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A cartography of the maize regulatory genome
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Genome-wide mapping of transcriptional enhancer candidates using DNA and chromatin features in maize

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Abstract

Background: While most cells in multicellular organisms carry the same genetic information, in each cell-type only a subset of genes is being transcribed. Such differentiation in gene expression depends, for a large part, on the activation and repression of regulatory sequences, including transcriptional enhancers. Transcriptional enhancers can be located tens of kilobases from their target genes, but display characteristic chromatin and DNA features, allowing their identification by genome-wide profiling. Here we show that integration of chromatin characteristics can be applied to predict distal enhancer candidates in *Zea mays* thereby providing a basis for a better understanding of gene regulation in this important crop plant.

Result: To predict transcriptional enhancers in the crop plant maize (*Zea mays* L. ssp. mays), we integrated available genome-wide DNA methylation data with newly generated maps for chromatin accessibility and histone 3 lysine 9 acetylation (H3K9ac) enrichment in young seedling and husk tissue. Approximately 1,500 intergenic regions, displaying low DNA methylation, high chromatin accessibility and H3K9ac enrichment, were classified as enhancer candidates. Based on their chromatin profiles, candidate sequences can be classified into four subcategories. Tissue-specificity of enhancer candidates is defined based on the tissues in which they are identified, and putative target genes are assigned based on tissue-specific expression patterns of flanking genes.

Conclusions: Our method identifies three previously identified distal enhancers in maize, validating the new set of enhancer candidates and enlarging the toolbox for the functional characterization of gene regulation in the highly repetitive maize genome.
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Background

Successful differentiation of zygotes into different cell-types that make up a complex multicellular organism requires flexibility to respond to environmental cues, but also a tight control of gene expression during developmental processes. Regulation of gene expression, among others, depends on a complex network of sequence-specific transcription factors (TFs) as well as protein factors that can read or write chromatin modifications [1,2]. In addition, gene expression regulation depends on genetic information encoded within cis-regulatory regions such as transcriptional promoters and enhancers, which contain multiple TF binding sites and display particular DNA and chromatin features [3]. In the last decade, genome-wide approaches in animals have identified thousands of enhancers (see e.g. [4]). Mutations in enhancers are known to cause developmental defects, cancer or other diseases [5–8], emphasizing the crucial role of enhancers in gene expression regulation. High-throughput genome-wide enhancer identification in plant species only started recently, and only a small number of enhancers were thoroughly studied in plant species [9,10], including enhancers for booster1 (b1) [11,12], teosinte branched1 (tb1) [13,14], pericarp color1 (p1) [15] in maize, Block C for FLOWERING LOCUS T in Arabidopsis thaliana (Arabidopsis) [16,17], and the enhancers for the chlorophyll a/b-binding protein gene AB80 and pea plastocyanin gene in Pisum sativum [18,19]. So far, few genome-wide approaches to identify cis-regulatory sequences in plants have been reported, i.e. in Arabidopsis, Oryza sativa (rice) and maize [20–22]. Although multiple studies in plants reported genome-wide profiles for different chromatin features, only one, in Arabidopsis, aimed at discovering enhancers [20].

Enhancers can be located up- or downstream of their target genes, and physically interact with their target genes to regulate gene expression [23,24]. They are typically short DNA sequences of 50 to 1,000 bps that are bound by TFs and characterised by an accessible chromatin structure, especially when they are actively involved in regulating gene expression [25,26]. Based on extensive studies in animals, active enhancers are associated with low DNA methylation and histone modifications such as acetylation of lysines 9, 14 and 27 of histone H3 (H3K9ac, H3K14ac and H3K27ac) [27–30]. Mono-methylation of lysine 4 of histone H3 (H3K4me1) is enriched at enhancers regardless of their activity [27,28]. Low DNA methylation has been reported to positively correlate with enhancer activity and also used to predict enhancers [29,31]. Although limited data is currently available, similar DNA and chromatin features were observed at known plant enhancers, indicating that these marks may, at least partially, be conserved between animals and plants [9].
Another feature reported for animal enhancers is bi-directional transcription, producing so-called enhancer RNA (eRNA). eRNA expression levels positively correlate with enhancer target gene expression levels [4,32], which can help to link enhancers to their target genes. The function of eRNAs is not yet clear, but some of them have been reported to play a role in the recruitment of TFs to enhancers or in the formation of enhancer-promoter interactions [33,34]. The purpose of this study was a genome-wide identification of active intergenic enhancers in maize, and to find their most likely target genes by integrating tissue-specific chromatin features and differential gene expression levels. To do so, we identified regions with low DNA methylation levels using published bisulfite-sequencing (BS-seq) data [35], and measured chromatin accessibility using DNase-seq, H3K9 acetylation using Chromatin Immunoprecipitation (ChIP)-seq, and differential expression using RNA-seq in V2 stage inner stem tissue (V2-IST) and husk tissue. We identified approximately 1,500 intergenic enhancer candidates and defined their tissue-specificity based on the presence or absence of DNase I hypersensitivity and H3K9ac enrichment signals. Our pipeline was validated by the detection of three previously identified (putative) enhancers, regulating the expression of \( b1 \), \( bx1 \) and \( tb1 \).

**Results**

**Selection of H3K9ac as best suited histone modification to identify active enhancers in maize**

In mammals, several histone modifications such as H3K27ac, H3K9ac and H3K4me1 were shown to mark active enhancers [27,28,30]. To define which of these histone modifications indicate best active enhancers in maize, we examined the enrichment of H3K27ac, H3K9ac and H3K4me1 at the hepta-repeat enhancer and other cis-regulatory sequences present at the \( B-I \) allele of the \( b1 \) gene. ChIP was performed on inner stem tissue from V5 maize seedlings (V5-IST) and husk tissue. The hepta-repeat enhancer of \( B-I \), located 100 kb upstream of the \( b1 \) transcription start site (TSS), is inactive in V5-IST and active in husk leaves [36]. Previously, the hepta-repeat enhancer and regulatory sequences ~45 kb upstream of \( b1 \) were shown to be enriched with H3K9K14ac when active [36]. The results presented here (Figure 1) indicated that the enrichment in both H3K9ac and H3K27ac was significantly higher in husk compared to V5-IST at the hepta-repeat enhancer (R3 and R6), ~45kb upstream regulatory sequences (g) and the untranslated 5' region of \( b1 \) (UTR). Based on these results, both H3K9ac and H3K27ac appeared to mark active enhancers. In contrast, H3K4me1 enrichment levels were relatively low throughout the intergenic \( b1 \) region in both V5-IST and husk tissues. In addition, at the
coding region, H3K4me1 enrichment levels were higher in low \( b1 \) expressing V5-IST than in high expressing husk tissue. Therefore, in contrast to animal systems [27,37], H3K4me1 is probably not suited to identify enhancers in maize. Since the enrichment at the enhancer region in husk relative to V5-IST tissue was highest for H3K9ac, we chose this histone modification to identify active enhancers genome-wide.

Figure 1. ChIP-qPCR analysis at \( b1 \) for H3K27ac, H3K9ac and H3K4me1. (A) Schematic representation of the \( b1 \) locus. Vertical arrows with letters indicate the regions examined by ChIP-qPCR. The \( b1 \) hepta-repeat enhancer is indicated with seven black triangles, the \( b1 \) coding region by a black box, and the TSS by a bent arrow. Grey bars represent TEs and other repetitive sequences. (B) Enrichment of H3K27ac, H3K9ac and H3K4me1 at the \( b1 \) locus relative to the enrichment at the maize \textit{actin 1} locus (\textit{actin}). Error bars represent the standard error of the mean for two (H3K9ac, H3K4me1) or three (H3K27ac) biological replicates.

An integrated pipeline to identify tissue-specific enhancers in maize

DNase-seq, H3K9ac ChIP-seq and RNA-seq experiments were carried out in two tissues, V2-IST and husk, isolated from the reference inbred line B73 (Additional file 1: Figure S1). These tissues were selected to identify tissue- as well as developmental stage-specific enhancers. Our study included material grown at two different locations (DNase-seq and H3K9ac ChIP-seq were performed at the Max Planck Institute for Plant Breeding Research and the University of Amsterdam, respectively), therefore, we performed RNA-seq experiments for each tissue in six
biological replicates, three per location. Comparison of gene expression levels between replicates in Reads Per Kilobase of transcript per Million mapped reads (RPKM) revealed high correlations among replicates between the two locations (Additional file 1: Figure S2). This high correlation between replicates and locations indicated the data were comparable and implied that the chromatin states of the plants from both locations were similar. Gene expression levels and significant differential expression levels were calculated, taking the variability among six replicates into account. The genes determined as significantly differentially expressed thus showed statistically significant differences in their expression levels at both locations.

