Comparative Study on the Transcriptome of Maize Mature Embryos from Two China Elite Hybrids Zhengdan958 and Anyu5

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Abstract

Zhengdan958 and Anyu5 are two elite maize hybrids of China, which manifest similar paternal lines (Chang7-2) but different maternal lines (Zheng58 and Ye478). Zhengdan958 has a 10–15% yield advantage over Anyu5. In this study, we applied digital gene expression technology to analyze transcriptomes of mature embryos from the two hybrids and their parents, aimed to investigate molecular mechanism of heterosis and genetic effects of maternal lines. Results showed that 71.66% and 49.70% of differentially expressed genes exhibited non-additive expression in Zhengdan958 and Anyu5, respectively. The number of non-additive genes involved in abiotic and biotic stress responses in Zhengdan958 was higher than that in Anyu5, which was in agreement with their phenotypic performance. Furthermore, common over-dominance and under-dominance genes (137 and 162, respectively) between the two hybrids focused on plant development and abiotic stress response. Zhengdan958 contained 97 maternal expression-level dominance (maternal-ELD) genes, and the number was higher than that of Anyu5 (45). Common up-regulated maternal-ELD genes were significantly enriched in meristem and shoot development while common down-regulated maternal-ELD genes were involved in pyruvate metabolic process, negative regulation of catalytic activity and response to stress. Therefore, non-additive genes mainly contribute to heterosis in Zhengdan958, including many genes for plant development, abiotic and biotic stress responses. Maternal effects may play important roles in maize heterosis.

Introduction

Heterosis is the phenomenon that F1 hybrids exhibit phenotypes that are superior to those of their parents [1–2]. Plant breeders use heterosis to breed superior varieties in many important crop species. Among them, maize is a typical model crop for studying heterosis. More than 95% of China maize acreage is planted with hybrids. Duvick estimates that maize hybrids exhibit a 15% yield advantage relative to superior open-pollinated varieties and that worldwide heterosis accounts for an additional 55 million metric tons of grain yield annually [3]. Despite the wide use of heterosis in plant breeding, the molecular basis of heterosis is not well understood.
In maize breeding, different inbred lines are divided into stiff stalk heterosis group (SS group) and non-stiff stalk heterosis group (NSS group) on the basis of level of grain yield heterosis. Generally, crosses within heterosis groups show lower grain yield heterosis than crosses between groups. Zheng58, Chang7-2, and Ye478 are the most popular inbred lines in China; Zheng58 and Ye478 belong to SS group (designated as maternal group), and Chang7-2 is an important inbred line of NSS group (designated as paternal group). Chang7-2 exhibits strong hybrid vigor as well as Zheng58 and Ye478. For example, Zheng58 and Chang7-2 are the parental lines of the famous commercial hybrid-Zhengdan958 that is currently the most widely grown variety in China. Ye478 and Chang7-2 are the parental lines of another commercial hybrid-Anyu5, which was widely grown in the 1990s. However, Zhengdan958 shows greater yield, higher planting density, and more stress-tolerance than Anyu5. Lai’s study indicated that Ye478 is one of the parental lines of Zheng58, which inherited 43% of its genomic content from inbred line Ye478 [4]. Thus, genetic difference between maternal lines Zheng58 and Ye478 may account for the difference between Zhengdan958 and Anyu5 in terms of grain yield and resistance.

The availability of high-throughput gene expression profiling technologies currently allows researchers to study the gene expression profile of hybrids relative to the inbred parents [5]. In general, most of these studies focused on characterizing gene expression patterns in single heterotic hybrid compared with the two parents. In this study, we investigated the gene expression profiles of two maize hybrids with common paternal line and different maternal lines using digital gene expression (DGE) technologies, in order to elucidate the molecular mechanism of heterosis and genetic effect of maternal lines.

Materials and Methods

Sample collection and RNA extraction

The two maize hybrids Zhengdan985, Anyu5 and their parents Ye478, Zheng58, and Chang7-2 were grown under similar field conditions with three replications. Ten mature seeds of each material were collected at 40 days after fertilization and frozen in liquid nitrogen. Mature embryos were extracted from seeds by hand using scalpels. Three replicate embryo samples of each genotype were mixed equally, and total RNA was extracted with Trizol (Invitrogen).

Library construction and sequencing

The mRNAs were purified from 6 g of total RNA (300 ng/μl) using poly (T) oligo-attached magnetic beads. After mRNA’s binding, cDNA synthesis was performed. Double strand cDNA was introduced into the cDNA fragment digested by NlaIII endonuclease and the binding fragment containing sequences of CATG site and adjacent poly (A) tail in 3’ end. After the precipitation of 3’ cDNA fragment, Illumina adaptor 1 was added to 5’ end. Both adaptor 1 and CATG site can be recognized by Mme I, which cut at downstream CATG site and produce fragment of 17 bp tags with adaptor 1. Adaptor 2 was added to the 3’ end of these tags after getting rid of fragment with beads in 3’ end. These sequences were prepared for Solexa sequencing. Transcriptome sequencing was performed by sequencing by synthesis method on the Illumina platform at BGI-Shenzhen (China). The data size per sample was 500 Mbp.

Gene expression classification

Clean-tags were obtained by filtering the adaptor sequences and removing low-quality sequences (containing ambiguous bases). The clean tags were then mapped to the reference genome and genes of the maize available at ftp://ftp.maizesequence.org/pub/maize/release-5b
Only the tags with perfect match or one mismatch were further considered and annotated on the basis of reference genes. The expression level of each gene was estimated by the frequency of clean tags and then normalized to number of transcripts per million clean tags (TPM), which is a standard method that is extensively used in differential gene expression analysis [7]. The expression level of each gene was measured by the normalized number of matched clean tags.

