Genome-wide differences in gene expression and alternative splicing in developing embryo and endosperm, and between F1 hybrids and their parental pure lines in sorghum

Meishan Zhang1 · Ning Li2 · Weiguang Yang1 · Bao Liu2

Received: 3 June 2021 / Accepted: 20 September 2021 / Published online: 30 November 2021 © Springer Nature B.V. 2021

Abstract

Key message Developing embryo and endosperm of sorghum show substantial and multifaceted differences in gene expression and alternative splicing, which are potentially relevant to heterosis.

Abstract Differential regulation of gene expression and alternative splicing (AS) are major molecular mechanisms dictating plant growth and development, as well as underpinning heterosis in F1 hybrids. Here, using deep RNA-sequencing we analyzed differences in genome-wide gene expression and AS between developing embryo and endosperm, and between F1 hybrids and their pure-line parents in sorghum. We uncover dramatic differences in both gene expression and AS between embryo and endosperm with respect to gene features and functions, which are consistent with the fundamentally different biological roles of the two tissues. Accordingly, F1 hybrids showed substantial and multifaceted differences in gene expression and AS compared with their pure-line parents, again with clear tissue specificities including extents of difference, genes involved and functional enrichments. Our results provide useful transcriptome resources as well as novel insights for further elucidation of seed yield heterosis in sorghum and related crops.

Keywords Gene expression · Alternative splicing · Imprinting · Development · Heterosis · Sorghum bicolor

Introduction

Sorghum (Sorghum bicolor L.) ranks fifth in global cereal production, and its seeds provide important resources for food, feed and biofuel worldwide (Mace et al. 2013). Owing to its relatively small genome (730 Mb), sorghum has also been used as a model species for genetic and genomic studies in sugarcane and other C4 grasses (Paterson et al. 2009). In cereal crops, endosperm and embryo are major components of seeds and hence are direct determinants of yield. As in all angiosperms, sorghum seed development is initiated following double fertilization, which gives rise to diploid embryo and triploid endosperm, respectively, with the two tissues exhibiting dramatic differences in many respects. The main function of embryo is to perpetuate genetic information to the next generation, while that of cereal endosperm is mainly to nourish the developing embryo and germinating seedling (Lu et al. 2013; Xin et al. 2013). In flowering plants, developing endosperm is the primary tissue of imprinted gene expression, i.e., unequal expression of maternal and paternal alleles depending on parent of origin (Gehring and Satyaki 2017; Zhang et al. 2011). In a previous study, we identified 101 sorghum imprinted genes including 85 maternally expressed imprinted genes (MEGs) and 16 paternally expressed imprinted genes (PEGs) from 14 DAP (days after pollination) sorghum endosperm based on RNA-sequencing (Zhang et al. 2016).

Hybridization is a prominent process in the evolution, domestication and genetic improvement of sexually reproducing crop species. Heterosis or hybrid vigor results from genome-wide interactions between paternal and maternal alleles and de novo regulatory changes in the F1 hybrids (Birchler et al. 2010; Huang et al. 2016). Crop growth and
development are regulated by complex global gene expression networks. Gene expression patterns in F1 hybrids can be divided into two broad categories, i.e., additive and non-additive expression, compared with that of the pure line parental average. Nonadditive gene expression is defined as expression of a gene in a hybrid, which is significantly deviated from the average value of the parental lines (mid-parent value), and which has been shown as significantly associated with heterosis (Paschold et al. 2012). The proportions of genes showing nonadditive expression differ greatly across species, tissues and organs, experimental procedures, and surveyed gene sets (Springer and Stupar 2007). In recent years, several studies have used RNA-seq to analyze globally differentially expressed genes (DEGs) between a hybrid and its parents, provided new insights into the genetic mechanisms of heterosis (Chen et al. 2018; Paschold et al. 2012; Zhang et al. 2017a). For example, RNA-seq analysis of the primary root of maize inbred lines B73 and Mo17 and their reciprocal F1 hybrids revealed that 42–55% of all expressed genes were differentially expressed between at least one of the parents and one of the hybrids, and about 10% of expressed genes were nonadditively expressed in both hybrids (Paschold et al. 2012). A transcriptome analysis between super-hybrid rice and its parents of young panicles identified a total of 5910 DEGs, which were significantly enriched in carotenoid biosynthesis and hormone signal transduction pathways (Chen et al. 2018).