After pre-processing of the data, our enhancer prediction pipeline consisted of three steps of data integration (Figure 2). First, enriched chromatin or DNA features were identified for three genome-wide data sets. In addition to calling DNase-seq and H3K9ac ChIP-seq peaks from our own data sets, we identified low and unmethylated DNA regions (LUMRs) by re-analysing published bisulfite (BS)-seq data [35]. By taking an overlap between the three data sets, regions displaying all three features were selected as enhancer candidate regions. We focused on intergenic enhancer candidates, excluding promoter regions, as chromatin profiles of enhancers located in proximity of, and within coding regions are more likely to overlap with chromatin profiles of genic regions, making it difficult to disentangle the underlying regulatory regions. Enhancer candidates predicted in only one tissue were defined as tissue-specific candidates. Transposable elements (TEs) were included in our analysis as some of them had been shown or suggested to act as enhancers in maize and other organisms [13,38]. The second step involved determining the degree of tissue-specificity of the candidates identified in both tissues by ranking the candidates based on signal intensity differences between the two tissues. This was done for both chromatin accessibility and H3K9ac enrichment, followed by summing the ranks and re-ranking. The last step assigned target genes to enhancer candidates, assuming that enhancers most likely regulate genes located directly up- or down-stream and that gene expression and active chromatin marks at enhancers are positively correlated.
Figure 2. Overall workflow of this study. Firstly, chromatin accessibility data from DNase-seq, H3K9ac enrichment data from ChIP-seq, and DNA methylation data from BS-seq were analysed individually. Secondly, the data on accessible regions, H3K9ac-enriched regions and low DNA methylated regions were integrated to predict enhancers. Thirdly, the enhancer candidates were ranked based on signal intensity differences of the chromatin accessibility and H3K9ac enrichment data between V2-IST and husk tissue. Finally, enhancer candidates were linked to their putative target genes based on their tissue specificity and on the differential expression of flanking genes determined by RNA-seq data. For shared candidates, adjacent genes being expressed in both tissues were associated.
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**Distribution of chromatin features in the uniquely mappable part of the maize genome**

To identify chromatin accessibility, H3K9ac enrichment, and low DNA methylation within the genome, we partitioned the genic and intergenic regions of the genome in six sub-categories: promoters, exons, introns, flanking and distal intergenic regions, and TEs (Figure 3A). Gene annotations were taken from the maize B73 annotation version 4 (AGPv4 assembly [39]) from Ensembl Plants [40]. Only intergenic TEs were considered in our study; TEs present in introns were counted as "introns". Promoter regions were defined as 1 kb upstream to 200 bp downstream from the TSS, therefore including the first nucleosome downstream of the TSS. The composition of the B73 maize genome was quantified by counting the numbers of mega bases in each genomic region (Figure 3B). Since 85% of the maize genome is highly repetitive [41], an important fraction of the next generation sequencing reads could not be mapped uniquely (Additional file 1: Table S1), which prevented enhancer identification in repetitive genomic regions. We determined the uniquely mappable parts of the genome by performing an all-against-all alignment for theoretical 93 bp single-end reads, allowing a maximum of 2 mismatches using the Uniqueome pipeline [42], which estimates the fraction of uniquely mapped reads for each nucleotide (Figure 3C). In the uniquely mappable genome, the proportion of TEs was reduced to approximately a quarter of the assembled genome.

**9,212 intergenic DHSs are potential cis-regulatory elements**

DNase I hypersensitive sites (DHSs) are genomic regions that are more sensitive to DNase I endonuclease activity compared to flanking regions due to a lower nucleosome density [43]. The mapping of DHSs by DNase-seq is a powerful approach to identify cis-regulatory regions, including enhancers, and has been used in many organisms including plants [20,25,44–46].
Figure 3. Genomic composition and distribution of features. (A) Definition of genomic regions. Promoters are defined from 1 kb up- to 200 bp downstream from the TSSs, flanking regions are 4 kb upstream from the promoters and 5 kb downstream from the Transcription Termination Sites (TTSs). TE refers to transposable elements, and distal refers to intergenic regions that are more than 5kb away from genic regions and are not TEs. (B) Composition of the entire maize genome according to AGPv4 and (C) the uniquely mappable genome. Distribution of (D and F) DHSs, (H and J) H3K9ac, (L) LUMRs and (N-O) enhancer candidates over the different genomic regions, and (E, G, I, K and M) the fractions (Mbp/Mbp, from 0 to 1, y-axes) the different features (x-axes) occupy at the various genomic regions in the uniquely mappable genome. The grey bars indicate the fraction of overall occupancy in the uniquely mappable genome.
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DNase-seq experiments were performed in two biological replicates for both V2-IST and husk tissue (Additional file 1: Table S1). To take the intrinsic digestion bias of DNase I into account, we also included a control sample generated by digesting B73 genomic DNA with DNase I. After mapping the reads obtained from each library, DHSs were identified for each library using MACS2 peak calling [47].

Data reproducibility between biological replicates was examined by counting the number of overlapping DHSs identified for all the possible combinations of replicates (Additional file 1: Table S2). This comparison showed that 54 to 92% of DHSs overlapped by at least 1 bp between replicates. The overlap between the two V2-IST replicates was the lowest (54% of the 35,906 V2-IST_2 peaks were overlapping with the 21,309 V2-IST_1 peaks) as 1.5 times more peaks were identified in the V2-IST_2 sample. The overlap between peaks identified in V2-IST and in husk samples appeared quite large (e.g. 80% of the peaks identified in V2-IST_1 were also observed in Husk_1), indicating that most DHSs are not tissue-specific. To select for high confidence DHSs in both V2-IST and husk tissue, only DHSs overlapping by at least 70% of their lengths between replicates were kept for further analysis. For signal intensity analysis, the reads in all biological replicates were pooled per tissue to estimate the overall coverage of the reads.

We correlated DNase I hypersensitivity and gene expression levels in gene bodies and their immediate 1 kb flanking regions for additional validation of the data set. For each tissue, genes were binned according to their gene expression levels, and the average DNase I hypersensitivity, measured in number of read counts per million mapped reads (RPM), was calculated for each bin using bwtools [48] (Figure 4A and B). A positive correlation between expression levels and DNase-seq coverage over genic regions was observed, especially directly upstream of the TSSs and transcription termination sites (TTSs). Chromatin at gene bodies was rather inaccessible among the gradient of gene expression. Presence of DHSs at TSSs and a positive correlation with expression levels observed in our dataset confirm previous observations in both animals and plants [21,26,49–51].

The number of DHSs per genomic region was counted to examine their fraction per genomic region (Figure 3D, F). When comparing the distributions of DHSs to a randomised distribution within the mappable genome (Additional file 1: Figure S3A and B), we observed a clear over-representation of DHSs at promoters (p-value < 0.001; permutation test). Still, 43% of DHSs, in total 9,212 out of 21,445, were in intergenic regions excluding promoters (Figure 3D, F): 7,802 in V2-IST, 7,123 in husk and 5,130 shared between both tissues (Table 1A). In addition,
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the fraction of the genome scored as DHS (in Mbp) was calculated for each genomic category. In total, DHSs occupied about 2% of the mappable genome in both tissues (Figure 3E, G). DHSs occupied 10 and 8% of the total mappable promoter regions in V2-IST and husk, respectively.

Table 1: Intergenic regions of interest. (A) DHSs and (B) H3K9ac-enriched regions and number (and percentage in parentheses) of these regions overlapping with other features (LUMRs, H3K9ac or DHSs) in each tissue (V2-IST and Husk), common to both tissues (Common) and the sum of the tissues (Total).

(A) Number of intergenic DHSs

| Tissue | All | Overlapping with LUMRs | Overlapping with H3K9ac |
|--------|-----|------------------------|------------------------|
| V2-IST | 7802 | 7653 (98.1%) | 344 (4.4%) |
| Husk   | 7123 | 6997 (97.4%) | 1505 (21.1%) |
| Common | 5130 | 5013 (97.7%) | 202 (3.9%) |
| Total  | 9212 | 9057 (98.3%) | 2030 (22.0%) |

(B) Number of intergenic H3K9ac enriched regions peaks

| Tissue | All | Overlapping with LUMRs | Overlapping with DHSs |
|--------|-----|------------------------|----------------------|
| V2-IST | 3115 | 3093 (99.3%) | 323 (10.4%) |
| Husk   | 6213 | 6170 (99.3%) | 998 (16.1%) |
| Common | 2668 | 2652 (99.4%) | 184 (6.9%) |
| Total  | 6511 | 6466 (99.3%) | 1454 (22.3%) |
ChIP-seq identifies 6,511 intergenic H3K9ac-enriched regions

ChIP-seq H3K9ac data were obtained from two and three biological replicates for V2-IST and husk tissue, respectively. The reads were aligned to the AGPv4 B73 reference genome and H3K9ac-enriched regions were identified, taking the input sample into account, by peak calling for each replicate using MACS2 [47].