Identification of differentially expressed genes (DEGs) and gene annotations

For each sequenced library, software package edgeR was used to adjust the read counts by one scaling normalized factor [8]. Differential expression analysis was performed by DEGseq program package for comparisons among samples [9]. Significant DEGs were defined with the Benjamini and Hochberg corrected P-value < 0.05 [10].

Gene ontology (GO) enrichment analyses were conducted according to Benjamini’s study [10]. When the GO database was searched, the GO categorization of all DEGs was displayed as three categories for cellular component, molecular function, and biological process. A GO term was considered significantly enriched if it had a hypergeometric test P < 0.05, a Benjamini and Hochberg (BH) FDR < 0.1.

Classification of gene expression patterns

Gene expression in hybrids compared to their parents can be classified into additive and non-additive expression. Among non-additive expression, the hybrid gene expression displays a significantly higher level or lower level than both parents, and the two instances have been previously designated above high-parent and below low-parent expression [11] or under-and over-dominance [12]. Moreover, there is such a phenomenon that hybrid gene expression levels which equal that of one parent and differ from that of the other parent in non-additive expression patterns. This phenomenon is termed expression-level dominance (ELD).

Results

Transcriptome analysis of Zhengdan958, Anyu5 and their parental lines

In this study, we sequenced five mature embryos DGE libraries from three inbred parents (Zheng58, Ye478 and Chang7-2) and two F1 hybrids (Zheng58×Chang7-2 = Zhengdan958; Ye478×Chang7-2 = Anyu5) by massive parallel sequencing on the Illumina platform at BGI-Shenzhen, China. The number of tags for each gene was calculated and normalized to TPM. Given the robustness of subsequent data analysis, only transcripts with more than one TPM were considered as expressed genes. Using these criteria, we obtained 12,859 unique tag-matched genes for Zhengdan958, 13,191 unique tag-matched genes for Anyu5, 13,294 unique tag-matched genes for Zheng58, 12,965 unique tag-matched genes for Ye478, and 12,551 unique tag-matched genes for Chang7-2 (S1 Table). For convenience, we hereafter use AA, BB and CC representing Zheng58, Ye478 and Chang7-2, respectively. Thus, AC indicates Zhengdan958, and BC denotes Anyu5.

Transcriptome differentiation between two maize hybrids Zhengdan958 and Anyu5

To study the effect of maternal lines and molecular mechanism for heterosis in Zhengdan958 and Anyu5, we analyzed their transcriptomes against paternal inbred line Chang7-2. The two
hybrids and Chang7-2 share the common paternal line and different maternal lines. Therefore, it is helpful to have a better understanding of maternal effects on hybrids.

A total of 847 DEGs among AC, BC and CC transcriptomes were identified. Among them, 424 DEGs (50.06%) were significantly up-regulated (AC > BC > CC), and 423 (49.94%) DEGs were significantly down-regulated (AC < BC < CC) (Fig 1). By GO enrichment analysis, we found up-regulated transcripts contained more DEGs for response to abiotic stress, biotic defense response, and photosynthesis than down-regulated transcripts (Fig 2A). For example, a total of 32 and 25 DEGs associated with response to abiotic stress were significantly enriched in up-regulated transcripts and down-regulated transcripts, respectively. Only three down-regulated genes played roles in response to biotic stress, whereas 12 up-regulated genes were found in this process. In terms of genes associated with photosynthesis, only one gene was found in down-regulated transcripts while six genes were significantly enriched in up-regulated transcripts.

We further classified up-regulated and down-regulated transcripts into non-additive and additive genes. Among 847 genes that were differentially expressed among the two hybrids and their paternal parent, 240 (28.34%) were additive genes while 607 (71.66%) were non-additive genes in Zhengdan958 (Fig 1A and 1C). In Anyu5, 426 and 421 genes displayed additive and non-additive expression, respectively (Fig 1B and 1D).

Subsequently, we pursued possible functions of genes showing non-additive expression (Fig 2B). The results showed non-additive genes in Zhengdan958 and Anyu5 were significantly enriched in carbohydrate biosynthetic and metabolic process and photosynthesis (12 and 13 genes in Zhengdan958 and Anyu5), plant development (21 and 15 genes in Zhengdan958 and Anyu5), abiotic stress response (11 and 14 genes identified in Zhengdan958 and Anyu5), and

**Fig 1.** Pie chart of differentially expressed genes (DEGs) in up-regulated (A, B) and down-regulated (C, D) transcripts in Zhengdan958 (A, C) and Anyu5 (B, D). Add and nonadd represent additive and non-additive genes. Maternal-ELD: maternal expression-level dominance. AA and BB represent Zheng58 and Ye478, respectively.

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biotic defense response (nine and four genes in Zhengdan958 and Anyu5). Non-additive expression mainly contributed to heterosis in Zhengdan958, and covered many genes associated with plant development, abiotic and biotic stress responses.

Over-dominance, under-dominance and maternal expression-level dominance (maternal-ELD) expression pattern in Zhengdan958 and Anyu5

Non-additive genes can be further classified into over-dominance, under-dominance and maternal-ELD genes. Both in Zhengdan958 and Anyu5, we found over-dominance and under-dominance expression accounted for the majority of up-regulated transcripts and down-regulated transcripts, respectively. For up-regulated transcripts, there were 209 (82.28% of 254 non-additive genes) and 146 (71.22% of non-additive 205 genes) over-dominance genes in Zhengdan958 and Anyu5, respectively (Fig 1A and 1B). For down-regulated transcripts, under-dominance genes (78.19%, 276 out of 353 non-additive genes in Zhengdan958; 82.41%, 178 out of 216 non-additive genes in Anyu5) were the most prevalent in the non-additively expressed genes (Fig 1C and 1D).

