Mapping responsive genomic elements to heat stress in a maize diversity panel

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Abstract

Background: Many plant species exhibit genetic variation for coping with environmental stress. However, there are still limited approaches to effectively uncover the genomic region that regulates distinct responsive patterns of the gene across multiple varieties within the same species under abiotic stress.

Results: By analyzing the transcriptomes of more than 100 maize inbreds, we reveal many cis- and trans-acting eQTLs that influence the expression response to heat stress. The cis-acting eQTLs in response to heat stress are identified in genes with differential responses to heat stress between genotypes as well as genes that are only expressed under heat stress. The cis-acting variants for heat stress-responsive expression likely result from distinct promoter activities, and the differential heat responses of the alleles are confirmed for selected genes using transient expression assays. Global footprinting of transcription factor binding is performed in control and heat stress conditions to document regions with heat-enriched transcription factor binding occupancies.

Conclusions: Footprints enriched near proximal regions of characterized heat-responsive genes in a large association panel can be utilized for prioritizing functional genomic regions that regulate genotype-specific responses under heat stress.

Keywords: Maize, Heat stress, Response eQTL, Response eGene, Chromatin footprints

Background

Changes in gene expression are one mechanism in plants for tolerating changing environmental conditions [1]. Heat is an abiotic stress that can affect plants in many ways. Minor changes in average temperatures can result in lower average yield [2]. Severe heat stress events during reproductive periods can result in major changes in the seed set [3, 4]. Heat stress events at earlier vegetative stages can result in leaf senescence and reduced growth rate [5, 6].

Transcriptome profiling in plants has identified many genes that exhibit altered transcript abundance in response to abiotic stress [7–11]. The heat shock factor (HSF) transcription factors (TFs) are activated in response to heat stress [12] and can regulate the
expression level of many genes through interactions with cis-acting elements. Prior studies have found evidence for enrichment of the HSF binding site motif at upregulated genes in Arabidopsis [13], maize [14], and rice [15]. Some HSFs are stably expressed and become activated in response to heat stress, while many other members of this family are themselves transcriptionally activated in response to heat stress [12, 16, 17]. Several other transcription factors and cis-regulatory elements have also been implicated in the response to heat stress [18]. The prior knowledge of potential mechanisms that create altered gene expression in response to heat stress makes this a good system to study variable responses to heat stress. In addition, this provides a useful system to consider genotype x environment (GxE) interactions for a specific molecular response. GxE interactions are important for developing crop varieties with high levels of productivity in fluctuating environments but our understanding of the molecular mechanisms that drive GxE for specific environments is often limited.

One difficulty in elucidating the mechanisms of gene expression regulation has been our ability to identify and annotate cis-regulatory elements that influence gene expression patterns and levels. In some cases, researchers simply utilize all regions upstream of the transcriptional start site (typically 1–2kb) to search for potential motifs or regulatory elements. Comparisons among related species are also utilized to identify conserved noncoding sequences (CNSs) that may be important for regulation [19, 20]. Recently, many groups have focused on using chromatin properties to identify putative cis-regulatory elements [21–24]. Specifically, regions of accessible chromatin or particular histone modifications can be useful in documenting functionally important cis-regulatory elements [25]. However, for gene expression responses to abiotic stress, it is unclear what proportion of cis-regulatory elements are premarked (accessible prior to the stress) or only exhibit altered accessibility following the stress exposure.

Local adaptation in wild species and breeding to improve the resilience of crop plants likely utilize natural variation in gene expression responses to abiotic stresses [26]. This variation can be attributed to varying activity of trans-acting factors such as transcription factors or due to cis-acting regulatory variation that can alter how specific genes are regulated. Our understanding of the evolutionary sources of variable cis-regulatory elements is limited. The mechanisms by which genes gain, or lose, responsiveness to an environmental stimulus is unclear. It is assumed that a large portion of cis-regulatory information is provided through transcription factor binding sites (TFBSs). While single-nucleotide polymorphisms (SNPs) can change TFBS sequences resulting in altered cis-regulatory information (reviewed by [27]) it is less clear how novel TFBSs and regulatory circuits for response to specific environments are created. It is also possible that insertion / deletion (InDel) polymorphisms, including transposable element (TE) insertions, play a major role in changing the composition of cis-regulatory elements. Several studies have compared the gene expression responses to heat stress in different genotypes [8, 14, 28–30]. While many genes exhibit similar responses to heat stress in different genotypes, there are also examples of genes that exhibit significant allelic variation for responses [8, 14]. The use of hybrid genotypes and allele-specific expression analyses has found evidence for both cis- and trans-regulatory variation. A comparison of the alleles suggests that structural variation may play important roles in driving variable responsiveness but this has not been investigated in detail.
In this study, we utilized a panel of 102 resequenced maize genotypes grown in control and heat stress conditions to study variation of gene expression and associated regulatory genomic elements in response to heat. Both genomic and transcriptomic data were utilized to document eQTLs and their regulated genes. In particular, *cis*-eQTL that influence variable responsiveness to heat stress were identified to document allelic variation for heat-induced expression. This approach was demonstrated as an efficient way to document hundreds of *cis*-allelic variations for heat stress response. The analysis of *cis*-regulatory variation for heat responses using eQTL was complemented with a genome-wide transcription factor footprinting assay in B73. MNase-defined cistrome-Occupancy Analysis (MOA-seq) data [24] was generated in both control and heat stress conditions to document regions with altered chromatin accessibility and transcription factor occupancy. We find that many heat-responsive genes exhibit changes in chromatin accessibility and transcription factor occupancy in the heat stress condition relative to control. The combination of these two approaches provided opportunities to identify high-quality candidate causal regions that could explain the allelic variation for heat responsiveness.

Results

Genome-wide markers associated with photosynthetic parameters

A subpanel of 102 diverse maize genotypes including stiff stalk, non-stiff stalk, and iodent sub-populations representing the Wisconsin Diversity Panel (WiDiv) in maize [31, 32] were selected for characterizing genotype responses to heat stress (Fig. 1a). A set of 1,132,322 variants (1,032,834 SNPs + 99,488 InDels) was employed for this

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**Fig. 1** Phenotypic responses to heat stress within a maize diversity panel. 

**a** Genotypic variants segregated 102 genotypes (including 5 flint/popcorn, 9 iodent, 9 mixed, 32 non-stiff stalk, 20 stiff stalk, 3 tropical, and 24 unknown) employed in this study using principal component analysis. 

**b** Plants showed a distinct pattern using two photosynthetic parameters—Y(NO) and Y(II)—measured under control and heat conditions. 