To examine the reproducibility between replicates, overlapping H3K9ac-enriched regions were counted for all replicate combinations, showing 62 to 96% overlap within a tissue (Additional file 1: Table S3). As for the DNase-seq data, H3K9ac-enriched regions with an overlap in length of at least 70% between all replicates were kept for further analysis, and reads in replicates were pooled for coverage calculation in each tissue. We correlated gene expression levels with H3K9ac enrichment levels across gene bodies and their 1 kb flanking regions (Figure 4C, D) and observed a peak of H3K9ac enrichment immediately after the TSS and increased levels across the gene bodies compared to gene flanking regions. At the TSS peak region, gene expression and H3K9ac levels showed a parabolic correlation, showing saturation for higher bins and signal reduction for the highest one. In gene bodies, H3K9ac was lower for the three highest bins than for the three following bins. Previous studies in yeast and maize have reported a genome-wide loss of nucleosomes at highly expressed genes [26,52]. Reduced nucleosome levels could explain the reduction in H3K9ac observed at highly expressed maize genes.

Correlations between enrichment levels of H3K9ac 3’ of the TSS and gene expression levels have been previously reported [30,53,54]. Our data suggest that H3K9ac enrichment levels reached saturation for genes with high expression levels.

To estimate the number of potential intergenic enhancer candidates from the H3K9ac data sets, the genomic distribution of H3K9ac-enriched regions was examined by counting the numbers of H3K9ac-enriched regions in the different types of genomic regions (Figure 3A, H and J). As seen for DHSs, a clear over-representation of H3K9ac-enriched regions at promoters was observed when compared to a randomised distribution (p-value < 0.001; permutation test, Additional file 1: Figure S3C and D). In both tissues, nearly 70% of all H3K9ac-enriched regions located at promoters; this enrichment is more pronounced than for DHSs (approximately 40%), suggesting a presence of H3K9ac at promoters in the absence of DHSs. The number of intergenic H3K9ac-enriched regions, excluding promoters, was 6,511 in total; 3,115 in V2-IST, 6,213 in husk, and 2,668 shared between both tissues (Table 1B).
Figure 4. Average DNase I hypersensitivity and H3K9ac enrichment at genic regions. Average signal (in RPM) for DNase I hypersensitivity in (A) V2-IST and (B) husk, and for H3K9ac enrichment in (C) V2-IST and (D) husk at genes and their 1 kb flanking regions. Genes were binned based on their expression levels, from no expression (light colour) to high expression (dark colour): the lowest expression level bin contains all genes with an expression lower than 1 RPKM. The thresholds (in RPKM) are at 1.94, 4.17, 8.58, 16.64, and 36.28 for V2-IST and 1.88, 4.00, 8.34, 15.83, and 32.99 for husk tissue.

The overall H3K9ac-enriched regions occupy 2% and 7% of the uniquely mappable genome for V2-IST and husk, respectively (Figure 3I, K). The fraction in husk is larger than in V2-IST because there were 1.5-fold more H3K9ac-enriched regions in husk and these regions were also longer (Additional file 1: Figure S4A, medians of 603 bp and 1,015 bp in V2-IST and husk, respectively). The latter aspect is partly due to merging H3K9ac-enriched regions from three replicates for husk and two for V2-IST. Interestingly, despite the increase in H3K9ac-enriched regions in husk compared to V2-IST, no difference in the distribution of gene expression levels between the two tissues was observed (Additional file 1: Figure S4B). This observation
suggests that the number of active genes is similar between the two tissues and independent from the identified number of H3K9ac-enriched regions.

**46,935 intergenic regions with low DNA methylation are potential enhancer candidates**

Low DNA methylation was selected as third feature to identify enhancers because of its positive correlation with enhancer activity in mammals and plants [29,36,55–58]. To count the number of potential enhancers in the B73 maize genome, publicly available BS-seq data obtained from B73 coleoptile shoots were used [35]. Studies in Arabidopsis have revealed that DNA methylation levels in CG (mCG) and CHG (mCHG) contexts (H being A, C, or T) are highly stable in different vegetative tissues [59,60]. Furthermore, locus-specific [36] and genome-wide studies in maize ([61]; R.O. and M.S, and N.M.S. unpublished observations) provided little evidence for changes in mCG or mCHG levels in different vegetative tissues, justifying the use of the coleoptile shoot data set. We identified regions with 20% or lower DNA methylation in CG and CHG contexts independently, followed by defining LUMRs as regions that were low in both mCG and mCHG. Data on DNA methylation in CHH context (mCHH) was not included in the enhancer prediction step since, compared to the average levels of mCG and mCHG (86% and 74% respectively), mCHH levels are generally low in maize (2%), like in other plant species [35,62,63]. The distribution of LUMRs within the genome was investigated by counting their number in each genomic region (Figure 3L). The distribution of LUMRs in the uniquely mappable genome revealed an enrichment at genic regions, especially in exons, and at promoters (p-values < 0.001; permutation test for all genomic categories), but a scarcity at TEs (p-value = 1; permutation test for TEs); this observation is coherent with the fact that most TEs are highly methylated [35,64,65]. Investigation of the LUMR fractions revealed that nearly 50% of the genic regions are lowly methylated, which increases to nearly 60% for promoter regions and exons, while almost all TEs are highly methylated (Figure 3M). To identify potential intergenic enhancer candidates, we focused on intergenic LUMRs, excluding promoters. We identified 46,935 intergenic LUMRs as potential enhancer candidate regions.

**Integration of features for enhancer candidate prediction**

To predict enhancer candidates, we integrated the DHS, H3K9ac and LUMR data sets discussed above. Firstly, we calculated how many LUMRs and DHSs, or LUMRs and H3K9ac-enriched...
regions overlapped by at least 1 bp with each other. The overlap between the chromatin features was investigated in both tissues and revealed that more than 97% and 99% of the intergenic DHSs and H3K9ac-enriched regions, respectively, overlapped with LUMRs (Table 1). DHSs are generally shorter than LUMRs (Additional file 1: Figure S4A; median of 484 and 452 bp for V2-IST and husk, versus 834 bp, respectively). While most DHSs or H3K9ac-enriched regions co-localised within LUMRs, only about 20% of the total DHSs and H3K9ac overlapped with each other (Table 1).

Active enhancers are expected to be indicated by a coincidence of chromatin accessibility, H3K9ac enrichment and low DNA methylation [29,36]. We therefore filtered LUMRs based on the presence or absence of DHSs and H3K9ac-enriched regions, and defined LUMRs overlapping with both DHSs and H3K9ac-enriched regions as active enhancer candidates (Figure 2). Respectively, 398 and 1,320 candidates in V2-IST and in husk were identified, of which 223 were shared between the tissues, resulting in 1,495 enhancer candidates in total (Additional file 2: Dataset 1 and Additional file 3: Dataset 2). 256 V2-IST and 775 husk candidates were located more than 5 kb away, and 208 V2-IST and 623 husk candidates were located more than 10 kb away from their closest flanking genes. In V2-IST and husk tissue, the median distances between the candidates and their closest genes were 11.4 kb and 8.4 kb, while the largest distances were 438 kb (Zm00001d004626) and 498 kb (Zm00001d030489), respectively. Intersection of our candidates with a published data set of sequence comparisons between rice and maize genomes indicated that 41 (10%) V2-IST and 241 (18%) husk candidates contained conserved non-coding sequences (CNSs). The overlap between enhancer candidates and CNSs is higher than expected for randomized features ([66], p-value < 0.001; permutation test).

**Enhancer candidates and transposable elements**

Interestingly, 133 (33%) V2-IST and 370 (28%) husk candidates overlapped by at least 1 bp with TEs (Table 2). In most cases, enhancer candidates intersecting with TEs (TE-enhancer) overlapped more than 80% of their length or were entirely located within TEs. The number of TE-enhancers is the highest for Long Terminal Repeat (LTR) retrotransposons, followed by helitrons and Terminal Inverted Repeat (TIR) TEs, consistent with the fraction of the genome the three orders of TEs contribute to the TE space of the maize genome [39]. This TE space is calculated taking the average length for TEs and their number into account (136,000 LTRs with average length of 9,282 bp, 21,000 helitrons with an average length of 3,605 bp, and 14,000
TIRs with an average length of 621 bp). A small number of TIR elements (seven) are embedded entirely within enhancer candidates, possibly representing rare cases where the insertion of a small TE into open chromatin does not disrupt enhancer function. Indeed, these seven TIRs range from 83 to 199 bp; one overlaps with an H3K9ac peak, six do not overlap with either a DHS or H3K9ac peak; all are enriched in mCHH (Additional file 1: Figure S5A and B). To further assess the potential of TEs to create enhancers, for the remaining analyses we focused on the subset of TEs that contained at least 80% of an enhancer (Table 2).

The average distance between TEs and their closest genes did not vary between all TEs and TEs containing enhancer candidates (mean distance of 40.4 kb and 42.5 kb, respectively; Additional file 1: Figure S6A and B). The TEs that contain candidates tend to be longer than other TEs. To assess if enhancer candidates are likely to overlap with promoters that create functional transcripts for the TEs, we examined the distribution of the candidates within TEs. They were distributed randomly within the TEs, while functional TE promoters are expected to be located at the TE ends, indicating most candidates within TEs are unlikely to be located at the functional promoter site of TEs (Additional file 1: Figure S6C).