Both in Zhengdan958 and Anyu5, maternal-dominance genes composed a small portion of non-additive genes in up-regulated or down-regulated transcripts (Fig 1). For Zhengdan958, 16.54% of genes (42 out of 254 up-regulated non-additive genes) represented maternal-ELD while for Anyu5, the percentage of maternal dominance was even lower (8.78%, 18 out of 205 up-regulated non-additive genes). Similar to up-regulated transcripts, a higher portion of maternal-ELD (15.58%, 55 out of 353 non-additive genes) displayed in Zhengdan958 while only 12.50% (27 out of 216 non-additive genes) were Ye478 maternal-ELD in down-regulated transcripts.

There were 137 over-dominance genes and 162 under-dominance genes that were held in common between Zhengdan958 and Anyu5 (Fig 3A and 3B). We also found the expression of six genes was activated while ten genes were repressed both in Zheng58 and Ye478 maternal-ELD (Fig 4). This indicated that these genes were conserved and important for heterosis.

We firstly investigated GO functional categories of common over-dominance and under-dominance genes. Those common over-dominance genes were significantly enriched in plant development (six genes), abiotic stress response (six genes), photosynthesis (four genes), and carbohydrate biosynthetic and metabolic processes (four genes) (Fig 3C and S2 Table). For
common under-dominance genes, there were eleven, nine and six genes associated with plant development, abiotic stress response and carbohydrate metabolic process, respectively (Fig 3C and S3 Table).

Fig 3. Venn diagram analysis of over-dominance (OD) (A) and under-dominance (UD) (B) genes in Zhengdan958 and Anyu5, and bar plot of GO enrichment analysis of common over-dominance and under-dominance genes between the two hybrids (C). AC and BC denote Zhengdan958 and Anyu5, respectively.

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The functions of specific over-dominance and under-dominance genes in the two hybrids were analyzed (S4 and S5 Tables). In Zhengdan958, there were seven, four and two over-dominance genes associated with plant development, abiotic stress response and brassinosteroid, respectively. In Anyu5, those specific over-dominance genes were mainly enriched in abiotic stress response. For specific under-dominance genes in Zhengdan958, there were seven genes associated with plant development, four genes for abiotic stress response, two for defense response to bacterium, and two for plant hormone such as indolebutyric acid and abscisic acid. The significant biological process was tetrahydrofolate metabolic process for Anyu5 under-dominance genes.

Analysis of the functional annotations of common genes between Zheng58 and Ye478 maternal dominance were conducted. Up-regulated common genes between Zheng58 and Ye478 maternal-ELD were significantly enriched in regulation of meristem development, shoot system development and cellular amino acid metabolic process (Fig 4C). These genes were in cell components including chloroplast and cytosol. Down-regulated common genes were significantly enriched in the pyruvate metabolic process, negative regulation of catalytic activity and response to stress, and were located in pyruvate dehydrogenase complex (Fig 4C).

We also discovered functional categories of specific maternal-ELD in the two hybrids. GO enrichment analysis indicated that in up-regulated genes, Zheng58 specific maternal-ELD genes were significantly enriched in plant development such as seed germination and leaf morphogenesis, carotenoid biosynthetic process, response to heat and gene silencing. Up-regulated Ye478 maternal-ELD genes were enriched in significant biological processes including photosynthetic and respiratory electron transport chain, defense response to bacterium incompatible interaction and toxin catabolic process (Tables 1 and 2). In down-regulated transcripts, Zheng58 maternal-ELD genes were significantly enriched in cell growth, photorespiration, regulation of ethylene biosynthetic process, response to abscisic acid and tetrahydrofolate metabolic process, while the Ye478 maternal-ELD genes were significantly involved in pollen germination, pollen tube growth, response to phosphate starvation, carotene, xanthophyll and chlorophyll biosynthetic process and epigenetics such as regulation of gene expression epigenetic and chromatin remodeling (Tables 1 and 2). These indicated that Zheng58 and Ye478 specific maternal-ELD genes showed obvious functional distinctions, which may be important players for shaping the difference of heterosis in Zhengdan958 and Anyu5.

**Discussion**

Heterosis is the outstanding performance of hybrids compared to either of their parents. Biological processes during maize mature embryo development after fertilization may be of importance for heterosis. To elucidate genetic effect of heterosis and maternal parents, mature embryos of two commercial maize hybrids Zhengdan958 and Anyu5 were compared using DGE technologies in this study. In a previous study, Zheng58 exhibits 57% difference with Ye478 in genomics [4]. However, our findings indicated that Zheng58 showed 18.17% difference with Ye478 in the mature embryo transcriptome. This result showed that a large portion of differential genes were not expressed in mature embryos. In the present study, several DEGs detected by digital gene expression profiling were already validated by quantitative RT-PCR [13].