**c** Associated loci were detected under control and heat conditions using GWAS for Y(NPQ). Two thresholds including 2.2$\times$10$^{-6}$ (blue) and 1.1$\times$10$^{-7}$ (red) were applied for detecting significantly associated loci; 

**d** Same as **c** for Y(NO); 

**e** Same as **c** for Y(II)
population (see “Methods” for filtering details). The genotypes that we used for this study include 3 to 32 inbred lines in each of seven previously characterized sub-populations of maize (Fig. 1a). The selection criteria used to select the subset of inbreds from the larger WiDiv panel retain high levels of diversity while reducing the overall population structure. The first two principal components (PCs) did provide some separation based on these known sub-populations. In total, the first five PCs calculated from the genotype matrix explained 17.1% of population structure variation in the selected 102 genotypes and were used as covariates to control for population structure in subsequent analyses. Seedlings (14 days after sowing) for each of these genotypes were subjected to a 4-h heat stress at 40 °C. The third leaf for each plant was also assessed for three chlorophyll fluorescence-based parameters including (1) light-adapted effective quantum yield of PSII [Y(II)], and two non-photochemical dissipation routes, (2) non-regulated energy dissipation at PSII [Y(NO)], and (3) regulated energy dissipation at PSII [Y(NPQ)] in control and heat stress conditions. Higher Y(II) values indicated less photosynthetic stress, high Y(NPQ) values indicated ongoing photochemical stress that was being mitigated through carotenoid-based energy dissipation, and high Y(NO) values indicated ongoing photochemical stress that was not being mitigated. On average, plants exhibited lower Y(NO) values and higher Y(II) values under the heat stress condition compared to the control condition, while the center of recorded Y(NPQ) did not show a significant shift between control and heat conditions (Fig. 1b; Additional file 1: Fig. S1). All three traits [Y(II), Y(NPQ), and Y(NO)] exhibited higher broad-sense heritability ($H^2$) in control (0.692, 0.682, and 0.716) compared to heat-stressed plants (0.538, 0.563, and 0.642), indicating more variation for these traits in heat stress condition compared to control plants.

The BLUP (best linear unbiased predictor) was applied to represent the phenotypic value of each genotype per trait in each condition in order to remove underlying variation. BLUPs from the chlorophyll fluorescence-based traits (Additional file 2: Table S1) were used to perform GWAS (genome-wide association study) using the same set of >1M variants in this population. We identified candidate association loci under two levels of thresholds ($2.2 \times 10^{-6}$ and $1.1 \times 10^{-7}$, see “Methods”) using the FarmCPU model [33]. Distinct patterns of GWAS significant hits were observed for traits taken from control and heat stress conditions (Fig. 1c–e). In total, 20 candidate genes were identified to be associated with at least one of the significant loci (Additional file 2: Table S2). Several interesting candidate hits included SNPs located within introns of the $farl8$ (FAR1-like transcription factor 8, $Zm00001d017164$) gene that were associated with Y(NO) under the control condition. Orthologs of $Zm00001d017164$ in Arabidopsis—$AT1G52520$ and $AT1G80010$—were demonstrated to show visible phenotypes under abiotic stress [34]. A SNP located ~1kb downstream of the transcription factor—$bbr3$ (BBR/BPC-transcription factor 3, $Zm00001d049800$)—was uniquely identified to be significantly associated with Y(II) measured under the heat condition. $Zm00001d049800$ encoded as a member of the BASIC PENTACYSTEINE (BPC) proteins and its ortholog $AT1G14685$ in Arabidopsis was found to be involved in the salt stress response regulation [35].
Transcriptome responses to heat stress in a subset of WiDiv panel

We proceeded to evaluate the transcriptional response to a heat stress event in the same panel of genotypes. Seedlings (14 days after sowing) were subjected to control conditions or 4 h of heat stress (40 °C) and leaf tissue collected from a pool of 2–3 individuals for each treatment / genotype combination was used for RNA-seq. The RNA-seq data was aligned to the maize B73 AGPv4 reference genome and all samples exhibited high overall alignment rates, ranging from 84.5 to 98.6% (Additional file 2: Table S3). After removing genes with no read counts in any samples, we obtained a gene count matrix with reads assigned to a set of 39,511 maize AGPv4 gene models. In order to assess the consistency between samples during the collection, three biological replicates of B73 were separately collected at early, middle, and late collection time points in each condition. The average pairwise Spearman correlations between replicates were 0.987 and 0.972 in control and heat respectively (Additional file 1: Fig. S2), indicating a high repeatability of samples during collections. The overall variability in the transcriptomes of all samples were assessed using a principal component analysis (Fig. 2a). The first principal component separated samples based on treatment, suggesting a significant impact from the heat stress (Fig. 2a). The second principal component provided a modest separation based on genotypes but did not suggest major contributions of the known population structure.
We hypothesized genotypes with less transcriptome shift (based on principal component 1) in response to heat stress might exhibit a more similar distribution of BLUPs for photosynthetic parameters between control and heat condition. The absolute difference of PC1 between control and heat conditions reflected the transcriptome plasticity of each genotype and could be used to classify genotypes as having large, medium, or small transcriptome responses. The BLUP for each genotype was determined as the genotypic effect after controlling experimental factors for the measured phenotype from either of conditions. We calculated correlations between BLUPs in both control and heat conditions for genotypes with large, medium, or small transcriptome responses to heat stress (Fig. 2b). Overall, for each measured phenotype, we observed an increased correlation coefficient with a larger transcriptome plasticity, suggesting a positive correlation between phenotypic and transcriptomic response in this study (Fig. 2b; Additional file 1: Fig. S3). A GO (Gene Ontology) enrichment analysis on genes with significant upregulation in response to heat in B73 revealed several significant GO enrichments that were associated with heat-responsive functions (i.e., GO:0009408 “response to heat”; GO:0010286 “heat acclimation”; GO:0031072 “heat shock protein binding”) (Additional file 2: Table S4), providing evidence of a strong heat stress response in the B73 samples. In addition to these enrichments for some terms expected for a heat response, there were also a number of other GO terms with significant enrichments, including some without apparent connections to heat stress responses (Additional file 2: Table S4).

The analysis of the full dataset revealed 2628 genes that exhibited consistent upregulation in response to heat in the majority of genotypes (see “Methods” for details). A comparison of the mean response to heat stress and the coefficient of variation (CV) for expression level in heat stress for the 2628 genes highlighted a subset of genes with strong expression response and limited variability in the response for different genotypes (Fig. 2c). Many of the genes with the highest expression response and relatively low variation among genotypes were previously identified as heat shock proteins (HSPs). There were 40 HSPs identified in the maizeGDB database (https://www.maizegdb.org) and 25 of these HSPs were detected to be consistently upregulated with high expression values and low CVs under the heat stress (Fig. 2c). Six additional genes (Zm00001d004243, Zm00001d022630, Zm00001d031436, Zm00001d033990, Zm00001d039933, and Zm00001d048592) that were not annotated as HSPs also exhibited strong and consistent responses to heat stress and might play important roles in response to heat stress. The heat shock factors (HSFs) were a class of transcription factors that were known to play important roles in gene expression responses to heat stress and activated HSPs. Of 31 previously reported non-redundant HSFs in maize [17], 17 of them exhibited detectable expression in our dataset. Eight of these HSFs (ZmHsf02, ZmHsf25, ZmHsf05, ZmHsf08, ZmHsf26, ZmHsf11, ZmHsf03, and ZmHsf13) were consistently upregulated in response to heat stress across a hundred of genotypes (Fig. 2d). Interestingly, ZmHsf06 was downregulated across all genotypes used in this study and also exhibited reduced expression in response to heat stress at multiple timepoints in B73 in a previous study [36] (Additional file 1: Fig. S4). The results suggested that the heat stress event resulted in a robust response to heat stress in this diverse panel and provided an opportunity to further examine variable responses in more diverse genotypes.
Characterization of variable gene expression responses to heat stress