Alternative splicing (AS) is another common mechanism to cause changes in gene expression as well as transcript diversity. AS is a process occurring in all higher eukaryotes whereby the same precursor mRNA generates multiple transcripts with variable quantities. AS is a major mechanism to achieve proteomic diversity with defined gene numbers (Asth 2004; Nilsen and Graveley 2010; Song et al. 2019). In human, more than 95% of multiexonic genes are alternatively spliced (Syed et al. 2012). According to recent studies, the occurrence of AS varies tremendously among plant species, ranging from 37 to 63% in intron-containing genes. At least some of these variations can be attributed to differences in depth of the RNA-seq data, technology platforms, and statistical analyses (Mei et al. 2017; Satyawan et al. 2017; Shen et al. 2014; Zhang et al. 2019). Meanwhile, extensive tissue-specific, developmental stage-specific splicing patterns, or altered AS behaviors, induced by environmental stresses also contribute to the diversity of AS (Mei et al. 2017; Thatcher et al. 2014; Wang et al. 2018). A previous study based on transcriptomic data indicated that 18,741 (45%) and 13,327 (38.5%) of expressed genes were alternatively spliced in maize and sorghum, respectively, but the proportions of different splicing events varied among tissues (Wang et al. 2018). In maize, 50.7% of multiexonic genes were alternatively spliced and 7.0% of AS events were differentially regulated between 9 DAP (days after pollination) embryo and endosperm, and a higher frequency of tissue-regulated events was detected in embryo than in endosperm (Lu et al. 2013). A pairwise analysis of different maize tissues showed that embryo had the highest number of AS events when compared with the rest 14 tissues (Thatcher et al. 2014). Intron retention (IR) is the most frequent AS event in plants, whereas exon skipping (ES) accounts for the most common event in mammalian species (Ast 2004; Nilsen and Graveley 2010; Syed et al. 2012). Some alternatively spliced transcripts play a significant role in development, stress responses and tissue-specific physiology (Lin and Zhu 2021; Liu et al. 2018; Mei et al. 2017). In contrast, most splice variants may not be functional, and degraded by RNA surveillance mechanisms such as the nonsense-mediated mRNA decay (NMD) pathway (Chaudhary et al. 2019; Chen et al. 2019; Satyawan et al. 2017). In some cases, degradation of AS isoforms through the NMD pathway serves to regulate the concentration of transcripts in the cells (Drechsel et al. 2013).

Although extensive studies have been conducted to identify AS events in different plant species, tissues and environmental conditions, little attention has been paid to AS events in crop plant imprinted genes at genome-wide scale. In a previous study, we compared the variant splicing differences in maize for several imprinted genes between embryo and endosperm in a set of pre-selected genes, and found that more IR events occurred in embryo than in endosperm (Zhang et al. 2018). In this study, based on genome-wide comparative transcriptome analyses, we aimed to identify differentially expressed genes (DEGs) in developing embryo and endosperm tissues of a pair of reciprocal sorghum F1 hybrids and their parental pure lines. We also performed AS analysis, including the identification and characterization of AS events, differential splicing between tissues and among genotypes, and AS features of sorghum imprinted genes. Experimental validation as well as gene ontology enrichment analyses were also conducted.

Materials and methods

Plant tissue collection

The sorghum (Sorghum bicolor L.) parental pure lines, BTx623 (B) and Y49 (Y) were grown in the field at Jilin Agricultural University experimental farms near Changchun. A pair of reciprocal inter-strain F1 hybrids, BY (BTx623 as female pollinated with Y49) and YB (Y49 as female pollinated with BTx623) were produced by manual pollination. Embryo and endosperm samples were dissected manually from the 14 days after pollination (DAP) immature seeds. Embryos were washed three times with RNA-free water to remove possible endosperm contamination on the surface.
of embryo. Materials were quickly frozen in liquid nitrogen and store in – 70 °C until RNA extraction. Each sample was collected from at least five different plants.

**RNA isolation**

Embryo and endosperm samples from 4 genotypes in biological triplicate were used for RNA isolation. The total RNA was isolated using Plant RNA Reagent (Invithogen, #12322-012 http://www.lifetechnologies.com) and subsequently purified with QIAGEN RNeasy Plus Mini kit (#74134) according to the manufacturer’s instructions.

**RNA-seq data generation and read alignment to the reference sorghum genome**

RNA-seq data of endosperm was downloaded from SRA, the accession number was PRJNA302593, which we uploaded 2016. For the embryo, the Agilent 2100 BioAnalyzer (Agilent Technologies), and Nanodrop were used to measure quality and quantity of each RNA sample for library construction. The library construction, cluster generation and HiSeq 2500 sequencing were conducted in accordance with the standard protocols. All samples were treated and sequenced as parallel experiments. Adapters and low-quality reads were removed from raw sequences by using fastq_quality_filter from FASTX-Toolkit (http://hannonlab.cshl.edu/fastx_toolkit/), with parameters ‘-p 80 -q 20’, leaving a total of 238.67 million 150 bp paired-end clean reads (71.6 Gb clean data). Reads were mapped to the sorghum reference genome (https://phytozome.jgi.doe.gov/pz/portal.html) by the HISAT2 under default parameter. The raw data information and mapping efficiency are given in Table S1.

**Gene expression estimation and putative transcript assembly**

We used StringTie (Pertea et al. 2016) to estimate the GTF file for each sample with default parameters, and then merged all GTFs into one final GTF with StringTie Merge. Transcripts were normalized using the transcript per million base pairs (TPM). HTSeq (Anders et al. 2015) was used to count reads per gene with parameters “htseq-count -f bam -r pos -s no -q -”, and DESeq2 (Love et al. 2014) was used to identify the differentially expressed genes (DEGs) and transcripts.

**Identification of AS events**

We identified the AS events using the ASTALAVISTA program (Foissac and Sammeth 2007). The different types of AS events were analyzed as previously described (Trincado et al. 2018).

