were also used to compare genotypes with different biomass content, aiming to identify transcripts related to carbon partitioning and to precursors of fiber components. Vicentini and collaborators [54] compared two cultivars showing 4% difference in lignin content. They identified more than 2,000 differentially expressed genes (DEGs), with four main distinct expression profiles and more than 100 groups of genes with similar expression. Among the DEGs, authors reported enrichment of the phenylpropanoid pathway, glutathione-S-transferases, trehalose metabolism, cell-wall proteins, response to biotic stresses and plant hormones. Instead of using clonal replicates of single genotypes to represent a given phenotypic group, Kasirajan and colleagues [32] compared two groups of genotypes with contrasting lignin content. They found DEGs more expressed in the high-fiber genotypes that were present in the phenylpropanoid pathway - lignin precursors - and associated with carbohydrate metabolism. However, by only using elite germplasm these articles exploit little existing variability for fiber content and, consequently, for biomass yield.

There are also other approaches to use the expression data provided by these high-throughput methods. One strategy is not to focus on expression at the gene level, but to look for differentially expressed transcripts and characterize splicing events. A second procedure is to assess the variation in
expression levels among the alleles of a gene. In that case, differences in the expression magnitude of two alleles can indicate allele-specific expression (ASE). This phenomenon can be explained by cis-regulation on promoter regions, frameshift mutations and epigenetic modifications that result on higher expression of one allele [24]. To evaluate ASE, polymorphisms have to be detected and allelic quantification should be obtained from RNA-Seq reads [14, 35, 23]. Then, for each polymorphism, a statistical test can be performed to detect allelic imbalance, by checking for deviations from equivalent expression between the alleles [12, 36]. ASE has been commonly assessed in large scale projects, mostly in human genetics [27, 37, 36, 35]. For example, a higher genic dosage caused by structural variations resulting from tumors was directly associated to increased allelic imbalance [36]. Recently, Lee and collaborators [37] found genes with allele-specific expression related to autism spectrum disorder risk. This approach has been used also in plants [14, 18, 42] and can be explored in other species.

As stated previously, ASE studies jointly use genotypic and expression data, which is feasible for sugarcane. Mancini and collaborators [7] discuss the main advances in sugarcane genetics and genomics. One of the most important is the use of SNPs to estimate the doses of the sugarcane alleles [11]. The high abundance of such markers is important for detecting a large number of polymorphisms, which are used to build genetic maps, discover QTLs and genomic regions associated with a given trait. It also opens the possibility for integration with expression data. A diverse set of Saccharum accessions was established in the Federal University of São Carlos (UFSCar), where researchers of the sugarcane breeding program laid out the Brazilian Panel of Sugarcane Genotypes [29, 3]. It is composed by 254 genotypes, representing wild species, cultivars with historic relevance and more recent cultivars. Some authors have already benefited from the genotyping of the panel [15, 29, 11, 3]. Using quantitative genotyping pipelines to obtain SNPs, the relative allelic proportions can be also estimated in this complex crop [11, 25, 34]. Then, the combination of such data with RNA-Seq provides enough information to evaluate ASE in sugarcane. However, a careful examination of the data is needed, as biases in the procedures - mapping and genotyping - can result in false ASE [24].

In this context, we point that it is feasible to understand, at the transcript level, differences between groups of accessions contrasting in their biomass content. In addition, investigating allelic imbalance can provide complementary results to the conventional analyses of gene profiles [36]. We explored gene expression data from leaves of twelve Saccharum accessions, phenotypically clustered in high- and low-biomass groups. First, we aimed to explore the variation between and within the groups in terms of differential gene expression. Next, we investigated the extent to which ASE occurred in both wild and cultivated accessions. We present and discuss our main findings regarding these objectives in two thesis chapters. The first chapter contains the investigation of differential gene expression, which was published in BMC Genomics [6]. We kept the integrity of all sections of this manuscript, including all the main and supplementary information. The second chapter focuses on the development of a model to test for ASE in complex polyploids such as sugarcane. It is also organized as a manuscript to be submitted.

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### 2 Differential Expression in Leaves of *Saccharum* Genotypes Contrasting in Biomass Production Provides Evidence of Genes Involved in Carbon Partitioning

#### Abstract

**Background:** The development of biomass crops aims to meet industrial yield demands, in order to optimize profitability and sustainability. Achieving these goals in an energy crop like sugarcane relies on breeding for sucrose accumulation, fiber content and stalk number. To expand the understanding of the biological pathways related to these traits, we evaluated gene expression of two groups of genotypes contrasting in biomass composition.

**Results:** First visible dewlap leaves were collected from 12 genotypes, six per group, to perform RNA-Seq. We found a high number of differentially expressed genes, showing how hybridization in a complex polyploid system caused extensive modifications in genome functioning. We found evidence that differences in transposition and defense related genes may arise due to the complex nature of the polyploid *Saccharum* genomes. Genotypes within both biomass groups showed substantial variability in genes involved in photosynthesis. However, most genes coding for photosystem components or those coding for phosphoenolpyruvate carboxylases (PEPCs) were upregulated in the high biomass group. *Sucrose synthase* (SuSy) coding genes were upregulated in the low biomass group, showing that this enzyme class can be involved with sucrose synthesis in leaves, similarly to *sucrose phosphate synthase* (SPS) and *sucrose phosphate phosphatase* (SPP). Genes in pathways related to biosynthesis of cell wall components and *expansins* coding genes showed low average expression levels and were mostly upregulated in the high biomass group.