To map regulatory variants that might influence gene expression levels or responses to heat stress in this panel, we retained genes with counts per million (CPM) > 1 in at least 10% genotypes, resulting in 20,255 and 20,306 genes expressed in control and heat conditions, respectively. Of these genes, 19,642 genes were commonly expressed in both conditions, while 613 and 664 genes were uniquely detected as expressed in control or heat conditions. Of 664 genes uniquely detected in heat condition, three were heat shock factors (ZmHsf03, ZmHsf11, and ZmHsf26). We performed eQTL mapping separately using control or heat expression data to map the genomic elements influencing transcript levels of expressed genes. Significantly associated SNPs located within 1Mb distance from the targeted gene were classified as cis-eQTL, while SNPs located >1Mb from the target gene were classified as trans-eQTL. The 1Mb cut-off for cis-eQTL has been used in prior studies of maize eQTL [37, 38] and is far enough to have quite low linkage disequilibrium (r^2<0.05) in our population. Potential trans-eQTL hotspots were assessed by counting the number of significant trans-interactions for each 10-kb window in the genome (Additional file 1: Fig. S5). Several chromosomal regions with at least 10 trans-interactions were commonly detected for both heat and control conditions. Three 10-kb regions on chromosomal 9 were detected to be interacted with gene expression distantly under heat stress. However, the detected “hotspots” had somewhat limited numbers of targets (<20) and the power to detect trans-eQTL hotspots in this study might be limited by our sample size. Consistent with prior studies [39], identified cis-eQTLs in this study could explain more variance of gene expression than identified trans-eQTLs in both control and heat conditions (Additional file 1: Fig. S6). The cis-eQTL analysis identified 10,548 and 10,391 significant cis-eQTL regulated genes (eGenes) for control and heat condition, respectively. The cis-eQTLs detected for genes that were only expressed in heat stress conditions were particularly interesting, as they reflected variable levels of gene expression activation under heat stress. There were 518 of 664 heat expressed only (heo) genes that had significant cis-eQTL (heo-cis-eQTL) and we subsequently referred to these as heo-eGenes (Additional file 2: Table S5; Additional file 1: Fig. S7). The heo-eGenes included a number of transcription factors, such as zhd20, iaa37, abi33, gata17, sbp18, wrky35, blih110, daq27, and ereb196. In addition, two heat shock factors—ZmHsf03 and ZmHsf26—were detected as heo-eGenes.

To identify cis-regulatory variation associated with heat responsiveness per gene in 19,642 commonly expressed genes under both control and heat conditions, a linear mixed model was incorporated with covariates to detect the interaction effect between each pair of cis-eQTL and eGene. To avoid redundancy testing, only the most significant cis-eQTLs in either condition (control or heat) were retained for each gene (two different cis-eQTLs might be selected for testing if they had equivalently significant p-values; see “Methods”), resulting in 15,588 tested gene/cis-eQTL pairs. Of tested cis-eQTLs, the majority had positive correlations for effects between control and heat conditions (Fig. 3a). However, a set of 273 cis-eQTLs with significant differences in effects between control and heat conditions were defined as responsive eQTLs (reQTLs) (Fig. 3a). These reQTLs provided examples of genes with significant differences in response to heat stress due to cis-regulatory variation for responsiveness. Compared to all cis-eQTLs identified in either of conditions or selected top significant cis-eQTLs for reQTLs identification,
reQTLs per se tend to be distributed around TSS (transcriptional start site) regions of regulated genes (Fig. 3b).

The reQTL regulated genes (reGenes) were further assessed and classified into two categories based on the strength of responsiveness under heat stress: reference alleles with greater response or alternative allele with greater response to heat stress. Both of these groups could be further subdivided based on whether the more responsive allele was up- or downregulated in response to heat stress. Of total 258 identified reGenes (Additional file 2: Table S6), 54.4% of these reGenes were examples with reference alleles that have consistently higher response to heat compared to the alternate allele while the other
45.6% exhibited consistently higher response for the alternate allele (Fig. 3c). Four genes related to heat responses (hsp1, hsp22, hsp101, and ZmHsf06) were identified, suggesting potential cis-regulatory elements involved in the regulation of heat responsiveness of these genes. Only limited number of cis-eQTL overlapped with GWAS hits and perhaps this was due to the distinct categories of SNPs could be enriched between GWAS and eQTL mapping [40]. However, the transcription factor—farl8 with SNPs inside its intron regions was significantly associated with Y(II) in control-specific condition and was also identified as a reGene to be regulated by a reQTL within its intron. Given the responsive difference per reGene between genotypes with reference alleles and alternative alleles, responsive patterns of reGenes could be classified into different scenarios according to the median expression value. There were 165 reGenes that exhibited stronger upregulation response for either the reference allele (78) or the alternate allele (87) and these included examples in which both alleles were upregulated (but to different levels) as well as examples in which only one allele was upregulated. For example, an altered InDel from ATA to TACTC in the putative 3′ UTR region of Zm00001d050304 was associated with higher expression under heat stress (Fig. 3d). Similarly, a SNP in the ~1kb promoter region of Zm00001d046322 (Fig. 3e) was strongly associated with higher heat response upregulation in genotypes with reference alleles compared to genotypes with alternative alleles. More distinct regulation patterns were observed as opposite regulations between genotypes with reference and alternative allele (Fig. 3f,g).

One of our goals was to identify sequence variation that might contribute to variable gene responsiveness under heat stress. However, only the most significant eQTL SNPs were selected for performing the analysis to detect the reQTL and its associated reGene. Although the multiple testing burden could be ameliorated this way, the tested reQTL might not reflect a direct causal relationship. Haplotype analysis revealed large haplotype blocks that could be associated with variable responses of most reGenes. For example, Zm00001d042183 was annotated as being involved in the triacylglycerol degradation pathway and we located a significant cis-eQTL on 1443 bp upstream from its TSS for reQTL mapping. The analysis of all SNPs near this gene revealed multiple highly associated SNPs that exhibited high levels of linkage disequilibrium (Fig. 3h). The reGene Zm00001d030549 that was putatively involved in the chlorophyll degradation pathway provides another example of multiple highly associated SNPs within a large haplotype block (Fig. 3i). Even though reGenes and heo-eGenes identified from our approach showed variable responsiveness between genotypes associated with different alleles, multiple SNPs exhibited similar correlations with gene expression responsiveness that fall in a large haplotype region that complicated the ability to determine the underlying causal variant.