**Reverse transcription and experimental validation with Semi-quantitative RT-PCR**

About 800 ng total RNA from each sample was used for cDNA synthesis using a Reverse Transcriptase kit (TransGen Biotech) following the manufacturer's instructions. The PCR primers sets that flank the differentially spliced region are used to amplify splice variants from cDNA prepared from different samples, with three biological replicates. The primer information is listed in Supplementary Information, Table S3. PCR was performed using rTaq DNA polymerase (TAKARA) in 20 µl reaction volume. The PCR profile was: initial denaturation at 94 °C for 2 min followed by 25–32 cycles (according to the primer pair) at 94 °C for 40 s, 55–60 °C (according to the primer pair) for 40 s, 72 °C for 40–60 s (according to the PCR products length), and a final extension at 72 °C for 8 min. The amplified products were visualized in a 1.8 % agarose gel stained with ethidium bromide. The sorghum actin gene (Genbank accession X79378) was used as an internal reference gene.

**Gene ontology (GO) analysis**

GO term enrichment analysis was performed using the hypergeometric statistic (R function phyper). GO categories with FDR corrected p value (q value) < 0.05 were considered as enriched.

**Results**

**Assessment of the RNA-seq data**

We performed high-throughput RNA-seq using the Illumina HiSeq 2500 platform for embryo and endosperm tissues dissected from sorghum seeds at 14 DAP (days after pollination) of four genotypes (two parental pure lines BTx623 [B] and Y49 [Y], and their reciprocal F1 hybrids, BY and YB). We obtained a total of 487.43 million 150 bp paired-end clean reads which were comprised of 77.9 Gb of high-quality raw data (Table S1). The sequence reads were aligned to the BTx623 v3 reference genome (https://phytozome.jgi.doe.gov/pz/portal.html) by HISAT2. An overview of the distribution of the aligned reads along each of the 10 sorghum chromosomes showed that, as expected, most of the reads were located in the chromosome arm regions, while transposable elements (TEs) were mainly aggregated to gene-poor, pericentromeric regions (Fig. 1). Moreover, approximately 80% of the mapped reads were in exonic regions and only a small number of the reads were mapped to intronic and non-genic regions (Table S1). The aligned reads located in exonic and intronic regions were used to detect gene expression and alternative splicing (AS) for each
We assembled RNA-seq alignments into potential transcripts for each sample with StringTie. Gene and transcript expression levels were also estimated by stringtie, and measured as transcript per million base pairs (TPM). The cutoff value for determining the expressed genes or transcripts was TPM > 0.5.

We found that although most genes were commonly expressed in both tissues of all genotypes, transcript isoform abundance seems higher in embryo than in endosperm, consistent with results in a maize transcriptome analysis (Lu et al. 2013). We detected a total of 40,175 genes expressed in the developing sorghum seed transcriptome, with 38,205 and 21,331 genes expressed in embryo and endosperm, respectively, which corresponded to 60.5–86.5% of the annotated genes, producing 72,912 and 43,461 transcripts, with 19,361 (48.2%) genes being commonly expressed in both tissues (Table S2). In addition, we detected expression of 15,091 and 2889 unannotated genes, which produced 43,077 and 21,516 transcripts in sorghum embryo and endosperm, respectively. Of note, we found that the total numbers of expressed genes in F1 hybrids were higher than those in the parental lines in both tissues (Table S2).

**Differentially expressed genes (DEGs) between embryo and endosperm, and between hybrids and parents**

Differential expression analyses between the tissues and among the genotypes were calculated by DESeq2. Tissue-specific expressed genes (i.e., genes expressed in either embryo or endosperm but not in both) or tissue-preferential expressed genes (i.e., genes whose expression level in one tissue was significantly higher than in the other) were categorized based on the following criteria: (i) FDR corrected p value (q value) was less than 0.01; (ii) genes with > 10 reads in one tissue and ≤ 1 read in another are defined as tissue-specific; (iii) genes showing > 2-fold differences in expression level are defined as tissue-preferential. According to these criteria, we identified 1459 (3.8%) embryo-specific genes, 12,779 (33.4%) embryo-preferential genes, 2307 (10.8%) endosperm-specific genes and 3847 (18.0%) endosperm-preferential genes, respectively (Fig. 2a, b; Supplementary Dataset). Clearly, a greater proportion of the genes were specifically or exclusively expressed in endosperm than in embryo, whereas the proportion of
preferentially expressed genes was 2.8 times more in embryo than in endosperm (Fig. 2a). However, for imprinted genes (defined as ≥ 5-fold differences in read counts between parental alleles) (Zhang et al. 2016), the great majority is endosperm-specific or -preferential, with embryo-preferential genes accounting < 10% of the total number of imprinted genes, and none imprinted gene showed embryo-specific expression (Fig. 2a).

Next, we analyzed distribution of the four types of DEGs (embryo-specific genes, embryo-preferential genes, endosperm-specific genes and endosperm-preferential genes) in the different genotypes, including a pair of reciprocal F1 hybrids (BY and YB) and their two parental pure lines (B and Y) (Fig. 2b, c). We found that tissue-specific expressed genes were less shared by the genotypes, i.e., they are more prone to showing genotype-specific expression. Approximately 19 and 25% of the embryo- and endosperm-specific genes, respectively, were common across all four genotypes. Additionally, ~ 50 and ~ 35% of the embryo- and endosperm-preferential expressing genes, respectively, were shared by all four genotypes (Fig. 2b). Notably, we found that the numbers of embryo-specific expressing genes in hybrids were significantly higher than those in their parents, whereas the other three types of between-tissue DEGs showed similar gene expression between hybrids and parental lines (Fig. 2b). Finally, there were more genes expressed at a higher level in embryo than in endosperm (Fig. 2c), which is consistent with a previous study in maize (Lu et al. 2013).