**Non-additive genes shapes the China commercial hybrid Zhengdan958 transcriptome**

Non-additive expression that significantly differs from the mid-parent value is mainly attributable to trans-regulated interaction in a hybrid. At the transcriptome level, an increasing
number of studies in plants showed that non-additive gene expression is quite prevalent in various types of hybrid situations [14–15]. The increase in activity of non-additive genes was associated with increased photosynthesis and determination of cell size and number through development, suggesting that non-additive genes played an important role in biomass heterosis [16]. Meyer et al. [17] proposed that non-additive expression led to higher metabolic activity because of an efficient utilization of resources, and therefore improved performance in hybrids. At the proteome level, Hoecker et al. [18] pointed out that non-additive protein accumulation...
in young primary roots of maize hybrids might be associated with heterosis manifestation. In addition, many transcripts/e-traits that were controlled many eQTL with opposite allelic effects displayed higher heritability in RIL population than their parents, indicating non-additive genetic variation [19]. Other studies also found a similar number of genes that displayed additive and non-additive expression [20], and the additive gene expression was predominated in heterotic hybrid [21–22]. This discrepancy could be due to the different experimental designs, statistical approaches, different plant materials, different tissues and/or different developmental stages [23]. In this study, non-additive expression patterns were overrepresented in Zhengdan958, and accounted for more than 70% of DEGs. In the hybrid Anyu5, a similar number of additive genes and non-additive genes were identified (426 additive genes and 421 non-additive genes). This data showed the proportional difference of non-additive genes between Zhengdan958 and Anyu5 resulted from the discrepancies of trans-regulated interaction in the two hybrids.

Stress adaption capability is an important factor contributing to heterosis. GO enrichment analysis indicated the total number of non-additive genes involved in abiotic stress response and biotic stress defense in Zhengdan958 were higher than that in Anyu5. This may be in agreement with phenotypic performance that Zhengdan958 shows more adaptation resistance than Anyu5. Therefore, the predominance of non-additive genes in Zhengdan958 may result in yield advantage over Anyu5, and exhibited a trans-regulatory mechanism to act in hybrid mature embryos. A large portion of non-additive genes in this study did not indicate whether this gene expression pattern is solely responsible for heterosis, but it must contribute to hybrid effects to a large extent.

In this study, over-dominance genes accounted for 34.43%-34.68% of non-additive genes in Zhengdan958 and Anyu5. We found about 22.57%-32.54% of common over-dominance genes between the two hybrids, were enriched in abiotic stress responses such as phototropism.

Table 1. GO enrichment analysis of specific maternal expression-level dominance genes of Zheng58.

| GO ID          | Biological process                           | P value | FDR value | Gene ID               |
|---------------|----------------------------------------------|---------|-----------|-----------------------|
| **Up-regulated** |                                              |         |           |                       |
| GO:00030154   | cell differentiation                         | 0.0011  | 0.0042    | GRMZM2G126128         |
| GO:0009845    | seed germination                             | 0.0013  | 0.0049    | GRMZM2G034896         |
| GO:0009965    | leaf morphogenesis                           | 0.0042  | 0.0120    | GRMZM2G126128         |
| GO:0016117    | carotenoid biosynthetic process              | 0.0026  | 0.0078    | GRMZM2G126128         |
| GO:0006952    | defense response                             | 0.0130  | 0.0276    | GRMZM5G819919         |
| GO:0009408    | response to heat                             | 0.0151  | 0.0313    | GRMZM2G034896         |
| GO:0016458    | gene silencing                               | 1.62E-04| 9.04E-04  | GRMZM2G169998         |
| GO:0030001    | metal ion transport                          | 0.0088  | 0.0201    | GRMZM2G001803         |
| **Down-regulated** |                                          |         |           |                       |
| GO:0009825    | multidimensional cell growth                 | 0.0006  | 0.0022    | GRMZM2G075701         |
| GO:0080167    | response to karrin                          | 0.0099  | 0.0202    | GRMZM6G734131         |
| GO:0009853    | photospiration                              | 0.0012  | 0.0043    | GRMZM2G168281         |
| GO:0010364    | regulation of ethylene biosynthetic process  | 1.14E-05| 1.44E-04  | GRMZM2G039280         |
| GO:0009737    | response to abscisic acid                    | 0.0396  | 0.0698    | GRMZM2G063931         |
| GO:0046653    | tetrahydrofolate metabolic process           | 1.91E-06| 8.40E-05  | GRMZM2G168281         |
| GO:0006120    | mitochondrial electron transport NADH to ubiquinone | 1.98E-04| 0.0010    | GRMZM5G804358         |

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photoperiodism flowering and response to desiccation and UV and hyperosmotic salinity response, which were important for maize hybrid vigor (S3 Table). Many studies indicated that over-dominance would lead to heterosis in Arabidopsis [16] and rice [24]. Among these common over-dominance genes, GRMZM2G105400 was significantly enriched in phototropism, anthocyanin accumulation in tissues in response to UV light and response to blue light as well as regulation of hormone levels, cell tip growth, root hair elongation and polysaccharide biosynthetic process. Phototropism allows plants to change the growth direction responding to the location of the light source [25], which is important in maize root and coleoptile development [26–27]. Many studies have shown that three groups of molecules were involved in the mechanism of phototropism in Arabidopsis. The first one was photoreceptor which responded to UV, blue and green light; the second group of molecules were involved in phototropism functions in the signaling pathway activated by photoreceptors, and the third group participated in the process that produced the plant curvature after signals were received [25]. In the third group, studies supported that plant hormone auxin played an important role in differential growth responses in Arabidopsis and maize [28–29]. Therefore, GRMZM2G105400 played a crucial role in phototropism in Zhengdan958 and Anyu5, which was important for hybrid vigor.