We explored the possibility that certain TE families could be a source of enhancers throughout the genome by looking for examples in which multiple members of the same TE family contained enhancer candidates (Additional file 4: Dataset 3). In most cases, only a single member of a TE family overlapped with enhancer candidates, with the exception of some very large TE families. Enrichment of TE families at enhancer candidates was tested by assuming a binomial distribution, and applying Bonferroni correction for multiple testing. Only three TE families showed significant enrichment for enhancer candidates (RLG00010, RLG00357, RLG01570; annotations are available from Gramene [67] and the TE classifications from the Maize transposable element (TE) database (http://maizetedb.org)). The LTR Gypsy family RLG00010 was most significantly enriched (p-val < 0.001), overlapping with seven V2-IST and 23 husk enhancer candidates. This represents a significant fraction of all TE-enhancers in the two tissues (7 and 8.6% for V2-IST and husk, respectively). The RLG00010 family was selected for further analysis.

The same trends were observed for RLG00010 members overlapping with enhancer candidates as for all TEs: a similar distribution of distances of TEs to their closest flanking gene (Additional file 1: Figure S6B and D), and a longer average length for TEs overlapping with candidates (10,895 bp compared to 8,517bp; Additional file 1: Figure S6A and E). Typical examples of RLG00010 TEs overlapping with enhancer candidates are shown in Additional file 1: Figure S5C. To examine if RLG00010 family members overlapping with enhancer
candidates were enriched for specific consensus sequences relative to other family members, several *de novo* motif analysis tools were used [68–71]. When comparing the results from different algorithms, the GGCCCA motif stood out as recurring (found by MEME with p-val < 0.0039, DREME with p-val < 0.043, RSAT Plants with E-value of 2.9e⁻⁷). This motif, also named site II motif, has been discovered in promoter regions of various genes that are highly expressed, for example ribosomal and DEAD-box RNA helicase genes [72–74]. TCP and ASR5 transcription factors are examples of proteins shown to bind the GGCCCA motif [75,76]. Scanning for the motif using FIMO [77] revealed that most enhancer candidates contained the GGCCCA motif irrespective of an overlap with the RLG00010 family (Additional file 1: Table S4). In fact, compared to random intergenic sequences, enhancer candidates showed an about 2-fold enrichment for the motif (p < 0.001). In contrast, the motif was not enriched in the RLG00010 family as such irrespective of their association with candidates.

### Table 2. Summary of overlap between enhancer candidates and TEs.

|                  | TE within enhancer candidate | Enhancer candidate entirely within TE | Enhancer candidate 80% within TE | Any Overlap |
|------------------|-------------------------------|---------------------------------------|----------------------------------|-------------|
|                  | LTR TIR Helitron               | LTR TIR Helitron                       | LTR TIR Helitron                 | LTR TIR Helitron |
| **V2-IST**       | 398                           | 0 1 0                                 | 83 1 10                          | 90 1 12 | 110 10 17 |
|                  | (0.2%)                        | (20.9%) (0.2%) (2.5%)                 | (22.6%) (0.2%) (3%)              | (27.6%) (2.5%) (4.3%) |
| **Husk**         | 1320                          | 0 7 1                                 | 202 9 56                         | 212 9 62 | 261 28 88 |
|                  | (0.5%) (0.1%)                 | (15.3%) (0.7%) (4.2%)               | (16.1%) (0.7%) (4.7%)           | (19.8%) (2.1%) (6.7%) |

Number of enhancer candidates overlapping fully, by 80% of their length, or least 1 bp with the TEs indicated. Percentages (in parenthesis) indicate the percent of intergenic enhancer candidates within each category. LTR stands for Long Terminal Repeats, TIR for Terminal Inverted Repeats. “All” indicates all enhancer candidates.

**Characterisation of enhancer candidates**

In humans, enhancers generally show a bi-directional pattern of DNA, chromatin, and transcript features. Histone modifications such as H3K27ac, as well as eRNA transcription, are located at both sides relative to single DHS peaks [4]. We set out to analyse whether DNA and chromatin features at our candidate enhancers displayed directionality. The read coverages for DNase-seq, H3K9ac ChIP-seq, and DNA methylation in all three contexts were extracted for each DHS located in enhancer candidates and their 1 kb up- and downstream flanking regions (431 candidates in V2-IST and 1,437 in husk) (Figure 5). Note that the number of DHSs was higher
than that of enhancer candidates because multiple DHSs could be located in one candidate. The averages of the read coverages are presented in Figure 6. Empirical observations indicated that H3K9ac was often enriched at only one side of DHSs (see e.g. Figure 7 and Additional file 1: Figure S7). Therefore, the orientation of DHSs was defined based on H3K9ac enrichment levels 300 bp from DHSs, the sides with the higher H3K9ac enrichment value, if present, being defined as 3’ end. The observed asymmetry was further validated by plotting the H3K9ac enrichment values from both sides of the DHSs with and without the previously defined orientations for all DHSs (Additional file 1: Figure S8). For DHSs showing H3K9ac enrichment at either side of at least 0.5 RPM, 241 and 841 out of respectively 431 in V2-IST and 1,437 in husk showed asymmetric H3K9a enrichment as indicated by an at least 2-fold change in H3K9ac enrichment between the two flanking regions.

The enhancer candidates were clustered into four categories based on H3K9ac enrichment patterns using the k-means clustering algorithm and the categories were numbered according to their appearance in the heatmaps (Figure 5). For each category, average patterns were

Figure 5. Heatmaps of chromatin, DNA and transcript features at enhancer candidates. DNase I hypersensitivity, H3K9ac enrichment, mCG, mCHG, and mCHH levels, presence of TEs, and transcript levels at and around (+/- 1 kb) DHSs in enhancer candidates. DHSs were scaled to equal size. The colour scales are in RPM for DNase I hypersensitivity, H3K9ac enrichment and transcript levels, and in methylation frequency (0-1) for DNA methylation. For TE sequences, red and white show the presence or absence of TEs, respectively. DHSs were clustered based on H3K9ac enrichment using a k-means (k=4) clustering algorithm. The categories identified were numbered from 1 to 4 from the top to the bottom. All the DHSs were oriented based on H3K9ac enrichment intensity values 300 bp away from the DHS boundaries; the side with higher H3K9ac enrichment was defined as 3’ end.
determined (Additional file 1: Figure S9). Heatmaps and profiles showed that H3K9ac can be primarily enriched on one side of the DHSs (category 1 and 2), within DHSs (category 3), or present at both sides but clearly enriched at one of them (category 4) (Figure 5 and Additional file 1: Figure S9). Comparing DNase-seq or H3K9ac ChIP-seq read coverages with the distribution of mCG and mCHG levels, but also the average profiles, indicated that high chromatin accessibility and H3K9ac enrichment levels were exclusive with high DNA methylation levels (Figure 5, 6 and Additional file 1: Figure S9). The average profiles show a plateau and steep decline of mCG and mCHG at the 5’ side of DHSs (Figure 6). In categories 1, 2 and 4, at the 3’ side of enhancer candidates, mCG and mCHG levels increased gradually (Figure 6, Additional file 1: Figure S9). These patterns indicate a sharp transition in DNA methylation level at the 5’ DHS boundaries, and a more gradual transition at the H3K9ac boundaries. However, a sharp transition at the 5’ ends of candidates may be masked in the average profile by the varying size of the H3K9ac-enriched regions. In line with this, the profile of category 3 candidates, having H3K9ac at the DHSs itself, showed sharp boundaries at both sides of the candidates. Levels of mCHH were lower than mCG and mCHG levels, as expected [35]. In line with earlier studies [61,62], mCHH marked boundaries between lowly and highly DNA methylated regions as shown by the relatively high level of mCHH, represented by a small mCHH peak in the average profiles, at the 5’ boundaries of the DHSs (Figure 5, 6 and Additional file 1: Figure S9). Additional heatmaps and profiles were created to illustrate the locations of TEs and transcripts for the four categories. The heatmaps suggest that TEs covered all selected regions, showing a slight depletion across DHSs but no apparent pattern across other features (Figure 5). In animal models, enhancers are characterised by bi-directional transcription and the transcribed regions are, among others, enriched with H3K27ac [4]. In our data, transcript levels were generally low at candidates except for a few showing transcripts within and/or outside of their DHS (Figure 5), making the detection of bi-directional transcription very challenging. In addition to this absence of detectable levels of bi-directional transcription, the clear asymmetric H3K9ac distribution at a majority of maize enhancer candidates suggested that the candidates have more resemblance to TSSs than animal enhancers do [4].
Figure 6. Average profiles of the enhancer candidates in (A) V2-IST and (B) husk. Average signal intensities of DNase I hypersensitivity, H3K9ac enrichment in RPM and DNA methylation levels in methylation frequency at DHSs and their 1 kb flanking regions. DHSs were scaled to equal size. Prior to calculation of the average, all the DHSs were oriented based on H3K9ac enrichment intensity values 300 bp away from the DHS boundaries; the sides with higher H3K9ac enrichment were defined as 3' end. The profiles show a clear preferential enrichment of H3K9ac 3' of the DHSs and high levels of DNA methylation (CG and CHG context) around the DHSs and H3K9ac-enriched regions. The level of mCHH is low throughout the regions with a slight increase at the 5' side of DHSs.