**Additive and nonadditive gene expression patterns in embryo and endosperm of F1 hybrids**

To determine the relative expression levels of genes in the sorghum reciprocal F1 hybrids (BY and YB) as compared to their parental pure lines (B and Y), and to identify genes displaying expression levels that deviate from additivity in the hybrids, we analyzed the RNA-seq data and identified nonadditively expressed genes in the hybrids. Here, nonadditive gene expression is defined as expression level in a hybrid that is significantly deviated from the averaged parental value, i.e., mid-parent value \[ MPV, \text{ which in embryo } = \frac{M + P}{2}, \text{ in endosperm } = \frac{2M + P}{2}, \text{ q value } < 0.05 \].

In embryo, no significant differences between the reciprocal hybrids with respect of the proportions of genes showing additive vs. nonadditive expression were detected; both hybrids showed a predominant proportion of additive expression (99.5%) and very small proportion of nonadditive expression (0.5%) (Fig. 3). In contrast, in endosperm, the reciprocal F1 hybrids showed clear differences in the proportions of genes showing additive vs. nonadditive expression. Moreover, the proportions of nonadditively expressed genes in endosperm were markedly greater than those in embryo, with 10.5 and 7.2% genes showing nonadditive expression.
in endosperm of the reciprocal hybrids, BY and YB, respectively (Fig. 3). This suggests the influence of cytoplasm and other maternal effects were significantly exacerbated by relative nuclear genome dosage as the triploid endosperm has a 2:1 maternal vs. paternal nuclear genome contribution.

**Differential alternative splicing (AS) between tissues and among genotypes**

The RNA-seq data also enable global analysis of dynamic changes in alternative splicing (AS) patterns. We investigated spliced events in types of genes, between tissues and among the four genotypes using the algorithm ASTALAVISTA (Foissac and Sammeth 2007). Overall, of the 40,175 total expressed genes, 23,200 (57.8%) were multixonic genes, and 43.2% of these genes were alternatively spliced, resulting in 29,373 AS events in total (Table 1; Supplementary Dataset). We found the overall AS frequency is slightly higher than that reported previously for sorghum (Min et al. 2015; Wang et al. 2018), but similar to those reported in maize (Thatcher et al. 2014) and rice (Lu et al. 2010), and significantly lower than other plant species, especially Arabidopsis and soybean, in which AS is reported to occur in > 60% of multixonic genes (Marquez et al. 2012; Shen et al. 2014). Specifically, 43.6 and 41.1% of expressed multixonic genes were alternatively spliced and showing 27,708 and 17,489 AS events in sorghum embryo and endosperm, respectively (Table 1). Among these, a total of 13,724 AS events occurred in both tissues but embryo-specific (occurred in embryo but not in endosperm) AS events (13,984) were 3.7 times more than endosperm-specific (occurred in endosperm but not in embryo) AS events (3765) (Fig. S1). Then, we analyzed AS events, in separation, for the four types of the DEGs and of imprinted genes. We found that 24.6 and 56.2% of embryo-specific and –preferential genes were alternatively spliced respectively, however, the ratios were much lower in both endosperm-specific and –preferential genes than in embryo DEGs, showing only 14.5 and 25.7%, respectively (Table 1). Additionally, the frequency of AS in imprinted genes was 46.3%, which was higher than those of total expressed genes and endosperm DEGs (Table 1).

Next, we projected these comparisons with emphasis on genotype differences in both tissues. Results showed that (i) 11,173 and 3976 events were conserved among all four genotypes in embryo and endosperm, respectively; (ii) compared with parents, 5270 and 4267 AS events were unique to embryo and endosperm of at least one hybrid, and 944 and 452 of both hybrids; (iii) compared with parents, 12,083 and 9423 AS events were missing from embryo and endosperm.

**Table 1 AS events in sorghum embryo and endosperm**

|                        | Expressed genes | Expressed multixonic genes | AS genes (% of multixonic genes) | AS events of multixonic genes |
|------------------------|-----------------|---------------------------|----------------------------------|-------------------------------|
| Total                  | 40,175          | 23,200                    | 10,031 (43.2)                    | 29,373                        |
| Embryo                 | 38,205          | 22,037                    | 9599 (43.6)                      | 27,708                        |
| Endosperm              | 21,331          | 17,941                    | 7379 (41.1)                      | 17,489                        |
| Embryo-specific        | 1459            | 1108                      | 273 (24.6)                       | 465                           |
| Embryo-preferential    | 12,779          | 11,692                    | 6572 (56.2)                      | 20,347                        |
| Endosperm-specific     | 2307            | 1541                      | 223 (14.5)                       | 382                           |
| Endosperm-preferential | 3847            | 2843                      | 731 (25.7)                       | 1512                          |
| Imprinted genes        | 101             | 80                        | 37 (46.3)                        | 65                            |

\[Fig. 3\] Additive and nonadditive gene expression between F1 hybrids and parental lines in sorghum embryo and endosperm. The numbers above individual columns present numbers and proportions of genes detected in the corresponding tissues and genotypes relative to total expressed genes. The statistical significance was determined using p-value (false discovery rate, q value<0.05). BY (BTx623 × Y49) and YB (Y49 × BTx623) are reciprocal F1 hybrids, while B (BTx623) and Y (Y49) are their parental pure lines.
from at least one hybrid, and 5653 and 4406 from both hybrids (Fig. 4a).