Changes in chromatin structure associated with heat responsiveness in maize

Chromatin accessibility provided an additional avenue to document candidate cis-regulatory elements under heat stress. To document changes in accessible chromatin regions with evidence of DNA binding protein footprints in response to heat stress in maize, we generated MNase-defined cistrome-Occupancy Analysis (MOA-seq) data for three biological replicates of B73 plants grown in control and heat stress (4 h at 40 °C). We initially identified peaks of accessible regions using the full MOA-seq reads. Prior work
found that using only the middle 20bp of each MOA-seq read provided likely TF footprints based on known binding sites [24]. The analysis of the high-resolution (center 20bp of each read only) MOA-seq data resulted in a set of smaller peaks (median size of 34bp compared to 179bp for full MOA-seq reads). Over 130,000 MOA-seq footprints were identified in each of the biological replicates in either control or heat condition (Additional file 2: Table S7 and Data S1). TF footprints were frequently identified in regions near the TSS of many genes (Additional file 1: Fig. S8). Three biological replicates of control and heat MOA-seq samples in B73 were compared to identify 13,792 TF footprints that exhibited significant (adjusted p-value <0.05) difference between conditions (Additional file 2: Data S1). These differential TF footprints likely revealed regions of the genome with differential occupancy of TFs or other DNA binding proteins in heat-stressed samples relative to the control samples. The analysis of genes that exhibited heat-stress-inducible expression revealed examples in which there was a distinct difference in MOA-seq coverage shape (Fig. 4a) as well as examples with quantitative differences in TF footprints (Fig. 4b), but there were relatively few truly novel peaks that were only present in heat-stressed samples. There were 4943 differential TF footprints located in putative promoter regions (−2000bp from the TSS per gene)

Fig. 4 Utilization of TF footprints to identify heat response cis-regulatory elements. a An example of differential TF footprints near a B73 heat-responsive gene. Raw read coverages were normalized by RPGC (reads per genomic content) methods. b Same as a, but for another heat-responsive gene, Zm00001d016255 (ZmHSF13). c Normalized MOA-seq signals near 1kb extended regions of putative summits of heat-enriched TF footprints in three biological replicates under control and heat conditions. d Compared 793 commonly upregulated genes with 11,501 background genes for presence and absence of heat-enriched TF footprints. e Comparisons of interaction terms between SNPs located inside and outside of defined TF footprints located within different categories of reGenes and their 1-kb flanking regions. The definition of RefControl, RefHeat, AltControl, and AltHeat were the same as Fig. 3c. ** indicates p-value < 1e −2 and ns indicates insignificant
and another 1486 located within gene regions. The differential TF footprints included more regions with increased accessibility in heat stress (11,140) compared to regions with reduced accessibility in heat stress compared to controls (2,652). The regions with increased MOA-seq coverage in heat stress compared to control might reflect increased binding or occupancy of transcription factors activated by heat stress. A metaplot of MOA-seq read depth at the heat-enriched TF footprints revealed that these regions already exhibited MOA-seq coverage in control samples but had substantially stronger signals in heat-stressed samples (Fig. 4c). Heat-enriched TF footprints were detected within 2000bp of the TSS in 8 HSFs and 18 HSPs. Interestingly, 1353 of 13,792 differential TF footprint regions were located within annotated transposable elements. Some transposable elements with heat-enriched TF footprints were also detected to be upregulated under heat stress in a previous study [30], such as RLG00007Zm000001d00242 (Gypsy retrotransposon), RLC000002Zm000001d10560 (Copia retrotransposon), and RLX06576Zm000001d00001 (unknown retrotransposon) (Additional file 2: Table S8).

The regions that contained heat-enriched MOA-seq footprint peaks were used to perform motif enrichment analysis to identify potential TF binding sites. We identified 26 motifs with putative TF binding activities in the cis-BP database [41] (e.g. AP2, E2F, Myb/SANT, GATA, DoF, bHLH, NAC/NAM) and another 24 motifs that do not contain previously characterized TF binding sites (Additional file 2: Table S9). Prior work had shown that HSFs played important roles in regulation of gene expression in response to heat stress and variants of the HSF binding motif were often enriched near heat-responsive genes [14]. We identified an enriched sequence (GAAGCTTTC) matching HSF motif—HSFB2A—in the heat-enriched TF footprints.

The same tissue samples that were used for B73 MOA-seq were also used to generate B73 RNA-seq data in the same conditions. To eliminate the experimental batch effect for comparing the changes in chromatin accessibility and potential variable regulatory elements in the panel of diverse genotypes, we focused on a subset of 793 genes that were commonly identified as upregulated genes and 11,501 genes that were commonly identified as expressed (mean CPM > 1) but not DEG in both RNA-seq datasets (the matched MOA-seq samples and the three biological replicates of B73 sampled as part of the population study). These 793 genes and 11,501 genes were classified as differentially expressed genes and control genes, respectively. We found that the differentially expressed genes exhibit significant enrichment (p-value < 2.2e−16) for heat-enriched TF footprints within their 2-kb flanking regions compared with control genes (Fig. 4d). We noted that the 2-kb flanking regions of six HSFs (ZmHsf03, ZmHsf05, ZmHsf08, ZmHsf11, ZmHsf25, ZmHsf26) that were consistently upregulated across genotypes contained heat-enriched TF footprints, suggesting potential regulatory regions of these HSFs. This suggested that the heat-enriched TF footprints were often associated with gene activation under heat stress. However, it was noteworthy that 42% of 793 genes that were upregulated in heat stress did not contain heat-enriched TF footprints. These included many examples of quite strong upregulation without evidence for altered TF footprints. The vast majority (>99%) of the 793 upregulated genes had TF footprints within 2kb. This revealed that differential TF footprints were enriched at upregulated genes but that some upregulated genes had consistent TF footprints even with altered expression levels. Together, these results suggested that identified chromatin accessibility changes near heat-upregulated
genes could highlight particular cis-regulatory elements (CREs) that might be involved in transcriptional responses to heat stress.

**Application of B73 TF footprints to identify candidate causal eQTL variants**

We sought to utilize the MOA-seq data to prioritize candidate variants near the reGenes as potential causative changes. There were 258 reGenes that were expressed in many genotypes in both control and heat conditions that exhibited significant variation for heat responsiveness as well as another 518 heo-eGenes that were only expressed in heat-stressed samples that exhibited significant cis-eQTL suggesting variable activation of these genes in response to heat stress. The reGenes were classified into four groups as RefControl \((n=42)\), RefHeat \((n=78)\), AltControl \((n=56)\), and AltHeat \((n=87)\) based on which haplotype exhibited a strong response to heat stress (Alt / Ref) and whether the more responsive allele was up- (Heat) or down- (Control) regulated in response to heat stress (Fig. 3c). Similarly, the heo-eGenes could be classified as RefHeat \((n=313)\) and AltHeat \((n=205)\) based on whether the reference or alternate allele exhibited greater expression in heat-stressed samples. The RefHeat or AltHeat groups of reGenes or heo-eGenes all exhibited upregulation in response to heat and many of these genes (30–37%) have a heat-enriched MOA-seq footprint within 2kb of the gene (Additional file 2: Table S10). These were enriched relative to other expressed genes. In contrast, the Ref-Control and AltControl genes that did not necessarily show upregulation in response to heat stress have only 14–18% of genes with a heat-enriched MOA-seq footprint, which was similar to all expressed genes.