Profiles of DNA and chromatin features at enhancer candidates and TSSs are similar

To rule out the possibility that our enhancer candidates were actually TSSs of unannotated genes, we compared the patterns of their DNA, chromatin features, and transcript features with those observed at annotated TSSs by randomly selecting 431 and 1,437 DHSs located at TSSs
for V2-IST and husk, respectively (Additional file 1: Figure S10). The selected regions were oriented according to the 5’ to 3’ orientation of flanking genes and analysed using the k-means clustering algorithm (k=3). In general, the heatmaps and average profiles of DHSs at TSSs displayed a strong DNA methylation signal at the 5’ ends of DHSs, and an enrichment in H3K9ac and an accumulation of transcripts at the 3’ ends of DHSs (Additional file 1: Figure S10 and S11). The heatmaps and the average plots of TSSs and enhancer candidates revealed similar patterns of chromatin accessibility and H3K9ac, but they differed in transcript levels (higher at annotated TSSs) and distribution of mCG and mCHG (high on both sides for candidates, while restricted to the 5’ side for annotated TSSs) (Figure 5, 6, Additional file 1: Figures S10 and S11). The median transcript level at the enhancer candidates was 6.6 times lower than that at coding sequences in V2-IST; the fold change could not be calculated for husk because the candidate expression levels had a median of zero RPKM (Additional file 1: Figure S12). One category (category 3), showed transcriptional activity and H3K9ac enrichment on both sides (Additional file 1: Figure S10). The DHSs in this category were either flanked by two oppositely orientated and closely spaced genes, or by alternative TSSs located in upstream regions. H3K4me3 histone modification was previously described for distinguishing TSSs from enhancers [21,78–80]. Analysis of published ChIP-seq data for H3K4me3 in maize third seedling leaf [61] indicated that 24% and 11% of the V2-IST and husk enhancer candidates, respectively, overlapped with H3K4me3 enriched regions (Additional file 1: Figure S13), which could hint at unannotated TSSs. The observed H3K4me3 enrichment at enhancer candidates was, however, on average weaker than at TSSs (Additional file 1: Figure S13), suggesting H3K4me3 may also differentiate TSSs and enhancers in maize. In addition, the H3K4me3 enrichment pattern did not entirely reflect the H3K9ac enrichment pattern at TSSs
Figure 7. Example of data on (A) DICE and (B) b1 repeat enhancer. From the top: AGPv4 annotation and candidate annotation from our prediction (V stands for V2-IST, H for husk candidate), DNase I hypersensitivity and H3K9ac enrichment signal (all replicates pooled) and peak position (indicated as blue and green bars, respectively) in V2-IST and in husk tissue, mCG, mCHG and mCHH levels and unique mappability in percentage. The numbers under gene names indicate relative gene expression levels (V2-IST/husk). Although the b1 locus is on chromosome 2; in the current version of the AGPv4 assembly, the b1 gene is located in contig 44 (B, on the right of the grey vertical line). The dark blue bars in the gene annotation tracks indicate previously annotated known enhancers and putative cis-regulatory elements. The vertical red boxes indicate enhancer candidates identified in this study. Peaks at those tracks might not be present in each replicate, affecting enhancer candidate prediction.
but was rather slightly shifted downstream of the H3K9ac peaks. Such a pattern has not been reported in humans [79] and was not observed in a previous study in rice [21].

In summary, despite a shared polarity with respect to flanking H3K9ac enrichment, the profiles of enhancer candidates differ from those at TSSs by the levels of transcript accumulation, DNA methylation and H3K4me3.

### Ranking and selecting a list of tissue-specific enhancer candidates

To facilitate linking enhancer candidates to putative target genes, we set out to determine the degree of tissue-specificity of our enhancer candidates by ranking the 398 V2-IST and 1,320 husk candidates based on the assumption that the levels of both DNase I hypersensitivity and H3K9ac enrichment are positively correlated with enhancer activity. The enhancer candidates were independently ranked based on the largest differences between the two tissues for DNase I hypersensitivity and H3K9ac levels. The strongest tissue-specific candidates were assumed to exhibit large differences in both DNase I hypersensitivity and H3K9ac enrichment, therefore the independent rankings for both features were summed for every candidate and the candidates were re-ranked (Additional file 2: Dataset 1 and Additional file 3: Dataset 2, column overall_rank). The ranking numbers were combined with a V for V2-IST or an H for husk as candidate IDs; the lower the number, the more tissue-specific the candidate. However, the rankings for DNase I hypersensitivity and H3K9ac enrichment did not correlate with each other (Additional file 2: Dataset 1 and Additional file 3: Dataset 2, column DNase_rank and H3K9ac_rank; shared candidates were ranked in both tissues). For example, the candidate ranked to the second place (candidate V2, Figure 8) for V2-IST showed a large difference in DNase I hypersensitivity signal between V2-IST and husk tissue as expected, while the H3K9ac enrichment stayed almost the same for both tissues. The 313th candidate in V2-IST (candidate V313), on the other hand, is characterized by a large difference in H3K9ac enrichment but not in DNase I hypersensitivity. The 194th candidate in V2-IST (candidate V194) showed a large difference between the tissues for both DNase I and H3K9ac enrichment signals but in an opposite direction. The lack of correlation between the ranks obtained from both chromatin features indicated that determining tissue-specificity using this combination of features does not work properly. Experimental examinations of a number of candidates will be necessary to determine the best feature (combination) to predict tissue-specificity. For now, enhancer candidates identified in only one of the two tissues were defined as tissue-specific, and the shared candidates between tissues as putative shared enhancers. With this definition, a total of
1,495 candidates were classified into 175 V2-IST-specific, 1,097 husk-specific and 223 shared candidates (Additional file 5: Dataset 4).

![Figure 8. Examples of candidate rankings. From the top: Identified candidate region with its ID (V stands for V2-IST, H for husk candidate) and coordinates, DNase I hypersensitivity and H3K9ac enrichment signal intensities in V2-IST and husk tissues. In these examples, the DNase I hypersensitivity and H3K9ac enrichment signal differences do not positively correlate to each other as assumed.]

**Predicting putative target genes of enhancer candidates based on expression levels of closest genes**

Lastly, we examined if our candidates could be linked to putative target genes. Multiple approaches have been reported using data on chromatin accessibility, transcript levels and/or histone modification patterns at both enhancers and genes, across different tissues or developmental time points [4,51,81,82]. We assumed that enhancers regulate the expression of either their adjacent up- or downstream gene, though it has been observed that other genes can
be located between enhancers and their target genes in animals and plants [17,83–85]. We correlated the defined tissue-specificity of candidate enhancers with the gene expression levels of the nearest flanking genes in both tissues. Only genes showing significant differential expression between V2-IST and husk tissue (Cuffdiff [86]) were considered as targets of tissue-specific enhancer candidates; for shared candidates, flanking genes that are expressed in both tissues were considered as potential target genes. If a flanking gene showed a significant difference in gene expression that matched the enhancer candidate specificity (e.g. higher gene expression in V2-IST for V2-IST candidates), then the candidate and the gene(s) were linked. With this method, 38 (22%) V2-IST-specific, 143 (13%) husk-specific and 101 (45%) shared enhancer candidates were linked to one putative target gene (Additional file 5: Dataset 4). We also identified 13 (2%) V2-IST-specific, 182 (17%) husk-specific and 103 (46%) shared candidates in which both flanking genes showed expression levels matching the features of the candidates. The other candidates could not be linked to a gene because either none of the flanking genes had a significant expression level difference in the expected direction for tissue-specific candidates (124 (71%) in V2-IST, 772 (70%) in husk) or, in case of shared enhancer candidates, neither of the flanking genes were expressed in one of the tissues (19 (9%) candidates).