We further investigated the different AS types in tissue-specific or tissue-preferential expressing genes, as well as in imprinted genes. The seven AS modes were illustrated (Fig. 4b). Overall, the four basic AS types accounted for ~79% of the total AS events in both tissues, and the distribution of different types of AS events was also similar between embryo and endosperm. Specifically, as described in other plant species, we found that intron retention (IR) accounted for the most abundant AS type in both tissues (~25%), though it was only slightly higher than the second type, alternative 3′ splicing site (A3SS). A similar trend was observed in 20 DAP sorghum BTx623 embryo and endosperm (Wang et al. 2018). Exon skipping (ES) events varied in different plant species from 3% in Arabidopsis (Marquez et al. 2012) to 25% in rice (Zhang et al. 2010) and 33% in maize (Thatcher et al. 2014). The number of ES events in our data fell in the middle, accounting for ~17% of the total AS events. Alternative 5′ splice site (A5SS) was the least frequent mode of the four main AS types (~13%) in both tissues (Fig. 4c). We also calculated the rare AS types. For example, both the skipping two exons, “ES1 + ES2” and “A5SS or A3SS” counted ca. ~2.5% of the total AS events. The mutually exclusive exon (MXE) AS type, which has been detected in rice (Lu et al. 2010; Zhang et al. 2010), but not in Arabidopsis (Marquez et al. 2012), accounted for ~0.6% of the AS in sorghum, ranked the least AS type in our data (Fig. 4c). This is consistent with the results in maize (Lu et al. 2013). For those complex AS models, we categorized them all together as the type of “others”, which likely resulted from different combinations of the four basic types or arose from different mechanisms (Syed et al. 2012). These complex models accounted for approximately 17% of the total AS events in embryo and endosperm of sorghum (Fig. 4c).

For the DEGs, no significant differences in the proportion of AS patterns were observed in tissue-preferentially expressed genes of both embryo and endosperm compared to all genes from the sorghum genome. However, the tissue-specific genes in both tissues showed obviously different splice patterns. Specifically, (i) in embryo-specific genes, IR was increased by 16%, whereas in endosperm-specific genes, IR was decreased by 28%, relative to those of the total expressed genes; (ii) the exon skipping type did not change in embryo-specific expressing genes but was increased by 27% in endosperm-specific expressing genes; (iii) in both embryo- and endosperm-specific genes, the “others” AS type was also decreased compared to the total expressed genes (Fig. 4c).

Interestingly, we found that all the four common types of AS, especially IR, were decreased in sorghum imprinted genes, while those of the rare types, like “skip 2 exons”, “A5SS or A3SS”, MXE and other complex types were

**Fig. 4** Summary of alternative splicing (AS) events in sorghum embryo and endosperm. **a** Venn diagrams show AS events among four genotypes in embryo and endosperm, respectively. **b** Visualization of seven AS modes. The exon-intron structure of the AS events are illustrated. IR: intron retention; A3SS: alternative 3′ splicing site; ES: exon skipping; A5SS: alternative 5′ splicing site; ES1+ES2: skipping two exons; MXE: mutually exclusive exon. **c** Distribution of various splicing patterns in DEGs of embryo and endosperm, as well as in sorghum imprinted genes. BY (BTx623 × Y49) and YB (Y49 × BTx623) are reciprocal F1 hybrids, while B (BTx623) and Y (Y49) are their parental pure lines.
increased, together led to nearly 30% of the AS events were accounted by the “others” type in imprinted genes (Fig. 4c). This suggests that compared with embryo, endosperm likely has unique AS splicing mechanisms in action.

We further analyzed the sequence length covered by each AS type in sorghum embryo and endosperm. We found that the four major types of AS events showed different sequence lengths. Specifically, (i) the retained intron length increased from 60 to 100 bp; (Fig. 5); (ii) both of A3SS and A5SS showed similar features of sequence length, with the most frequent length < 10 bp; (iii) the skipped exon lengths ranged from 1 to 200 bp and peaked at 60–80 bp, but this peak was not as sharp as the other types of AS events (Fig. 5). These features of AS are consistent with previous findings in other species (Campbell et al. 2006; Shen et al. 2014).

Generation of mature mRNAs from genes requires precise removal of introns from precursor mRNAs (pre-mRNAs) and joining of exons. Four loosely conserved core sequence elements in introns, the 5′SS with a conserved GT, the 3′SS with a conserved AG, therefore, usually splice at a 5′-GT-AG-3′ exon-intron junction (Reddy et al. 2013). We compared splice motif preference among the different AS types and between tissues. We found that the splicing junction site of 5′-GT-AG-3′ pair represented the highest proportion of all splicing sites, followed by the 5′-GC-AG-3′ pair. Specifically, of the IR and A5SS events, splice junctions (SJ)s were comprised of ca. 96% GT-AG and 4% GC-AG, whereas in A3SS and ES, SJs were comprised of 98 and 99% GT-AG, 2 and 1% GC-AG, respectively. (Fig. S2). This result was in line with previous studies in other plant species (Li et al. 2014; Marquez et al. 2012; Shen et al. 2014). Further Chi-square test indicated that there was no significant difference in the frequencies of the types of splicing sites between embryo and endosperm.