Maternal effects play roles in maize heterosis

Maize hybrids Zhengdan958 and Anyu5 possess the same paternal line but different maternal lines. Thus, the difference in the transcription profiling between Zhengdan958 and Anyu5 may

| GO ID       | Biological process                                      | P value   | FDR value | Gene ID       |
|-------------|---------------------------------------------------------|-----------|-----------|---------------|
| Up-regulated|                                                          |           |           |               |
| GO:0022904  | respiratory electron transport chain                    | 2.86E-05  | 1.90E-04  | GRMZM2G103103 |
| GO:0009767  | photosynthetic electron transport chain                 | 7.71E-05  | 2.79E-04  | GRMZM2G103103 |
| GO:0009816  | defense response to bacterium incompatible interaction  | 4.07E-05  | 1.91E-04  | GRMZM2G015889 |
| GO:0009407  | toxin catabolic process                                 | 3.23E-05  | 1.90E-04  | GRMZM2G102216 |
| Down-regulated|                                                     |           |           |               |
| GO:0009846  | pollen germination                                      | 3.23E-04  | 0.0011    | GRMZM2G162670 |
| GO:0016036  | cellular response to phosphate starvation              | 4.78E-04  | 0.0015    | GRMZM2G377761 |
| GO:0009560  | embryo sac egg cell differentiation                     | 5.44E-04  | 0.0016    | GRMZM2G135410 |
| GO:0009860  | pollen tube growth                                      | 7.15E-04  | 0.0021    | GRMZM2G162670 |
| GO:0016120  | carotene biosynthetic process                           | 7.37E-06  | 3.32E-05  | GRMZM2G377761 |
| GO:0016123  | xanthophyll biosynthetic process                        | 1.03E-05  | 4.37E-05  | GRMZM2G377761 |
| GO:0015995  | chlorophyll biosynthetic process                        | 0.0011    | 0.0028    | GRMZM2G377761 |
| GO:0040029  | regulation of gene expression epigenetic               | 4.92E-06  | 2.53E-05  | GRMZM2G135410 |
| GO:0006338  | chromatin remodeling                                    | 6.65E-05  | 2.39E-04  | GRMZM2G135410 |
| GO:0046520  | sphingoid biosynthetic process                          | 7.37E-06  | 3.32E-05  | GRMZM2G162670 |

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be mainly attributed to maternal lines Zheng58 and Ye478. DNA methylation profiles of the maize endosperm indicated that maternal alleles are more demethylated than paternal alleles, which indicates that maternal alleles are more active than parental alleles [30]. Nodine and Bartel [31] found that maternal parents contribute equally to transcriptome of early embryos as well as paternal lines in Arabidopsis. We found that Zhengdan958 contained more maternal dominance genes than Anyu5 (2.16-fold, 97/45). Significant GO terms indicated that maternal-ELD specific genes were enriched in important biological processes associated with heterosis, and displayed different functions in two hybrids. In addition, common maternal-ELD genes were focused on plant development and response to stress. Therefore, maternal lines may play important roles in maize heterosis by maternal dominance mechanism.

One objective of this study is to elucidate the genetic effects of maternal parents. Investigations of gene functions could be useful to understand maternal effects. Therefore, we discussed functions of several DEGs showing Zheng58 and Ye478 maternal-dominance. Common gene GRMZM2G018375 or thiamine thiazole synthase 1 (\textit{THI1}) was involved in response to stress in Zhengdan958 and Anyu5. Li et al. demonstrated that \textit{THI1} played important roles in both guard cell abscisic acid signaling and drought response in Arabidopsis [32]. Moreover, a large number of studies found that thiamine biosynthesis was up-regulated during plant adaptation responses to abiotic stress such as salt, cold, flood, heat, drought, and osmotic stress [33–36]. And the up-regulation of thiamine biosynthesis under salt and osmotic stress conditions was mediated by abscisic acid at the early seedling stages in Arabidopsis [37]. In this study, gene GRMZM2G063931 involved in response to abscisic acid process were displaying Zheng58 maternal-ELD as well as \textit{THI1}, indicating the regulation of thiamine biosynthesis during plant adaptation responses to abiotic stress may be mediated by abscisic acid in maize mature embryos.

The gene GRMZM2G034896 (Zheng58 maternal-ELD gene) was up-regulated and involved in response to heat and seed germination in this study. Its homolog in Arabidopsis, respiratory burst oxidase homolog protein B \textit{RBOHB}, also played essential roles in plant development such as seed germination and seedling elongation development [38] as well as response to abiotic stimuli and pathogen defense [39]. The Ye478 maternal-ELD gene GRMZM2G015889 was activated and associated with defense response to bacterium incompatible interaction. This is consistent with the function of its homolog serine/threonine-protein kinase \textit{PBS1} that was required in specific recognition of the bacterial protein in Arabidopsis [40]. Lee and Kim also found that \textit{OsPBL1} is involved in antiviral defense signaling pathways in rice [41]. An overexpression of the two DEGs in mature hybrid embryos may prompt stress adaption capability in seed development and possible roles for hybrid vigor.