**Identification of three known enhancers in maize**

In maize, five well-characterised and putative enhancers were reported, namely the \( b1 \) hepta-repeat, the enhancers of \( tb1 \), \( p1 \), and the putative enhancers DICE and \( Vgt1 \) that regulate the expression of the genes \( bx1 \) and \( ZmRAP2.7 \), respectively [11,13–15,23,85,87]. In our screen, we identified the confirmed and putative enhancers of \( b1 \), \( tb1 \) and \( bx1 \) (Figure 7 and Additional file 1: Figure S7), although these enhancers were mostly identified and characterized in maize lines other than B73, which could have affected their functionality. For example, the \( b1 \) hepta-repeat enhancer has been identified for the \( B-I \) epiallele and consists of seven copies of a 853 bp sequence in tandem, while B73 only carries a single copy of this sequence (90% identity with consensus repeat sequence; [12]). In our data set, \( b1 \) showed differential expression in the same direction as observed in the line the \( b1 \) repeat enhancer was discovered [23], already indicating there is some degree of conservation in the regulatory region. The \( tb1 \) enhancer was identified in the inbred line W22 [13,14], and DICE was shown to be required for high \( bx1 \) expression in Mo17 [85]. The enhancer candidates for \( b1 \) and DICE were not linked with \( b1 \) and \( bx1 \), respectively, because their known target genes were not the closest flanking gene. We identified neither the \( p1 \) enhancer nor \( Vgt1 \). In the case of the \( p1 \) locus, high repetitiveness of
Chapter 2

the region rendered the enhancer unmappable. For Vgt1, a clear DHS was present but H3K9ac-enrichment was not detected within the overlapping LUMR.

Four H3K9ac-enriched enhancer candidate regions identified by ChIP-seq, candidate H108, the b1 and tb1 enhancer and DICE, were selected for validation with ChIP-qPCR. For each region, primer pairs were designed to amplify sequences located at the summit of the peak of the ChIP-seq H3K9ac-enriched region (P), its slope (S), and outside of the peak (O; no enrichment by ChIP-seq) (Additional file 1: Figure S14). Results confirmed the presence and absence of H3K9ac enrichment at the identified candidate regions and their flanking regions, respectively. The differential H3K9ac enrichment observed for candidate H108 and the b1 enhancer fits their expected husk tissue-specificity based on the ranking. DICE had a high and low ranking in V2-IST and husk, respectively. In accordance, DICE showed higher H3K9ac enrichment levels in V2-IST than in husk. The tb1 enhancer showed H3K9ac enrichment in both V2-IST and husk. This is in accordance with what is observed for the pooled ChIP-seq data (Additional file 1: Figure S14C). Due to our stringent criteria, the tb1 enhancer was only called as a candidate in husk.

To examine if H3K4me1 is indeed not enriched at enhancers as suggested by the results depicted in Figure 1, enrichment for H3K4me1 was determined for the same regions as for H3K9ac enrichment (Additional file 1: Figure S14). Except for the enhancer of tb1, none of the analysed regions showed a clear H3K4me1 enrichment, confirming our previous observation and supporting the idea that H3K4me1 does not generally mark plant enhancers.

Discussion

The combination of DNase-seq, H3K9ac ChIP-seq and BS-seq data allowed us to identify approximately 400 and 1,300 enhancer candidates in V2-IST and husk tissue, respectively, and about 1,500 unique enhancer candidates in total. Interestingly, our enhancer candidates displayed an asymmetric enrichment of H3K9ac at DHSs, which differs from the histone acetylation enrichment at both sides of DHSs observed in animals [4,27,29]. Target genes were predicted for 255 V2-IST and 529 husk candidates. Importantly, our method successfully predicted three confirmed or putative enhancers in the maize genome, enhancers for the b1 (candidate H167) and tb1 (candidate H1233) genes, and the DICE enhancer (candidates V4 and H1318).

We investigated the enrichment of three histone modifications at the enhancer of b1: H3K27ac, H3K9ac, and H3K4me1, and showed that both H3K27ac and H3K9ac were enriched at the
hepta-repeat enhancer of \( bl \) in the active, but not the inactive state. These results are in accordance with previous studies in animals, but also in plants [20,28,30,36,37,88]. In contrast, H3K4me1, which was shown to be enriched at animal enhancers regardless of their activity [27,79], was not enriched at the \( bl \) hepta-repeat enhancer, but also not at DICE and candidate H108 (Additional file 1: Figure S14), while it was present at relatively high levels at transcribed regions of \( bl \) and \( actin1 \) (Figure 1). This distribution at enhancers may be typical for plants as it is supported by previous observations in Arabidopsis in which H3K4me1 was almost exclusively enriched in genic regions [89].

Regions with low DNA methylation overlap with DHSs and both were previously used to predict enhancers [29,90]. In our study, more than 97% of DHSs and more than 99% of H3K9ac-enriched regions overlapped with LUMRs, and enhancer candidates were identified by taking the overlap between LUMRs, DHSs and H3K9ac-enriched regions, resulting in about 1,500 unique enhancer candidates. Many more intergenic LUMRs were identified (about 47,000), and 26% of these carried only one, while 71% carried none of the other required chromatin features. We hypothesize that these remaining LUMRs represent enhancers in tissues other than the ones used in our study. This could in part explain the relatively low number of identified candidates compared to studies in animals in which a large number of developmental stages, cell types, and/or tissues were used [4,51,83]. In Arabidopsis more than 10,000 intergenic enhancer candidates were predicted using only two different tissues [20], as we did. However, the authors based their prediction solely on chromatin accessibility. Based on chromatin accessibility data only, we would predict about 9,000 candidate enhancers. Instead, we used a more stringent approach to identify active enhancers.

10% and 18% of V2-IST and husk candidates contained previously published CNSs between maize and rice [66], suggesting these candidate sequences and functions may be conserved across species. The rest of the candidates might be maize specific or rapidly diverging [91], explaining the lack of sequence conservation. About 30% of the enhancer candidates in both tissues overlapped by at least 1 bp with TEs (33% in V2-IST and 28% in husk), and in most cases TEs covered the entire enhancer candidate region. This raises questions regarding the origin of the regulatory potential of those enhancer candidates. Indeed, TEs have been reported as an important source of \emph{cis}-regulatory elements because TEs have evolved to mimic the regulatory sequences of the host to hijack its transcriptional machinery [14,38,92–94]. Three LTR Gypsy families were significantly enriched for enhancer candidates. Motif analysis of the enhancer candidates overlapping with the most enriched TE family, RLG00010, identified the GGCCCA motif, which is discovered in \emph{cis}-regulatory elements of genes with diverse functions...
Compared to random intergenic sequences, this motif was not only enriched in the RLG00010 enhancer candidates, but also in all other candidates. This suggests that GGCCCA is a general motif associated with enhancer function.

Although we identified three previously discovered putative or confirmed enhancers in maize, two others, Vgt1 and the enhancer of p1, were not detected. This can be explained by several factors: i) enhancer sequences can be located in repetitive regions, which are not uniquely mappable and therefore excluded from our analysis (true for the p1 enhancer), ii) enhancers may not always require the stringent criteria used to define enhancer candidates in this study (could be true for Vgt1, which featured an LUMR and DHS but no H3K9ac-enriched region), iii) enhancers may not be active in V2-IST or husk tissue and therefore undetected, and iv) enhancers may only be present in other lines than B73.

We identified about three times more enhancer candidates in husk tissue than in V2-IST (398 vs 1320), which is possibly due to a larger number of H3K9ac-enriched sequences in all genomic regions in husk compared to V2-IST (Figure 3H and J). There was however no difference in the distributions of gene expression levels between the two tissues (Additional file 1: Figure S4B), indicating that the number of genes expressed at particular levels is similar in V2-IST and husk and that the larger number of H3K9ac-enriched sequences is therefore not due to a higher number of genes being expressed in husk. The differences in the number of H3K9ac-enriched regions were substantial, even when considering possible technical bias introduced during the analysis. This observation highlights that the H3K9ac enrichment pattern changes between tissues and/or developmental stages, irrespective of the overall distribution of expression levels. The reasons for this change are currently unknown.

The heatmaps and average profiles of the chromatin and DNA features at the candidates revealed that H3K9ac was preferentially enriched on one side of the DHSs (Figure 5 and 6). This observation was unexpected considering earlier studies in animals describing histone acetylation (H3K27ac), but also methylation (e.g. H3K4me1) enrichment on both sides of DHSs at enhancers [4,27,29]. Symmetrical enrichment of histone modifications at animal enhancers has been associated with bi-directional transcription at enhancers [4]. Given the relative low coverage of our RNA-seq data at enhancer candidates, we were not able to assess whether eRNAs were produced bi- or uni-directionally. eRNAs are indeed known to be transcribed at a low level and in addition sensitive to degradation, making them difficult to detect with a technique such as RNA-seq [4,95]. The analysis of nascent transcript data (GRO-seq) for maize and Arabidopsis suggests the absence of transcription at plant enhancers [96], ...
further supporting the possible differences between plant and animal enhancers. A method like CAGE-seq could be used to further investigate the transcription of enhancers in plants.

Elevated levels of mCHH were detected 5' of the DHSs at enhancer candidates. mCHH islands have been observed to flank genic regions in maize, but also low DNA methylated intergenic CNSs [61,65]. The findings of Li et al. [61] showed that mCHH islands may act as boundaries between euchromatin and heterochromatin, preventing activation of TEs by nearby transcriptionally active genes. A similar function is likely at enhancers.