In most cases, there were multiple variants that were significantly associated with the heat response (for reGenes) (Fig. 3h,i) or expression levels in heat stress samples (for heo-eGenes). We assessed whether at least one of the most highly associated (top 3 \(p\)-values) variants was located within a heat-enriched MOA-seq footprint for the subset of genes that contained these footprints. Among the genes that had decreased expression in response to stress (RefControl or AltControl) with a heat-enriched MOA-seq footprint there were only 12.5% with a highly associated variant located within the small footprint region. In contrast, 24% of the RefHeat and 30% of the AltHeat reGenes with heat-enriched TF footprints had at least one highly associated variant that was located in the footprint region (Additional file 2: Table S10). There were 21.7% and 22.6% of the RefHeat and AltHeat heo-eGenes, respectively, that had a highly associated variant within the heat-enriched TF footprints. Through rerunning the reQTL mapping using all variants located in reGenes and their flanking regions, it revealed that significant difference of interaction effects for variants located inside of heat-enriched TF footprints and variants outside of the TF footprints for either RefHeat (Wilcoxon-test, \(p=8.67e^{-3}\)) or AltHeat reGenes (Wilcoxon-test, \(p=2.07e^{-3}\); Fig. 4e; Additional file 2: Table S11). For both reGenes and heo-eGenes, our findings suggested a robust power of heat-enriched MOA-seq signals derived from a single genotype on predicting responsive genomic regions. It was worth noting that in many of the AltHeat genes there was a significant response to heat stress for both the reference and alternate allele, but the alternate allele had a significantly stronger response. In these cases, the B73 allele might still have a heat-enriched footprint and the alternate allele might have a variant that allows for enhanced TF binding. Together, our results suggested overlaps between heat-enriched TF footprints and
identified responsive variants might pinpoint regulatory genomic regions in maize under heat stress.

**Transient expression assays confirm allelic variation for heat response**

We were interested in determining whether the 2-kb proximal regions of the reference and alternate alleles of reGenes would be sufficient to recapitulate the variable responses to heat stress. We selected three pairs of alleles—Zm00001d005114 (B73) / Zm00014a020759 (Mo17), Zm00001d017187 (B73) / Zm00035ab255850 (MS71), and Zm00001d042183 (B73) / Zm00039ab143500 (Oh43) for these experiments. Two of these examples exhibited significant differences for TF footprints in response to heat stress (Fig. 5a,b) while the other gene had ~19.4% increased signals in the TF footprints under heat stress but did not pass the significance filter (Fig. 5c). The two genes with significant differences in the footprints Zm00001d005114 and Zm00001d017187 showed differential response to heat stress for the reference and alternate alleles (Fig. 5d,e). For both of these genes, the reference allele had a much weaker response to heat stress as compared to the alternate allele. The third gene, Zm00001d042183, exhibited no
significant response to heat stress in genotypes carrying the reference allele, but had significant upregulation in genotypes with the alternate allele (Fig. 5f). A comparison of the allele-specific expression in three biological replicates of B73xMo17 or B73xOh43 F1 hybrids was used to validate the allelic ratio in control and heat stress plants for Zm00001d005114 and Zm00001d042183 that were heterozygous for the reference and alternate alleles in these hybrids (Fig. 5g,h). Both of these genes showed a significant difference in the ratio of reference:alternate allele expression in heat stress samples compared to plants grown in control conditions, confirming the cis-allelic variation for heat responses for these alleles.

The 2-kb promoter sequences (including the transcription start site and 5′ UTR) were cloned and fused to a luciferase reporter construct for assessing segregations of their activation roles for reGenes under heat stress. Alignments of the two promoters revealed significant conservation in the region immediately upstream of the transcription start site, but in each case the more upstream regions were not alignable, often due to polymorphic transposon insertions (Additional file 1: Fig. S9). Two of the examples included a differential heat-enriched MOA-seq footprint that overlaps the transcription start site (Additional file 1: Fig. S9a-b). The third example had several MOA-seq footprints that were only detected in heat-stressed samples but these were not identified as significantly heat-enriched (Additional file 1: Fig. S9c). Each of the three pairs of promoters exhibited differences in the level of heat activation using the dual luciferase reporter assay that mirrored the RNA-seq data for the reference and alternate alleles (Fig. 5i–k). In each case, the reference allele had a much smaller response to heat relative to the alternate allele. It is worth noting that the level of activation in response to heat stress was highly variable for the three different promoters with Zm00001d017187 exhibiting much stronger heat response than the other two promoters.

Discussion

Plants frequently encounter abiotic stresses and there is evidence for variable stress responses in many species [42]. This study focused on characterizing the sources of natural variation for gene expression responses to a heat stress event. Our characterization focused on a moderate (non-lethal) 4 h heat stress treatment at 40 °C. While many genes respond rapidly (<30 min) to heat stress, there are quite dynamic changes in the response at early time points [14]. In order to minimize potential complications due to sampling over a ~15-min period for a full population, we focused on an intermediate 4 h heat stress treatment. There are many genes with highly conserved responses to a heat stress in the population that we surveyed. Many of the genes with strong, consistent responses to the heat stress event are heat stress proteins that likely act as chaperones. This confirms that the panel of genotypes that was used in this study were all able to generate a strong heat response. While the genotypes can also respond to a heat stress, there are many genes with variable responses and these provide opportunities to study genotype by environment (GxE) interactions in the context of a specific abiotic stress.

Gene expression levels were highly variable within the population that was used for this study. Prior studies have documented many examples of cis- and trans-eQTL in maize [37, 43–45]. We find similar trends in terms of the proportion of cis- and trans-eQTL as well as the relative magnitude of effects. Relatively few major trans-eQTL
hotspots were identified in our analysis. There are a handful of genomic loci that influence expression of 10–20 other genes but no loci with hundreds of trans-eQTLs. Given that our population consists of maize inbreds that have been subject to artificial selection and are phenologically restricted this may not be all that surprising. It is also likely that our experiment was under-powered to detect trans-eQTL [46].

Mapping sources of cis-variation for heat stress response

A primary goal of our study was to document variation in cis-regulatory elements that provide variable gene expression responses to heat stress. In order to document genes with cis-eQTL for heat responses, we implemented response eQTL analysis that was largely adapted in human studies [47–49]. Other than assessing gene responsive variations between a few genotypes, our approach employed hundreds of genotypes to substantially increase the power of statistical testing for pinpointing potential responsive regulatory regions and avoid transcriptomic instability of a single genotype in response to external heat stress stimuli. This approach identified 258 reGenes that exhibit differential responses to heat stress. While this is a promising approach to document genes with significant variation in responsiveness that is associated with nearby genetic variation, there are a class of genes that cannot be identified using this approach. The set of genes that do not exhibit detectable expression in control conditions that are activated by heat stress (heat stress expressed only by genes) cannot be assessed for variation. We found another set of 518 heat-expressed only genes that have cis-eQTLs in heat stress, suggesting differential activation in response to the heat stress. These two sets of genes represented examples of cis-variable responses to heat stress. We found similar numbers of genes for which the reference or alternative allele exhibited a strong response to the heat stress and the genes include examples of both variable activation and variable repression in response to heat stress. It is worth noting that similar to all GWAS approaches there are limitations in detection of rare alleles that exhibit variable responses. We restricted our analyses to sequence variants with a minor allele frequency of at least 0.10 in this panel. However, the reQTLs that exhibited significant effect had a median minor allele frequency as 0.33.

Applying TF footprinting and transient assays to characterize cis-variable responses

The approaches to document genes with variable response often focus on using the most highly associated sequence variant in either control or heat expression data. However, many studies have shown that there are often multiple sequence variants present together in a haplotype with high levels of linkage disequilibrium [50] and it can be quite challenging to resolve which of these variants are causal for the expression difference. The majority of reGenes and heo-eGenes have multiple nearby genetic variants that are highly associated with variable expression responses. In order to develop prioritized candidate variants that might be responsible for the variable stress-responsive expression, we utilized a recently developed method [24] to perform genome-wide transcription factor footprinting searching.