Two DEGs (GRMZM2G063931 and GRMZM2G039280) associated with plant hormone displayed maternal-ELD of Zheng58, and were in a down-regulated state. GRMZM2G063931 (ubiquitin-conjugating enzyme E2) was involved in response to abscisic acid. Ubiquitin plays important roles in maintaining cellular homeostasis and enables effective adaptation to environmental changes. Seo et al. reported that the role of ubiquitination was associated with hormonal signaling [42]. Therefore, the maternal-ELD gene GRMZM2G063931 may regulate plant response to abiotic stress through hormone abscisic acid mediated by ubiquitination in Zhengdan958. GRMZM2G039280 (ethylene-overproduction protein 1) was significantly involved in regulation of ethylene biosynthetic process, and played roles in response to drought as well in this study. Moreover, Du et al. reported that \textit{OsETOL1}, the homolog of GRMZM2G039280, played distinct roles in drought and submergence tolerance by modulating ethylene production and energy metabolism in rice [43]. When ethylene biosynthesis pathway is repressed, it may induce changes in a wide range of developmental processes and adaptation responses [44].
In addition, the gene GRMZM2G135410 coding for chromatin remodeling and regulation of gene expression epigenetic proteins were expressed in the way of Ye478 maternal-ELD pattern, indicating a contribution to epigenetic regulation of hybrids. One gene GRMZM2G169998 showing the Zheng58 maternal-ELD pattern was also significantly involved in epigenetic modification process gene silencing. A recent study revealed that epigenetics might be helpful and complementary to conventional genetic explanations of the molecular mechanism of heterosis [45]. This might be true for embryo development after fertilization, because the interaction of the different parental genomes has to be coordinated and adjusted for generating heterosis [46].

In summary, the analysis of gene expression during mature seed embryos manifestation as well as the analysis of non-additive genes may contribute to a better understanding of the molecular networks regulating the heterosis and effect of maternal lines in Zhengdan958 and Anyu5.

Supporting Information
S1 Table. Expressed genes in Zhengdan958, Anyu5, Zheng58, Ye478 and Chang7-2.

S2 Table. GO enrichment analysis of common over-dominance genes between Zhengdan958 and Anyu5.

S3 Table. GO enrichment analysis of common under-dominance genes between Zhengdan958 and Anyu5.

S4 Table. GO enrichment analysis of specific over-dominance genes in Zhengdan958 and Anyu5.

S5 Table. GO enrichment analysis of specific under-dominance genes in Zhengdan958 and Anyu5.

Author Contributions
Conceived and designed the experiments: HL JM. Performed the experiments: JM JL YC LW FW HW. Analyzed the data: JM HL. Contributed reagents/materials/analysis tools: HL JL YC LW HW. Wrote the paper: JM HL.

References
1. East EM. Inbreeding in corn. Rep. Conn. Agric. Exp. Stn. 1908; 1907: 419–428.
2. Shull GH. The composition of a field of maize. J Hered. 1908; 296–301. doi:10.1093/jhered/os-4.1.296
3. Duvick DN. Heterosis: feeding people and protecting natural resources. In the genetics and exploitation of heterosis in Crops; Coors J.G.; Pandey S, Eds.; Amer Society of Agronomy: Madison, WI, USA. 1999; 19–29.
4. Lai J, Li R, Xu X, Jin W, Xu M, Zhao H, et al. Genome-wide patterns of genetic variation among elite maize inbred lines. Nat Genet. 2010; 42: 1027–1030. doi:10.1038/ng.684 PMID: 20972441
5. Pea G, Feron S, Gianfranceschi L, Krajewski P, Pe ME. Gene expression non-additivity in immature ears of a heterotic F1 maize hybrid. Plant Sci. 2008; 174: 17–24. doi:10.1016/j.plantsci.2007.09.005
1. Benjamini Y, Hochberg Y. Controlling the false discovery rate: a practical and powerful approach to multiple testing. J Roy Stat Soc. 1995; 289–300.

2. Hoecker N, Keller B, Muthreich N, Chollet D, Descombes P, Piepho HP, Hochholdinger F. Comparison of maize (Zea mays L.) F1-hybrid and parental inbred line primary root transcriptomes suggests organ-specific patterns of nonadditive gene expression and conserved expression trends. Genetics. 2008; 179: 1275–1283. doi: 10.1535/genetics.108.086278 PMID: 18562640

3. Swanson-Wagner RA, Jia Y, DeCook R, Borsuk LA, Nettleton D, Schnable PS. All possible modes of complementation contribute to transcriptome complexity, diversity, and dynamics. Science. 2009; 326: 1112–1115. doi: 10.1126/science.1178534 PMID: 19965430

4. AC1 Hoen P, Ariyurek Y, Thygesen HH, Vreugdenhil E, Vossen RH, de Menezes RX, et al. Deep sequencing-based expression analysis shows major advances in robustness, resolution and inter-lab portability over five microarray platforms. Nucleic Acids Res. 2008; 36: e141–e141. doi: 10.1093/nar/gkn705 PMID: 18927111

5. Robinson MD, McCarthy DJ, Smyth GK. edgeR: a Bioconductor package for differential expression analysis of digital gene expression data. Bioinformatics. 2010; 26: 139–140. doi: 10.1093/bioinformatics/btp616 PMID: 19910308

6. Schnable PS, Ware D, Fulton RS, Stein JC, Wei F, Pasternak S, et al. The B73 maize genome: complexity, diversity, and dynamics. Science. 2009; 326: 1112–1115. doi: 10.1126/science.1178534 PMID: 19965430

7. AC1 Hoen P, Ariyurek Y, Thygesen HH, Vreugdenhil E, Vossen RH, de Menezes RX, et al. Deep sequencing-based expression analysis shows major advances in robustness, resolution and inter-lab portability over five microarray platforms. Nucleic Acids Res. 2008; 36: e141–e141. doi: 10.1093/nar/gkn705 PMID: 18927111

8. Robinson MD, McCarthy DJ, Smyth GK. edgeR: a Bioconductor package for differential expression analysis of digital gene expression data. Bioinformatics. 2010; 26: 139–140. doi: 10.1093/bioinformatics/btp616 PMID: 19910308

9. Wang L, Feng Z, Wang X, Wang X, Zhang X. DESeq; an R package for identifying differentially expressed genes from RNA-seq data. Bioinformatics. 2010; 26: 136–138. doi: 10.1093/bioinformatics/btp612 PMID: 19855105

10. Benjamini Y, Hochberg Y. Controlling the false discovery rate: a practical and powerful approach to multiple testing. J Roy Stat Soc. 1995; 289–300.