Experimental validation of differential expression and AS events

To validate the DEGs and AS events detected in our transcriptomic data, we selected 20 AS genes (17 of which were also DEGs between tissues) and interrogated the accuracy via RT-PCR (primers were listed in Table S3). Results showed that most of the DEGs and AS events were very well supported by this independent experiment, with 82% of
the DEGs and 80% of the AS events were supported in both tissues (Fig. 6; Table S3).

**Gene ontology (GO) enrichment analyses of DEGs, nonadditive-expressing genes and AS genes**

To assess the possible functional relevance of the DEGs, we conducted Gene Ontology (GO) enrichment analysis by using a hypergeometric statistic (R function phyper). The significantly enriched (p value < 0.05) top GO terms in each category of the DEGs were presented in Fig. 7 and all enriched GO terms were listed in Table S4. We found that more GO terms were enriched in embryo than in endosperm when controlling q value at the 0.01 level. Those genes that exhibited embryo-specific or -preferential expression were involved in all three main GO categories: biological processes, molecular function, and cellular component. For example, the top significantly enriched GO subcategories in the biological process category were “cellular glucan metabolic process”, “DNA replication” and “signal transduction”. In the molecular function category, “ADP binding”, “ATP binding”, “heme binding”, and “nucleic acid binding” were significantly enriched. We also found that the embryo-DEGs were enriched in a few cellular component categories including “nucleus”, “kinesin complex” and “apoplast” (Fig. 7 and Table S4). In contrast, there were limited functional terms involved in endosperm-specific genes when controlling q value at 0.01. “Biological processes” of endosperm-DEGs were mainly enriched in “regulation of transcription, DNA-dependent” and “protein digestion”. In molecular function category, “transcription factor activity”, “enzyme inhibitor activity” and “aspartic-type endopeptidase activity” were significantly enriched (Fig. 7 and Table S4). GO enrichment was not detected for endosperm-DEGs in the “cellular component” category.

Analysis of GO enrichment for nonadditively expressing genes, using all sorghum genes as background, revealed that they were significantly overrepresented by some specific GO terms (Fig. 7). All the enriched GO terms were listed in Table S4. Importantly, GO terms “defense response” and “response to biotic stimulus” were significantly enriched in both tissues, indicating that nonadditively expressed genes in hybrids were possibly related to increased resistance or resilience to biotic and abiotic stresses of the hybrids. In addition, in embryo, the significantly enriched top GO terms were “transferase activity (transferring hexosyl groups)” and “ribonuclease T2 activity”, while the nonadditively expressed genes in endosperm were highly enriched in “carbohydrate metabolic process”, “oxidation reduction”, “iron ion binding”, “oxidoreductase activity”, and so on (Fig. 7 and Table S4).

We also conducted GO enrichment analysis for AS events in the different tissues and genotypes. Overall, fewer AS genes were involved in functional terms compared to the DEGs when controlling q value at the 0.01 level. Nevertheless, the AS genes of the “IR” type were highly enriched in protein metabolism process in both tissues. Importantly, the GO term “nutrient reservoir” (GO:0045735) was found to be significantly enriched in the genes of the “A3SS” and “A5SS” AS types in endosperm but not in embryo of all four genotypes, indicating functional difference of genes underwent specific types AS between the tissues. In addition, genes spliced by “A3SS”, “ES” and “A5SS” patterns of both tissues showed highly enriched in GO terms “nucleic acid binding” or “RNA binding” (q values, 0.01) (Fig. 7 and Table S4).

**Discussion**

Our RNA-seq analyses demonstrated that more than one third of genes in 14 DAP sorghum seeds showed tissue differential gene expression, with transcript abundance being higher in embryo than in endosperm, and more genes were preferentially expressed in embryo than in endosperm, whereas endosperm DEGs tended to show a higher proportion of specific expression (relative to embryo). We also found the great majority of sorghum imprinted genes exhibiting endosperm-specific or -preferential expression. Further comparative transcriptome analyses of the F1 hybrids and parental lines indicated that the genes exhibiting non-additive expression was significantly higher in endosperm than in embryo. These results suggest there exist distinct regulatory systems of gene expression between embryo and endosperm during seed development. A reasonable explanation of the multifaceted genome-wide gene expression differences between embryo and endosperm is that they are of different ploidy levels and exist in distinct transcriptional and chromatin states immediately after double fertilization. First, embryo is diploid, with one paternal genome and one
Fig. 6 RT-PCR validation of DEGs and AS events in sorghum embryo and endosperm. In the agarose gel images, arrows indicate the alternative transcripts. The length (bp) of PCR product was indicated. The gene IDs were shown above the gene model. In the gene model diagrams (not drawn to scale), blue bar indicates exon; red bar indicates the alternatively spliced regions; blue line indicates intron; the arrowhead on the gene structure diagrams indicates the position of PCR primers used for RT-PCR. 