MOA-seq was performed for B73 in both control and heat stress conditions. This approach identifies ~100,000 regions with TF binding for each replicate in either of conditions. However, ~60% peaks commonly presented in both control and heat TF
footprints using the merged data. A comparison of the control and heat-stressed samples using replicates identified over 13,000 TF footprints with significant variation in heat stress and control conditions with ~80% of these exhibiting increase in heat compared to control. Genes that are upregulated in response to heat exhibit a significant enrichment for TF footprints that are significantly increased in heat stress. However, there are several observations that should limit the simplistic view of novel TF binding being associated with heat-responsive gene expression. First, we found very few examples of truly novel TF footprints in heat stress compared to control. Instead the majority of significant differences in MOA-seq data represent examples in which there is quantitative variation such that a footprint is present in control conditions but becomes stronger in heat stress tissues. This likely suggests changes in average TF occupancy in a cell population following heat stress rather than novel binding events that are not detectable in control conditions. Similar observations were made regarding chromatin accessibility changes in rice plants subjected to varying water availability [51] or in maize plants subjected to heat stress conditions [52]. Second, there are many genes that are upregulated in response to heat stress, including some that are only expressed in response to heat that do not have evidence for variable TF footprints in surrounding regions. Many of these genes have consistent TF footprints in control and heat stress samples. This may suggest that TF occupancy does not change in response to heat stress. Instead, there may be variable activity of the TFs due to post-translational modifications that provide altered regulation.

The MOA-seq data generated from B73 provided insights into one allele that was present in our population. When B73 is the more responsive allele, we can use the MOA-seq data to prioritize potential genetic variants that exhibit significant association with expression responses. All of the reGenes for which B73 has the more responsive allele have at least one highly associated SNP or InDel within their 2-kb flanking regions that is located within a TF footprint. A subset of these are located within TF footprints that exhibit significant differences between heat stress and control samples.

In order to provide further evidence for causal variants that influence heat-responsive gene expression, it is necessary to perform functional assays. We developed a transient protoplast expression system to assess the heat-responsive activity of promoters. We initially focused on utilizing the 2-kb promoter region for the reference and alternate haplotypes to drive expression of a reporter. The dual luciferase assay was then used to compare the relative expression in control and heat-stressed protoplasts. We found that we could recapitulate the allelic variation observed in plants in these assays. In the examples tested to date, we have used the full haplotype of the promoter region and this has included multiple sequence variants. However, this assay can be further utilized to perform targeted sequence changes to monitor the functional impact for each of the specific sequence variants. This system will be useful as we seek to document the molecular basis allelic variation in gene expression responses to a heat stress event.

Heat stress is one of the abiotic stresses that plants encounter regularly. In this study, we have developed approaches to document and characterize natural genomic variation that can regulate heat stress responses of genes at the transcriptomic level. Changes in gene expression are one important mechanism plants utilize to survive abiotic stresses whereas we have limited understanding of the molecular mechanisms that generate
natural variation for allelic responses. Our study provides further insights into the prevalence and characterizing sources of natural variation for gene expression responses to heat stress in maize or other plant species.

**Conclusions**

Two complementary approaches, population transcriptomics and chromatin footprints, were utilized to identify genomic regions that generate allelic variation of gene expression responses to heat stress in maize. This allelic variation for regulating different heat stress responsiveness was confirmed using dual luciferase assays. A better understanding of the allelic variation that can create variable gene expression responses to heat stress provides avenues to improve our predictions of crop responses to abiotic stress and engineer novel gene expression responses to environmental stresses within species. These approaches also provide opportunities to better characterize the transcriptome responses that might produce significant genotype by environment interactions.

**Methods**

**Plant materials, growth, and treatment**

We implemented an approach to select a subset of 120 genotypes from the 509 inbreds from the WiDiv (Wisconsin Diversity) Panel that would retain most of the diversity represented in this panel while reducing the level of population structure. A set of SNPs for this population was retrieved from a previous publication [32] for genotype subtraction. Biallelic SNPs were retained upon multiple thresholdings (no missing data across 509 inbreds; MAF > 0.05; heterozygosity ≤ 0.2). To reduce the computational cost, we randomly selected 1000 SNPs per chromosome for each inbred. The entire set of selected SNPs were used to calculate Euclidean distances between inbreds. Ward's method of hierarchical clustering was then employed to determine the initial 120 clusters. One representative genotype was subtracted from each of 120 clusters (full list in Additional file 2: Table S1). This subset of 120 genotypes retains high levels of diversity but reduces the overall population structure. All genotypes were grown in the growth chamber for 14 days in 30 °C/20 °C 12h/12h day/night cycle. Positions of plants in the growth chamber were randomly shuffled during the plant growth to minimize the microenvironmental effect. On the 14th day, plants subjected to heat stress were treated for 4 h under 40 °C and plants from corresponding genotypes were under control conditions of 30 C in parallel. Control and heat environments were separately implemented in two identical growth chambers. Once the heat treatment was completed, the third leaf per plant was collected and 2–3 plants per genotype were merged to represent one genotype for RNA-seq data generation in each condition. Three biological replicates of B73 were separately inserted into the panel during sample collections for checking experimental repeatability.

**Phenotypic measurements**

The same set of genotypes used in RNA-seq data collection with three replicates were subjected to the same control and heat conditions for chlorophyll fluorescence measurement. Chlorophyll fluorescence parameters including photochemical and non-photochemical quenching were analyzed using a pulse-amplitude modulated chlorophyll
fluorescence camera (MAXI Version IMAGING-PAM M-Series, Heinz Walz, Germany) with a blue (450 nm peak $\lambda$) actinic and measuring light-emitting diode (LED) array and an IMAGE-K7 CCD camera with a Cosmear-Pentax zoom objective lens. All data was acquired in a dark room. Whole-plant samples were dark adapted for 15 min, and the third leaf was detached for a given sample and was directly taped to a non-fluorescent background underneath the LED array with near-infrared LEDs on only. Zoom and focus were adjusted to acquire data on four samples in one image, a control and heat-treated sample from two different genotypes. The camera aperture was set to 1.0 for all samples and the measuring light intensity and gain were at settings of 2 and 3 respectively (frequency = 1.0) to maintain a current fluorescence yield ($F_t$) between 0.100 and 0.180. Once $F_t$ came to equilibrium (10–20 s after turning on the non-actinic blue light source), a saturation light pulse was applied and the minimum fluorescence yield ($F_o$) and maximal fluorescence yield ($F_m$) were recorded. Samples were then exposed to 3 min of actinic light of approximately 950 μmol m$^{-2}$s$^{-1}$ to determine photochemical $Y_{II}$ and non-photochemical quenching parameters $Y_{NPQ}$. The other non-photochemical loss $Y_{NO}$ was determined using the equation of $Y_{II} + Y_{NPQ} + Y_{NO} = 1$ [53]. Data was collected within an area of interest outlining the entire leaf. The maximum value of $Y_{NPQ}$ at 2 min following actinic light treatment was used for treatment comparisons. Due to the capacity of measurement, the entire genotypes were split into 4 blocks using a split block experimental design with B73 as checks in each block under either control or heat condition. BLUPs (best linear unbiased predictions) of $Y_{NPQ}$, $Y_{II}$ and $Y_{NO}$ were separately estimated for control and heat data using a linear mixed model with all effects (genotype, block, replicate) treated as random. In addition, the broad-sense heritability ($H^2$) of each chlorophyll fluorescence parameter per condition was estimated as the proportion of genotypic variation to the total variation.