11. Hoecker N, Keller B, Muthreich N, Chollet D, Descombes P, Piepho HP, Hochholdinger F. Comparison of maize (Zea mays L.) F1-hybrid and parental inbred line primary root transcriptomes suggests organ-specific patterns of nonadditive gene expression and conserved expression trends. Genetics. 2008; 179: 1275–1283. doi: 10.1535/genetics.108.086278 PMID: 18562640

12. Swanson-Wagner RA, Jia Y, DeCook R, Borsuk LA, Nettleton D, Schnable PS. All possible modes of complementation contribute to transcriptome complexity, diversity, and dynamics. Science. 2009; 326: 1112–1115. doi: 10.1126/science.1178534 PMID: 19965430

13. Li H, Liu T, Cao Y, Wang L, Zhang Y, Li J, et al. Transcriptomic analysis of maize mature embryos from an elite maize hybrid Zhengdan958 and its parental lines. Plant Growth Regul. 2015; 76: 315–325. doi: 10.1007/s10725-015-0026-1

14. Auger DL, Gray AD, Ream TS, Kato A, Coe EH, Birchler JA. Nonadditive gene expression in diploid and triploid hybrids of maize. Genetics. 2005; 169: 389–397. PMID: 15489529

15. Pumphey M, Bai J, Laudencia-Chingcuanco D, Anderson O, Gill BS. Nonadditive expression of homologous genes is established upon polyploidization in hexaploid wheat. Genetics. 2009; 181: 1147–1157. doi: 10.1534/genetics.108.099941 PMID: 19104075

16. Fujimoto R, Taylor JM, Shirasawa S, Peacock WJ, Dennis ES. Heterosis of Arabidopsis hybrids between C24 and Col is associated with increased photosynthesis capacity. Proc Natl Acad Sci USA. 2012; 109: 7109–7114. doi: 10.1073/pnas.1204464109 PMID: 22493265

17. Meyer RC, Witucka-Wall H, Becher M, Blacha A, Boudichevskaia A, Dörmann P, et al. Heterosis manifestation during early Arabidopsis seedling development is characterized by intermediate gene expression and enhanced metabolic activity in the hybrids. Plant J. 2012; 71: 669–683. doi: 10.1111/j.1365-313X.2012.06021.x PMID: 22472524

18. Hoecker N, Lamkemeyer T, Sarholz B, Paschold A, Fladerer C, Madlung J, et al. Analysis of nonadditive protein accumulation in young primary roots of a maize (Zea mays L.) F1-hybrid compared to its parental inbred lines. Proteomics. 2008; 8: 3892–3894. doi: 10.1002/pmic.200800023 PMID: 18704907

19. West MA, Kim K, Kleibenstein DJ, van Leeuwen H, Michelmore RW, Doerge RW, et al. Global eQTL mapping reveals the complex genetic architecture of transcript-level variation in Arabidopsis. Genetics. 2007; 175: 1441–1450. doi: 10.1534/genetics.106.064972 PMID: 17179097

20. Guo M, Rupe MA, Yang X, Crasta O, Zinselmeyer C, Smith OS, Bowen B. Genome-wide transcript analysis of maize hybrids: allelic additive gene expression and yield heterosis. Theor Appl Genet. 2008; 113: 831–845. doi: 10.1007/s00122-006-0335-x PMID: 16868764

21. Stupar RM, Springer NM. Cis-transcriptional variation in maize inbred lines B73 and Mo17 leads to additive expression patterns in the F1 hybrid. Genetics. 2006; 173: 2199–2210. doi: 10.1534/genetics.106.060999 PMID: 16702414

22. Paschold A, Jia Y, Marcon C, Lund S, Larson NB, Yeh CT, et al. Complementation contributes to transcriptome complexity in maize (Zea mays L.) hybrids relative to their inbred parents. Genome Res. 2012; 22: 2445–2454. doi: 10.1101/gr.138461.112 PMID: 23086286

23. Hochholdinger F, Hoecker N. Towards the molecular basis of heterosis. Trends Plant Sci. 2007; 12: 427–432. doi: 10.1016/j.tplants.2007.08.005 PMID: 17720610

24. Zhang HY, He H, Chen LB, Li L, Liang MZ, Wang XF, et al. A genome-wide transcription analysis reveals a close correlation of promoter INDEL polymorphism and heterotic gene expression in rice hybrids. Mol Plant. 2008; 1: 720–731. doi: 10.1093/mp/sns022 PMID: 19826376
25. Sakai T, Wada T, Ishiguro S, Okada K. RPT2: a signal transducer of the phototropic response in Arabidopsis. Plant Cell. 2000; 12: 225–236. doi: 10.1105/tpc.12.2.225 PMID: 10662859

26. Muller JL, Wolverton C, Ishikawa H, Hangarter RP, Evans ML. Spatial separation of light perception and growth response in maize root phototropism. Plant Cell Environ. 2002; 25: 1191–1196. doi: 10.1046/j.1365-3040.2002.00899.x PMID: 12361060

27. Fuchs I, Philippar K, Ljung K, Sandberg G, Hedrich R. Blue light regulates an auxin-induced K+-channel gene in the maize coleoptile. Proc Natl Acad Sci USA. 2003; 100: 11795–11800. doi: 10.1073/pnas.2032704100 PMID: 14500901