**Conclusions:** Together, these results show differences in carbohydrate synthesis and carbon partitioning in the source tissue of distinct phenotypic groups. Our data from sugarcane leaves revealed how hybridization in a complex polyploid system resulted in noticeably different transcriptomic profiles between contrasting genotypes.

Keywords: Sugarcane; Gene expression; Transcriptomics; RNA-Seq; Polyploid.

#### 2.1 Conclusion

This work presented a broad view of the expression of many coding genes in sugarcane leaves of different genotypes. With regard to cell wall, most genes were upregulated in the high biomass group, but in general with low average expression levels. On the other hand, highly expressed genes involved in sucrose synthesis were upregulated in hybrids and *S. officinarum* genotypes. These results agree with current knowledge about the partitioning of carbohydrate to sucrose storage and maintenance of plant structure and metabolism in wild genotypes and modern cultivars. In addition, our research shows that investigating expression profiles in wild genotypes can enhance the understanding of genes selected through domestication and breeding. Expression profiles in other plant parts of wild and cultivated accessions are needed to provide knowledge about the action of the genes involved in carbohydrate metabolism and biomass production. Our data from sugarcane leaves revealed how hybridization in a complex polyploid system resulted in noticeably different transcriptomic profiles between contrasting genotypes.
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3 A HIERARCHICAL BAYESIAN MODEL TO ASSESS ALLELE-SPECIFIC
EXPRESSION IN MIXED-PLOIDY SPECIES REVEALS EXPRESSION BIASES IN
SUGARCANE

Abstract

Allele-specific expression (ASE) represents differences in the magnitude of expression between alleles of the same gene. Allelic imbalance in diploids occurs if the ratio of expression between both alleles shows deviations from the expected equivalent expression. However, this is not straightforward for polyploids, especially autopolyploids, as knowledge about the dosage of each allele is required for accurate estimation of ASE. This is the case for the genomically complex *Saccharum* species, characterized by high levels of ploidy and aneuploidy. We propose a model to test for allelic imbalance in *Saccharum* that can be easily expanded to other polyploids. As a test case we used genotyping data and RNA-Sequencing libraries from leaves of six sugarcane accessions. We used a hierarchical Beta-Binomial model to test if allele expression followed the expectation based on genomic allele dosage. The doses of the alleles were used in a prior Beta distribution for modeling the proportion of the reference allele from RNA counts. This proportion was then used in a Binomial distribution to model the number of RNA-seq reads showing this allele. We used the Bayesian Markov chain Monte Carlo procedure to draw samples from the *a posteriori* distribution. We called a polymorphism as showing ASE when the relative genomic dose was outside the highest density interval of the posterior distribution in a certain genotype. Part of the genes evaluated in each accession showed ASE and were related to a broad range of processes, mostly associated to the general metabolism, organelles, responses to stress and responses to stimuli. In addition, the frequency of genes with ASE in high-level functional terms was similar among the genotypes. Because the highest frequencies of ASE occurred in sugarcane hybrids, we fancy some influence of the interspecific hybridization in these genotypes. Although the number of polymorphisms we evaluated is still somewhat limited, our study is the first to assess genome-wide ASE in a high- and mixed-ploidy system using estimated doses of the alleles.

Keywords: Allelic imbalance; Polyploid; Allele dosage; Bayes; *Saccharum*

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4 CONCLUSIONS

In this thesis, we aimed to investigate differences among *Saccharum* genotypes phenotypically contrasting in their biomass content. In the first chapter we assessed gene expression profiles of twelve sugarcane genotypes grouped into high and low biomass groups. The gene expression data correctly represented the difference between the groups and revealed substantial variability among the high biomass accessions. The groups showed significant differences in the expression of genes involved in carbon partitioning, mostly sucrose synthesis and degradation. Within the groups we could identify the enrichment of defense and carbohydrate-related terms. In addition, we explored the expression and co-expression profiles of groups of genes that were members of pathways of interest. Finally, we also showed how expression profiles at the transcript level can bring new insights when assessing differences between the biomass groups.

We devoted the second chapter to investigate if genes showing allele imbalance could be related to distinct functional processes. As we aimed to investigate whether alleles were expressed accordingly to their estimated dosages, we proposed a model to account for prior knowledge of this information. We used a hierarchical Bayesian approach to go from a prior distribution of the allele proportion, based on genotyping information, to a posterior considering the relative expression of the allele. Our results reveal that allele-specific expression affects part of the investigated loci in *Saccharum* genotypes. However, we could not find clear functional patterns among genes showing allele-specific expression. Despite the innate limitations of the genotyping-by-sequencing approach, we successfully developed and applied a model to drive insights about allele-specific expression in the complex polyploid sugarcane.