**Genome-wide association study**

The same set of filtered genomic variants in the studies and BLUPs estimated from phenotype data were used for genome-wide association study (GWAS). The GWAS association model FarmCPU was employed using rMVP v1.0.6 [54] to detect genotype-phenotype association signals. The effective number of variants was determined as 453,641 using GECv2.0 [55] with default settings. Two levels of Bonferroni correction were applied to set the cut-off at $1.10 \times 10^{-7}$ (0.05/453,641) and $2.20 \times 10^{-6}$ (1/453,641).

**RNA-seq data generation and data processing**

RNA-seq data were generated using NovaSeq 6000 in paired-end 150bp mode. Raw reads per library were preprocessed using Trim-galore (Babraham Bioinformatics) with default settings. Preprocessed reads were aligned against the indexed B73 AGPv4 reference genome [56] using HISAT2 [57]. Alignment files were sorted and indexed using SAMtools (v1.9) [58]. The longest transcript was used to represent the individual gene model in the B73 AGPv4.41 version (downloaded from Ensemble). Raw counts per gene were calculated using HTSeq-count (v0.11.2) [59] and normalized into CPM (counts per million mapped reads) value per gene by library sequencing depth using DESeq2 [60]. Genes with absolute log$_2$ fold change greater than 1 and adjusted $p$-value less than 0.05 were considered as differentially expressed genes (DEGs) in B73 samples.
To confirm identities of accessions employed in this study, we employed Broad’s Genome Analysis Toolkit (GATK) [61]—“RNA-seq short variant discovery” pipeline to call SNPs using alignment files generated from HISAT2. Comparing nucleotides on same SNPs generated from RNA-seq and whole-genome resequencing data of identical accessions, we retained accessions with a consistent SNP rate greater than 80% in the study, resulting in 102 genotypes for further analysis.

Comparing gene expressions between control and heat conditions across genotypes

Genes with CPM > 1 in at least 10% genotypes in the panel in either control or heat condition were retained for the analysis. A paired Wilcoxon test was performed to compare expressions between control and heat conditions. The p-value per gene was adjusted for multiple testing corrections using the Benjamin-Hochberg methods. Genes with adjusted p-value < 0.01 were considered statistically significant. Significant genes with log2 transformed heat/control greater than 1 were considered as upregulated genes and less than 1 were considered as downregulated genes. Genes with the median value of log2 transformed ratio of heat to control greater than 0 and 90% quantile of the log2 transformed ratio of heat to control greater than 0 were considered as genes with consistent upregulation in the population.

GO enrichment analysis

Upregulated genes identified in B73 replicates inserted into the panel transcriptomic experiment were considered as the tested gene set. GO terms per gene were retrieved from the maize GAMER dataset [62] for maize AGPv4 gene models. GOATOOLS [63] was implemented to perform the GO enrichment analysis. Any GO term with a Bonferroni-corrected p-value less than 0.05 was considered significant.

Genotyping and eQTL mapping analysis

Missing genotype SNP calls of 509 inbreds were imputed using Beagle (version 05.Apr.21) [64]. SNPs of the subset inbreds consisting of 102 genotypes employed in this study were subtracted and retained with minor allele frequency (MAF) ≥ 0.1 and heterozygous rate ≤ 0.1. Highly correlated SNPs were pruned using PLINK [65] (--indep-pairwise 50 5 0.99). InDels from the same prior study were retrieved and only InDels with no missing data on studied genotypes were retained to merge with filtered SNPs together as the final set of 1,132,322 genotype variants. The first five principal components (PCs) were calculated using the R function prcomp. Each gene with CPM > 1 in at least 10% genotypes was retained for analysis in either control or heat condition. CPM values per gene under control or heat condition were transformed into a normal distribution using the R package bestNormalize (https://github.com/petersonR/bestNormalize) separately. Twenty-five hidden factors were separately inferred from control and heat expression data using PEER [66] for controlling variance. SNPs located nearby 1Mb of gene regions were classified as cis-eQTLs, otherwise as trans-eQTLs similar to cutoffs used in prior maize eQTL studies [37, 38]. The R package MatrixeQTL [67] was used for eQTL mapping in controlling 5 PCs and 25 hidden factors in either control or heat conditions. SNPs with Benjamini-Hochberg-corrected p-value < 0.01 in each condition were considered as eQTLs. Genes with significantly associated eQTLs were considered
as eGenes. The proportion of variance explained by each identified eQTL was calculated based upon the MatrixeQTL manual.

**Trans-eQTL hotspot identification**

Trans-eQTLs identified by eQTL mapping in each condition were used for detecting trans-eQTL hotspots. The maize genome was equally segmented into 10-kb bins. Any SNP within each 10-kb bin targeted greater or equal to 3 genes remotely (>1Mb) was considered as candidate trans-eQTL. Each 10kb bin with more than 10 targeted unique gene sets was considered as a trans-eQTL hotspot. Based on genes targeted by trans-eQTLs in control and heat conditions, we split the number of targeted genes by trans-eQTL hotspots into common targeted genes by two conditions, uniquely targeted genes by control and uniquely targeted genes by heat.

**Response eQTL (reQTL) detection**

We modified a prior published method [49] for prioritizing candidate cis-eQTLs for reQTL detections in this data. Specifically, we focused on genes that were commonly expressed in both of conditions and narrowed candidate cis-eQTLs for reducing multiple testing burden based on following criteria: (1) For each eGene per condition, we selected the top significant cis-eQTL. If multiple top significant cis-eQTLs existed, we randomly picked one for testing; (2) for eGene with the identical cis-eQTL in both conditions, we retained it for testing; (3) for eGene with non-overlapped cis-eQTLs between two conditions, we picked one cis-eQTL randomly if two cis-eQTLs from control and heat are in high LD (> 0.8), or we retained both of cis-eQTLs if they are in low LD (≤ 0.8, 4) for eGene with only detected cis-eQTL in one condition but not the other, we retained this cis-eQTL for the analysis. Selected candidate cis-eQTLs were input into a linear mixed model below for detecting reQTLs using the R package lmerTest [68]:

$$E_i = d_i + c_i + d_1^i + \cdots + d_30^i + c_i + \cdots + d_30^i \times c_i + d_i \times c_i + (1|G_i)$$

where $i$ was the $i$th sample, $E_i$ was considered as the normalized expression value for the $i$th sample, $d_i$ was the cis-eQTL allele dosage for $i$th sample, $c_i$ was experimental condition for the $i$th sample (control or heat), $d_1^i$ to $d_30^i$ represented 30 covariates employed in this study (5 principal components + 25 PEER hidden factors), $d_i \times c_i$ was the interaction between cis-eQTL and condition and $(1|G_i)$ was the random effect of individual genotype. The $p$-value of the interaction term was estimated using the Satterthwaite and Kenward-Roger methods and then adjusted using the Bonferroni methods. The cis-eQTL relative to its targeted gene with adjusted $p$-value less than 0.01 were considered as reQTLs.