28. Liscum E, Briggs WR. Mutations of Arabidopsis in potential transduction and response components of the phototropic signaling pathway. Plant Physiol. 1996; 112: 291–296. doi: 10.1104/pp.112.1.291 PMID: 8819327

29. Watahiki MK, Tatematu K, Yamamoto M, Yamamoto KT. The MSG1 and AXR1 genes of Arabidopsis are likely to act independently in growth curvature responses of hypocotyls. Planta. 1999; 207: 362–369. PMID: 9951732

30. Guo M, Rupe MA, Danilevskaya ON, et al. Genome-wide mRNA profiling reveals heterochronic allelic variation and a new imprinted gene in hybrid maize endosperm. Plant J. 2003; 36: 30–44. doi: 10.1046/j.1365-313X.2003.01852.x PMID: 12974809

31. Nodine MD, Bartel DP. Maternal and paternal genomes contribute equally to the transcriptome of early plant embryos. Nature. 2012; 482: 94–97. doi: 10.1038/nature10756 PMID: 22266940

32. Li C, Wang M, Wu XM, Chen DH, Lv HJ, Shen JL, et al. THI1, a thiamine thiazole synthase, Interacts with Ca2+-dependent protein kinase gene CPK33 and modulates the S-type anion channels and stomatal closure in Arabidopsis. Plant Physiol. 2015; doi: 10.1104/pp.15.01649

33. Ribeiro DT, Farias LP, De Almeida JD, Kashiwabara PM, Ribeiro AF, Silva-Filho MC, et al. Functional characterization of the thi1 promoter region from Arabidopsis thaliana. J Exp Bot. 2005; 56: 1797–1804. doi: 10.1093/jxb/eri168 PMID: 15897230

34. Ferreira S, Hjeme K, Larsen M, Wünsies G, Larsen P, Fey S, et al. Proteome profiling of Populus euphratica Oliv. upon heat stress. Ann Bot. 2006; 98: 361–377. doi: 10.1093/aob/mcl106 PMID: 16740589

35. Wong CE, Li Y, Labbe A, Guevara D, Nuin P, Whitty B, et al. Transcriptional profiling implicates novel regulatory domain of a plant NADPH oxidase and its functional implications. J Biol Chem. 2010; 285: 18562–18570. doi: 10.1074/jbc.M110.058909 PMID: 20949878

36. Rapala-Kozik M, Kowalska E, Ostrowska K. Modulation of thiamine metabolism in Arabidopsis seedlings under conditions of abiotic stress. J Exp Bot. 2008; 59: 4133–4143. doi: 10.1093/jxb/erm253 PMID: 18940932

37. Rapala-Kozik M, Wolak N, Kujda M, Banas AK. The upregulation of thiamine (vitamin B1) biosynthesis in Arabidopsis thaliana seedlings under salt and osmotic stress conditions is mediated by abscisic acid at the early stages of this stress response. BMC Plant Biol. 2012; 12: 1. doi: 10.1186/1471-2229-12-2 PMID: 22214485

38. Müller K, Carstens AC, Linkies A, Torres MA, Leubner-Metzger G. The NADPH-oxidase AtrbohB plays a role in Arabidopsis seed after-ripening. New Phytol. 2009; 184: 885–897. doi: 10.1111/j.1469-8137.2009.03005.x PMID: 19761145

39. Oda T, Hashimoto H, Kuvabara N, Akashi S, Hayashi K, Kojima C, et al. Structure of the N-terminal regulatory domain of a plant NADPH oxidase and its functional implications. J Biol Chem. 2010; 285: 14345–14345. doi: 10.1074/jbc.M109.058909 PMID: 19864426

40. Swiderski MR, Innes RW. The Arabidopsis PBS1 resistance gene encodes a member of a novel protein kinase subfamily. Plant J. 2001; 26: 101–112. doi: 10.1046/j.1365-313X.2001.01014.x PMID: 11359614

41. Lee KJ, Kim K. The rice serine/threonine protein kinase OsPBL1 (ORYZA SATIVA ARABIDOPSIS PBS1-LIKE 1) is potentially involved in resistance to rice stripe disease. Plant Growth Regul. 2015; 1–9. doi: 10.1007/s10725-015-0036-z

42. Seo KJ, Song E, Chung S, Lee JH. Roles of various cullin-RING E3 ligases involved in hormonal and stress responses in plants. J Plant Biol. 2012; 55: 421–428. doi: 10.1007/s12374-012-0902-4

43. Du H, Wu N, Cui F, You L, Li X, Xiong L. A homolog of ETHYLENE OVERPRODUCER, OsETOL1, differentially modulates drought and submergence tolerance in rice. Plant J. 2014; 78: 834–849. doi: 10.1111/tj.12508 PMID: 24641694

44. Jackson S, Chen ZJ. Genomic and expression plasticity of polyploidy. Curr Opin Plant Biol. 2010; 13: 153–159. doi: 10.1016/j.pbi.2009.11.004 PMID: 20031477

45. Chandler VL, Stam M. (2004). Chromatin conversations: mechanisms and implications of paramutation. Nat Rev Gen. 2004; 5: 532–544. doi: 10.1038/nrg1378
46. Jahnke S, Sarholz B, Thiemann A, Kühr V, Gutiérrez-Marcos JF, Geiger HH, et al. Heterosis in early seed development: a comparative study of F1 embryo and endosperm tissues 6 days after fertilization. Theor Appl Genet. 2010; 120(2), 389–400. doi: 10.1007/s00122-009-1207-y PMID: 19915820