- **a** Embryo-specific expression and intron retention (IR);
- **b** Endosperm-specific expression and IR;
- **c** Embryo-preferential expression and IR;
- **d** Endosperm-preferential expression and IR;
- **e** Endosperm-specific expression and alternative 3' splicing site (A3SS);
- **f** Embryo-specific expression and exon skipping (ES);
- **g** IR and alternative 5' splicing site (A5SS);
- **h** A sorghum actin gene used as control. BY (BTx623 × Y49) and YB (Y49 × BTx623) are reciprocal F1 hybrids, while B (BTx623) and Y (Y49) are their parental pure lines.
maternal genome, while endosperm is triploid, with two maternal genomes and one paternal genome. Second, while the major task of embryo is to ensure totipotency and perpetuation of genetic information to the next generation, that of endosperm is to nourish embryo development and growth of the germinating seedling, but genetically inconsequential to the next generation (Gehring and Satyaki 2017). Conceivably, such distinct functional partitioning entails substantially variable gene regulation circuits and hence their direct molecular readouts.

Numerous observations have suggested that, like animals, there are many alternative splicing (AS) events in plants, which occurred differentially in different tissues and developmental stages. Multicellular organisms may generate different splice forms of the same gene in different tissues, or even within different cells in the same tissue (McGuire et al. 2008). Our study demonstrated that the number of AS events and the frequency of AS types varied in embryo- or endosperm-specific genes and imprinted genes. The frequency of AS events was higher in embryo than in endosperm, a similar trend was found in soybean (Shen et al. 2014). These tissue-specific AS events may function in a coordinated manner in specific pathways or interaction networks as groups of genes that are coregulated at the transcriptional level (Shen et al. 2014). Moreover, in this study we found that sorghum imprinted genes showed different AS patterns from the rest “normal” genes, implying a role of AS in regulating imprinted gene expression.

The most common AS type in plants is intron retention (IR) whose rates vary among plant species and individuals of the same species, and even differ in the same genotype, ranging from 30 to 62% (Lu et al. 2013; Mei et al. 2017; Thatcher et al. 2014). One possible reason is that different statistical approach or classification methods were used. In some studies, IR only counts the mode that retains one intron, and those from different combinations of the four basic types are categorized into the “Others” AS type (illustrated in Fig. 4b); other studies not only count the mode that retain one intron, but also all of the IR-containing complicated cases, for instance, “IR1 + IR2”, “IR1 or IR2”, “A3SS and IR”, and “A3SS or IR” are all included in IR (Marquez et al. 2012; Zhang et al. 2017b). In the later categorizations, the percentage of IR events can reach to 40–47% of all AS events (Marquez et al. 2012; Zhang et al. 2017b). If count the simple IR only, the IR frequency will be down to about 26% in the studies of Arabidopsis (Marquez et al. 2012; Zhang et al. 2017b). Our analysis also indicated that IR was the most variable AS type in sorghum. A previous study in maize suggested that IR may act to fine-tune gene expression across seed development (Mei et al. 2017). However, some IR events were recently shown to be more likely to represent partially spliced transcripts due to their low abundance (Syed et al. 2012). Although the biochemical mechanisms of AS are complex and in large part remain poorly understood, AS is an important mechanism whereby plants modulate gene expression in development (Syed et al. 2012).

Gene ontology (GO) analysis of the DEGs between the two tissues revealed that genes associated with different GO terms are potentially related to the functions of a given tissue during seed development. There were 1459 and 2307 genes specifically expressed in embryo and endosperm, respectively; however, the greater number of GO terms enriched in embryo-specific genes than in endosperm-specific genes suggests embryo may perform more specialized functions. In contrast, nonadditive gene expression was more prevalent in endosperm than in embryo; accordingly, more enriched GO terms are involved in nonadditive genes of endosperm than in those of embryo. This may suggest that compared with embryo, gene regulation in endosperm is more permissive (i.e., regulated with less stringency), consistent with inconsequential of endosperm to the perpetuation of genetic information. Moreover, analysis of GO enrichment on AS genes indicated that GO terms with “nutrient reservoir activity” was enriched only in endosperm, consistent with the role of endosperm in nutrient storage, and again indicating fundamental functional difference between the two tissues (Merkin et al. 2012; Wang et al. 2018).