**MOA-seq library construction and data processing**

To assess the changeable chromatin accessibility under heat stress, we collected leaf tissues from plants grown in the same growth stage under the same control and 4 h 40 °C heat conditions as the large panel experiment. Collected tissues were processed for MOA-seq library constructions following a previous protocol [24]. For B73 samples, the genome index of the reference genome was built using STAR (v2.7.9a) [69]. Raw reads of MOA-seq data were preprocessed using SeqPurge (v2019_09) [70] with
parameters of “-min_len 20 -threads 20 -qcut 0 -prefetch 50000.” Overlapping paired reads were merged into single reads using FLASH (v1.2.11) [71]. Processed reads were aligned against the indexed reference genome using STAR (v2.7.9a) [69]. As STAR is designed to map RNA, we set the flag --alignIntronMax 1 for DNA, this presence and Introns from being allowed / expected by STAR. Alignment files were converted into bam formats using SAMtools (v1.9) [58]. Alignment fragments with less than 81 bp and MAPQ as 255 were retained for further analysis. To generate high-resolution maps, each read was shortened to 20bp centered around the middle of the read. Read shortening was performed using awk: for reads with uneven number of bases, the middle base was taken and then read extended 10bp to each site. For reads with even numbers of bases, one of the two middle bases was chosen randomly and the reads were extended 10bp to each site. Each filtered alignment file was converted into bigwig format using bedtools v2.29.2 [72]. The effective genome size was determined using the k-mer size of 60. Coverage of alignment files were normalized to visualize using bamCoverage RPGC (reads per genome coverage) normalization function. MOA-seq TF footprints were determined using the “macs3 callpeak” function (https://github.com/macs3-project/MACS) with parameters of “-s 20 --min-length 20 --max-gap 40 --nomodel --extsize 20 --keep-dup all --buffer-size 10000000.” Distance of TF footprints to TSS was calculated using annotatePeak.pl function [73]. We then employed the “DiffBind” (v2.12.0) function in R [74] to detect differential footprints between control and heat in B73 samples with FDR < 0.05.

Motif discoveries
Summits of heat-enriched TF footprints were redefined using the “macs3 refinepeak” function and merged B73 replicates in heat condition. For each summit, flanking regions were extended according to the median length of TF footprints and nucleotide sequences were extracted from the B73 AGPv4 genome in the given region. Identified heat-enriched MOA-seq footprint sequences were input for STREME (v5.4.1) [75] to identify enriched motifs with shuffled genomic sequences as the background. Identified motifs were compared with PWMs (position weight matrix) in cis-BP public TF database [38] using tomtom (version 5.0.5) [76]. Searched motifs with p-value less than 0.05 were considered as significant and only the most significant motif matching the database was considered as the target motif for the query PWM.

RNA-seq data generation and processing in maize hybrids
To evaluate allele-specific gene expression, we generated RNA-seq data of two selected maize hybrids including Mo17 × B73 and Oh43 × B73 grown together with B73 samples for MOA-seq data generation. All parental lines were also included in our heat stress panel experiment. SNPs between two alleles in one hybrid were retrieved from the raw SNP sets employed in the subpanel. The SNP set per hybrid was separately employed to mask the B73 AGPv4 reference genome using bedtools v2.29.2 [72]. Raw RNA-seq reads generated from each hybrid were aligned to the respective masked reference genome in B73 AGPv4 coordinates using the same steps as above. Allele-specific alignments per sample were then split using the SNPsplit (v0.5.0) [77] and allele-specific read counts were calculated for each split alignment file using HTSeq-count (v0.11.2) [59].
Dual luciferase reporter assay for candidate reQTLs/reGenes

We amplified 2-kb regions upstream of the start codon for reGenes pairs that displayed differential heat responsiveness between alleles. These regions were cloned into a minimal vector driving a Firefly Luciferase reporter through a combination of NEBuilder HiFi Assembly (E5520S) and traditional restriction enzyme/ligation-based approaches. Both approaches leveraged a backbone double-digested with SpeI-HF (R3133S) and SacI-HF (R3156S), dephosphorylated (Quick CIP, M0525S), and gel extracted (Omega E.Z.N.A. Gel Extraction kit, D2500-01) prior to ligation or HiFi assembly. HiFi reactions were designed and conducted as suggested in the product manual. Traditional clones were generated by amplifying regions of interest and creating corresponding SpeI/SacI restriction enzyme sites. PCR products were then purified (Zymo DNA Clean & Concentrator-5 kit, D4013), double-digested with SpeI/SacI, and ligated (T4 DNA Ligase, M0202S) into the prepared backbone. Colonies were screened and sequenced through Primordium plasmid sequencing (www.primordiumlabs.com) prior to further use.

The conditional reporter was co-transformed into maize leaf protoplasts alongside a Renilla Luciferase driven by a constitutive 35S reporter in three separate transformation events, using ~200k protoplasts per transformation. Protoplasts were generated as previously described [78], were transformed with a total of 10μg of plasmid DNA (8μg conditional reporter, 2μg constitutive reporter), allowed to recover for 16 h following transformation, and then evenly split and either subjected to a +10°C heat stress in a water bath or left at room temperature for 3 h. Protoplasts were spun down and snap frozen at the end of the stress event. Dual luciferase assays were conducted per manufacturer instructions (Promega Dual-Luciferase Reporter Assay System, E1960), and measurements were taken on a Promega GloMax Explorer Plate Reader (GM3500).

Supplementary Information

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Review history

The review history is available as Additional file 3.

Peer review information

Wenjing She was the primary editor of this article and managed its editorial process and peer review in collaboration with the rest of the editorial team.

Authors' contributions

Z.L. and N.M.S. conceived and designed the experiments. Z.A.M. performed the dual luciferase assay experiment, and Z.L. and D.P. collected the phenotype data. J.E. and T.H. performed MOA-seq library constructions and MOA-seq data processing. Z.L., Z.A.M., D.P., J.E., and T.H. analyzed the data, and Z.L., Z.A.M., D.P., and N.M.S. wrote the article. All author(s) read and approved the final manuscript.
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Availability of data and materials
Raw read RNA-seq data of heat stress panel have been deposited in the NCBI SRA database under the Bioproject ID PRJNA831425 [79]. MOA-seq data and associated RNA-seq data have been deposited in the NCBI SRA database under the Bioproject ID PRJNA499202 [80]. The UCSC genome browser: https://genome.ucsc.edu/s/shanwai1234/B73v4 MOAseq was set up for visualizing MOA-seq and associated RNA-seq coverage of B73 samples in B73 AGPv4 genome coordinates. Codes and associated CPM values per gene used in this study were deposited in the Github repository (https://github.com/shanwai1234/MaizeHeatStress) under the GNU General Public License v3.0 [81] and the Zenodo repository (https://doi.org/10.5281/zenodo.7239565) [82].

Declarations
Ethics approval
Not applicable.

Competing interests
The authors declared that they have no competing interests.

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