Comparison between the chromatin and DNA methylation profiles at enhancer candidates and TSSs revealed the presence of similar features, including chromatin accessibility, asymmetric H3K9ac enrichment and low DNA methylation. On average, the TSSs show a higher level of transcript accumulation, a lower level of DNA methylation 3’ of TSSs and a higher level of H3K4me3 than enhancer candidates (Figure 5, S9, S11 and S13). The difference in transcript levels and H3K4me3 enrichment between enhancers and TSSs has been observed by others [4,79].

For each enhancer candidate, a target gene was predicted following expression and proximity criteria. Our prediction method assumed that target genes were either the adjacent up- or downstream gene, and that target genes of tissue-specific enhancer candidates would be upregulated in the tissue in which the enhancer candidates were detected. Using our stringent criteria, 580 candidates were linked to genes, including tb1. In Drosophila, about 20% of the enhancers were predicted to control genes that were not directly adjacent to the enhancers [83] and a recent prediction in human and mice estimated that 69% of the enhancers contact genes that are not directly consecutive [82]. Whether this proportion is similar in maize remains to be determined, but examples of such enhancers have been reported, for example DICE, the putative enhancer of bx1 [85]. In addition, our approach disregarded the possibility that enhancer candidates would act as transcriptional repressors [97]. Future studies in maize are required to more precisely identify and validate the target genes of the enhancer candidates discovered.

**Conclusions**

This study provides a genome-wide glance at transcriptional enhancer candidates in maize by comparing DNA and chromatin features in two maize tissues and by providing details on some of their characteristics. The study identified about 1,500 enhancer candidates that were characterized by increased chromatin accessibility, low DNA methylation levels, and
asymmetric enrichment of H3K9ac. Three identified candidates were putative or confirmed enhancers ($b1$, $tb1$ and $bx1$ enhancers). In contrast to animals, plant enhancer candidates show asymmetric chromatin features. Validation of enhancer candidates remains to be achieved. Future improvements in predicting enhancer candidates are expected from the investigation of more histone modifications as well as TF binding sites, the integration of genome-wide chromosomal interaction data and a direct functional analysis of candidates, e.g. by targeted genome editing. A better understanding of the regulatory code in maize not only helps to better compare transcription regulation in highly complex genomes of different kingdoms but promises new targets for informed breeding in this important crop. Our data provides a framework for the maize community to characterize the regulation of genes of interest.

**Material and Methods**

**Experimental methods**

**Plant stocks and material**

The seed stock of the maize B73 inbred line used in this study was obtained from J. Gardiner (University of Arizona, Tucson, AZ) in 2013. It was obtained from the North Central Regional Plant Introduction Station in Ames Iowa, USA (Order No: 169545, Accession: PI550473, Lot: 94ncai02). It is from the same Accession (PI 550473) that was used for the maize B73 genome sequencing project [41], but a different lot number because it was requested several years later. The $B$-$I$ plant stock used in this study (W23) was obtained from V.L. Chandler (University of Arizona, Tucson, AZ). Maize plants were grown in the greenhouse at two different locations: The Max Planck Institute for Plant Breeding Research in Cologne (MPIPZ) and the University of Amsterdam (UvA). At the MPIPZ maize plants were grown for DNase-seq and RNA-seq. At the UvA maize plants were grown for H3K9ac ChIP-seq and RNA-seq. At both locations plants were grown in soil under 16-h light/8-h dark cycles at an average temperature of 23°C. The plants were harvested at the V2 stage (2 collars visible; V2-IST), V5 stage (5 collars visible; V5-IST) or when the silks started emerging from the husks. The two tissues used for the RNA-seq, DNase-seq, and ChIP-seq experiments were the inner stem tissue of V2 seedlings, which is composed of the seedling stem with the outer leaves and all exposed leaf blades removed, and the soft inner husk leaves surrounding the ear; the tough outer husk leaves were discarded (Additional file 1: Figure S1).
RNA-seq
RNA for RNA-seq experiments was isolated at both locations. To be able to examine reproducibility and comparability, per tissue, three biological replicates were analysed, each consisting of pooled material from three plants. The inner husk leaves and inner stem tissue of V2 seedlings were flash frozen in liquid nitrogen between 9 and 11 hours after dawn. After grinding in liquid N₂, 100 mg material was used for RNA extraction with TRIzol (ThermoScientific) following manufacturer's instructions except that the top aqueous phase was transferred to a new tube, 500 µl of isopropanol were added, followed by mixing and incubation for 10 min at RT. The entire sample was transferred in two steps to an RNeasy MINI spin column (Qiagen RNeasy kit) and centrifuged for 15 sec at 8000 x g. The flow-through was discarded and 700 µl of the Qiagen RW1 buffer was added. Two washing steps were performed using 500 µl of the Qiagen RPE buffer. RNA was eluted in 50 µl RNase-free water and the concentration was assessed spectrophotometrically (Nanodrop, ThermoScientific). Next, RNA samples were diluted to a concentration of 200 ng/µl and treated with DNase I (DNA-free kit, Ambion) according to manufacturer’s instructions. Samples were then extracted with 1 volume of phenol:chloroform:isoamyl alcohol (25:24:1 v/v) and centrifuged for 5 min at 13,000 x g, 4 °C. The same step was repeated twice. Next, 80% of the aqueous phase volume was transferred into a new tube and precipitated with 1/10th volume of 3 M Sodium Acetate pH 5.6, two volumes of 100% Ethanol, and 1 µl of glycogen (10 mg/mL), followed by centrifugation at 13,000 x g for 15 min at 4 °C. The pellet was subsequently washed twice with 70% ethanol and finally resuspended in 20µl of RNase-free water. The concentration was measured spectrophotometrically (Nanodrop, ThermoScientific) and 1µg of RNA was separated on a 1.2% agarose 1x MOPS (3-N-morpholinol propane sulfonic acid) gel to assess RNA quality. The concentration was adjusted to 400 ng/µl and 500 ng of total RNA was treated with the Ribo-Zero rRNA Removal Kit (Plant Leaf, Epicentre) to specifically remove ribosomal RNAs. RNA-seq libraries were prepared with the NEBNext Ultra™ Directional RNA Library Prep Kit for Illumina sequencing (New England Biolabs). Quality and quantity were assessed at all steps of the library preparation by capillary electrophoresis (Agilent Bioanalyser and Agilent Tapestation). Sequencing was performed with TruSeq v3 chemistry on a HiSeq2500. Approximately 15-20 million of 100 bp single-end reads were obtained for each library.
DNase-seq

Nuclei preparation. For each inner stem tissue sample (V2 stage) and inner husk leaf sample nuclei were extracted from 12 V2 stage maize seedlings and three husks according to the protocol of Steinmüller and Appel [98]. For each tissue, two biological replicate samples were used. Briefly, tissue was ground in liquid nitrogen, 5 g were transferred into an ice-cold 50 ml centrifuge tube, and 25 ml of cold nuclei isolation buffer (20 mM Tris-HCl pH8, 250 mM sucrose, 5 mM MgCl2, 5 mM KCl, 40% glycerol, 0.25% Triton X-100, 0.5 mM EGTA pH 8, 5 mM EDTA pH8, 0.1 mM PMSF, 0.1% 2-mercaptoethanol, 1:1000 Proteinase Inhibitor Cocktail (Sigma)) were added and the tube was flicked until the powder was in suspension. The tube was rotated at low speed at 4°C until the sample was completely thawed (about 30 min). The tissue suspension was filtered through successive layers of 60 µm and 20 µm nylon mesh (Nylon Net Filters, Millipore) into an ice-cold 50 ml centrifugation tube and centrifuged at 6,000 x g for 15 min at 4 °C. The supernatant was discarded and the pellet resuspended in 15 ml of ice-cold nuclei isolation buffer using a 1 ml cut-off pipette tip, followed by centrifugation at 6,000 x g for 12 min at 4 °C. The pellet was resuspended in 10 ml of ice-cold nuclei isolation buffer and centrifuged at the same conditions again, followed by resuspending the pellet in 1 ml of ice-cold nuclei storage buffer (20% glycerol, 20 mM Tris pH 7.5, 5 mM MgCl2, 1 mM DTT). To check the quality and abundance of the nuclei, a 20 µl aliquot was stained with 1 µl DAPI (1 mg/ml) and examined by fluorescent microscopy. The nuclei suspensions were flash frozen in liquid nitrogen and stored at -80 °C until further use.