In sum, the comprehensive gene expression data, and the elucidation of differential gene expression (including tissue-preferential expression) and differential AS patterns between tissues and between F1 hybrids and parents in sorghum provide foundational information for further functional genetic and genomic research on sorghum and other related cereal crops. In particular, the comparisons between F1 hybrids and their pure-line parents with respect to genome-wide gene expression differences between embryo and endosperm may have implications to further our understanding of the molecular mechanisms underpinning heterosis. Like in maize, heterosis occurs ubiquitously in sorghum seed production. Conceivably, endosperm makes a much greater contribution to seed production than dose embryo. Thus, our observations that a great majority of imprinted genes exhibiting endosperm-specific or -preferential expression and that nonadditive expression in F1 hybrids was significantly higher in endosperm than in embryo lend further support to previous findings in maize (Springer and Stupar 2007) that gene regulation is a major contributor to the formation of crop yield heterosis.
| GO:0001522 | pseudouridine synthesis |
| GO:0005975 | carbohydrate metabolic process |
| GO:0006734 | cellular glucan metabolic process |
| GO:0006896 | glycolysis |
| GO:0006260 | DNA replication |
| GO:0006281 | DNA repair |
| GO:0006355 | regulation of transcription, DNA-dependent |
| GO:0008357 | regulation of transcription from RNA polymerase II promoter |
| GO:0006468 | protein amino acid phosphorylation |
| GO:0006470 | protein amino acid dephosphorylation |
| GO:0006508 | proteolysis |
| GO:0006810 | transport |
| GO:0006952 | defense response |
| GO:0006979 | response to oxidative stress |
| GO:0007018 | microtubule-based movement |
| GO:0007185 | signal transduction |
| GO:0008033 | RNA processing |
| GO:0008283 | cell proliferation |
| GO:0008296 | spemidine biosynthetic process |
| GO:0008356 | asymmetric cell division |
| GO:0009110 | lipid biosynthetic process |
| GO:0009451 | RNA modification |
| GO:0009607 | response to biotic stimulus |
| GO:0007925 | response to hormone stimulus |
| GO:0009755 | photosynthesis, light harvesting |
| GO:0009836 | radial pattern formation |
| GO:0018439 | aromatic compound catalytic process |
| GO:0019538 | protein metabolic process |
| GO:0043866 | leaf development |
| GO:0051114 | oxidation reduction |

| GO:0001868 | nucleotide binding |
| GO:0003676 | nucleic acid binding |
| GO:0003677 | DNA binding |
| GO:0003700 | transcription factor activity |
| GO:0003722 | RNA binding |
| GO:0003747 | translation release factor activity |
| GO:0003777 | microtubule motor activity |
| GO:0003824 | catalytic activity |
| GO:0004185 | serine-type carboxypeptidase activity |
| GO:0004190 | aspartic-type endopeptidase activity |
| GO:0004593 | hydrolyzing O-glycosyl compounds |
| GO:0004501 | peroxidase activity |
| GO:0004672 | protein kinase activity |
| GO:0004722 | protein serine/threonine phosphatase activity |
| GO:0004857 | enzyme inhibitor activity |
| GO:0005215 | transporter activity |
| GO:0005506 | iron ion binding |
| GO:0005524 | ATP binding |
| GO:0008017 | microtubule binding |
| GO:0008230 | growth factor activity |
| GO:0008270 | zinc ion binding |
| GO:0009682 | pseudouridine synthase activity |
| GO:0016491 | oxidoreductase activity |
| GO:0016705 | oxidoreductase activity, reduction of molecular oxygen |
| GO:0016708 | oxidoreductase activity, oxygen, or NADH as donor |
| GO:0016747 | transferase activity, transferring acyl groups |
| GO:0016758 | transferase activity, transferring hexosyl groups |
| GO:0016762 | xyloglucan:xyloglucosyl transferase activity |
| GO:0016787 | hydrolase activity |
| GO:0016788 | hydrolase activity, acting on ester bonds |
| GO:0020378 | heme binding |
| GO:0022657 | transmembrane transporter activity |
| GO:0038979 | ribonuclease T2 activity |
| GO:0043531 | ADP binding |
| GO:0043565 | sequence-specific DNA binding |
| GO:0045735 | nutrient reservoir activity |
| GO:0046972 | metal ion binding |
| GO:0046983 | protein dimerization activity |
| GO:0051082 | unfolded protein binding |
| GO:0051087 | chaperone binding |

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| GO:0000786 | nucleosome |
| GO:0005576 | extracellular region |
| GO:0005634 | nucleus |
| GO:0005871 | kinesin complex |
| GO:0016020 | membrane |
| GO:0040406 | apoplastic |
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**Fig. 7** Log10 (q value) fold change heatmap showing gene ontology (GO) terms of DEGs and AS genes in sorghum embryo and endosperm. GO terms with significant enrichment (q value < 0.05) in at least one comparison were shown. Color in heatmap represent the log10 (q value). Em and En are abbreviations for embryo and endosperm, respectively. IR intron retention, A3SS alternative 3′ splicing site, ES exon skipping, A5SS alternative 5′ splicing site.

**Supplementary Information** The online version contains supplementary material available at https://doi.org/10.1007/s11103-021-01196-y.

**Acknowledgements** This study was supported by the National Natural Science Foundation of China (No. 31370341) and the Natural Science Foundation of Jilin Province, China (No. 20150101086JC). The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.

**Author contributions** MSZ, BL and WGY designed the research. Material preparation, data analysis and experimental validation were performed by NL and MSZ. MSZ, BL and NL wrote the manuscript. All authors read and approved the manuscript.

**Data availability** RNA-seq data of endosperm can download from SRA, the accession number is PRJNA302593. For the embryo, transcriptome data generated in this study have been deposited to NCBI under the accession number of PRJNA732112, and are available at: https://dataview.ncbi.nlm.nih.gov/object/PRJNA732112?reviewer=j64as52ku120aqj4d2la2ain2h.

**Declarations**

**Conflict of interest** The authors declare that they do not have any conflict of interest.

**Ethical approval** We declare that this manuscript complies with all ethical rules required by this journal.

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