DNase I digestion. DNase I treatment was adapted from Chandler et al. [99]. Nuclei suspensions were thawed on ice while preparing the solutions for DNase I digestion. One undigested control and 4 concentrations of DNase I (50, 100, 150, and 200 U/ml) were used (Additional file 1: Figure S15). 2.5 ml of DNase I buffer (50 mM Tris pH8, 250 mM sucrose, 100 mM KCl, 0.1 mM CaCl2, 5 mM MgCl2, 50 µg/ml BSA, 0.05 M beta mercaptoethanol) was prepared per sample. The DNase I dilutions were prepared by mixing DNase I (Roche) with DNase I dilution buffer (20 mM Tris pH7.5, 50 mM NaCl, 1 mM DTT, 100 µg/ml BSA, 50% glycerol). 1 ml of nuclei suspension was divided in 5 x 200 µl in 1.5 ml microcentrifuge tubes using cut-off pipette tips. The tubes were centrifuged at 1,500 x g for 5 min at 4°C and the supernatant was discarded. 100 µl of 100 mM EDTA pH 8, followed by 600 µl of phenol/chloroform/isoamylalcohol (25:24:1 v/v) were added to the tube for the undigested control and set aside at RT after thorough mixing. The other pellets were resuspended in 475 µl of cold DNase I buffer by rubbing the tubes against a plastic tube rack and letting incubate 3 min at 25°C. 25 µl of each of the DNase I dilutions were added to the respective tubes with
nuclei suspensions and incubated for 10 min at 25 °C. The reaction was stopped by adding 100 µl of 100 mM EDTA pH 8, mixing, and adding 600 µl of phenol/chloroform/isoamylalcohol. All samples, including the undigested control, were shaken by hand or using a tissue lyser (Qiagen) at 8 Hz for 5 min. A second phenol/chloroform/isoamylalcohol extraction was performed, followed by an RNase A treatment (2 µg/ml final concentration) at 37°C for 10 min. 600 µl isopropanol, 50 µl of 7.5 M ammonium acetate and 2 µl of 10 mg/ml glycogen were added followed by centrifugation at 16,000 x g for 30 min at 4°C. Two 70% ethanol washings were performed and the pellets were finally resuspended in 30 µl 10 mM Tris-HCl pH 8.5. The concentration of nuclei acids was then assessed spectrophotometrically (Nanodrop, ThermoScientific) and the entire sample (30 µl) was mixed with 6 µl Cresol Red loading buffer (1.75M sucrose (60%), 5mM cresol red, pH 8) and loaded on an agarose gel (1x TAE buffer, 1.5% agarose, 0.5 µg/ml ethidium bromide). Gel visualization under UV light indicated which digestion fulfilled the requirement that the DNA is only partially digested (Additional file 1: Figure S15). In our hands, these were the samples digested with 50 U/ml of DNase I. One should test several concentrations as the digestion efficiency can vary depending on the batch of DNase I enzyme and chromatin concentration. The DNA fractions between 100-300 bp were extracted from the gel using gel purification (NucleoSpin Gel, Macherey Nagel) and the DNA was eluted from the column in 15 µl 10 mM Tris-HCl pH 8.5. The DNA concentration was measured using QuantiT PicoGreen (Invitrogen) on a fluorometer (Synergy 4 Hybrid Multi-Mode Microplate Reader, BioTek). A DNA concentration range of 1-3 ng/µl was obtained.

**Naked DNA control.** Genomic DNA was extracted from 100 mg of inner husk tissue derived from three pooled husks using the DNeasy Plant Mini kit (Qiagen) and following manufacturer’s instructions. 1.7 µg of gDNA was digested with 50 U/ml of DNase I following the same protocol as described for chromatin.

**Library preparation and sequencing.** DNA samples were diluted to 1 ng/µl in a total volume of 10 µl followed by library preparation using the Ovation Ultralow DR Multiplex kit (NuGEN) according to the manufacturer’s protocol. 15 cycles of amplification were performed for the naked DNA sample and 16-18 cycles for the chromatin-derived samples. The libraries were sequenced on an Illumina Hi-Seq2500 platform and approximately 20-30 million 100 bp single-end reads were obtained for each library.

**ChIP-seq and ChIP-qPCR**

The ChIP procedure was based on the original protocol from Haring *et al.* [100] with minor modifications. In short, plant samples (5 inner stems from V2 plants or 3 gr of inner husk leaves
per sample) were fixed with formaldehyde. Chromatin was extracted and sonicated. The soluble fraction was then immunoprecipitated using antibodies against H3K9ac (Abcam, ab10812), H3K27ac (Abcam, ab4729), H3K4me1 (Abcam, ab8895) or rabbit serum (No antibody control, Sigma no. R9133) using protein-A coated magnetic beads (ChIP-seq, Diagenode, kch-802) or protein-A agarose beads (ChIP-qPCR, Sigma-Aldrich). Immunoprecipitated DNA was recovered, decrosslinked and column-purified (Qiagen, 28104). For each ChIP-seq library, three ChIP samples were pooled yielding about 50 ng of DNA prior to adapter ligation and PCR amplification. Adaptor ligation (TrueSeq Universal adapter, Illumina) and PCR amplification were performed for each pooled ChIP sample using the KAPA Hyperprep kit (KAPA, KK8500) as indicated by the manufacturer. The efficiency of the conversion process was assessed by comparing the input ChIP sample to the obtained ChIP-seq library on an Agilent High Sensitivity D1000 ScreenTape System. Efficient conversion corresponds to a visible 100 bp shift in fragment sizes and an unbiased increase in DNA concentration. For all samples, approximately 30 million 100 bp single-end reads were generated on an Illumina HiSeq2500 platform.

For ChIP-qPCR, the column-purified material (4 µl out of 80 µl) was mixed with 2 µl of each primer (10 µM; Additional file 5) and 4 µl of the 5X FIREPol Evagreen qPCR Mix plus (Solis Biodyne) in a total volume of 20 µl and run on an Applied Biosystem 7500 Real Time PCR system (50°C, 2'; 95°C, 10', 45 cycles: 95°C, 15''; 65°C, 1'). For each primer pair, a calibration curve was generated using DNA isolated from fixed, sonicated chromatin (100 ng/µl; dilutions 1/64, 1/256 and 1/1024) to test primer efficiency and calculate DNA quantities from ChIP samples. Enrichment is calculated as the mean quantity of the different biological replicates (2-5) and normalized over the quantity at the maize actin locus. All PCR primer sequences are listed in Additional file 6: Table S5.

**Computational analysis**

For all the analysis, the B73 maize genome sequence and annotation version 4 (AGPv4) [39] from Ensembl Plants [40] were used as the reference. Data on chromosomes 1 to 10, excluding contigs, were used for all the analysis. For statistical enrichment analysis, permutation tests were performed (n = 1000) [101]; the randomisation of features within the uniquely mappable part of genome was performed using BEDtools [102].
RNA-seq

The sequenced reads were trimmed at both ends based on sequencing quality (Q20) and remaining Illumina adaptor sequences were removed using Trimmomatic [103]. When the remaining read length was less than 35 bps, the read was removed from the analysis. The reads were aligned, allowing 1 mismatch, to the reference genome using TopHat2 [104] and Bowtie [105]. Transcript assembly and gene expression level calculation for each replicate were performed with a guided reference [40] using the Cufflinks pipeline (Cufflink, Cuffquant and Cuffnorm; [106]). The RPKM values and the significance of the differential expression levels for each gene were calculated taking the variance over the six replicates using Cuffdiff [86]. The RPM coverage in the genome was calculated using BEDtools [102].

DNase-seq and ChIP-seq

For DNA-seq data, to assess technical variation, two independent DNase-seq libraries were generated from one biological husk sample and the number of shared DNAse I hypersensitive sites (DHSs) were counted after MACS2 peak calling [47]. The two replicates shared 14,401 DHSs (66% and 88% of the peaks in replicate 1 and 2, respectively, Table S2). We concluded that the results from the technical replicates were comparable. The reads from the two technical replicates were therefore pooled and treated as one biological replicate in the further analysis.

H3K4me3 ChIP-seq data was obtained from the NCBI database (SRX1073672; [61]). The quality filtering of the sequencing data was done in the same way as described in BS-seq analysis section. The reads were aligned to the reference genome using BWA [107]. Non-uniquely mapped reads were filtered out with a MAPQ cut-off value of 20 using samtools [108]. Peaks were called for each biological replicate with a q-value cut-off of 0.001 using MACS2 [47]. During the peak calling, naked DNA digestion data and input control data were used as controls for DNase-seq and for ChIP-seq, respectively. Only peaks with 70% or larger overlap between replicates were kept for analysis. If there were three replicates, overlapping peaks in two replicates were identified first and then the third replicate was compared to the already-integrated peaks.

BS-seq

Raw data of genome-wide bisulphite sequencing experiments on wild-type B73 coleoptile shoot tissue (harvested 5 days after the start of germination) [35] was obtained from the NCBI database (GSE39232). FastX toolkit [109] was used to filter artefacts introduced by library
construction such as linker and/or adaptor sequences, and to filter reads of which the qualities of more than 80% of the bases were lower than a threshold of Q20. The reads were trimmed based on their per base sequence qualities and reads shorter than 70 bases after trimming were removed using PRINSEQ [110]. The read mapping to the reference genome and methylation base calling was performed using BS-seeker2 [111]. The low and un-methylated regions were identified for both CG and CHG data using MethylSeekR [31]. The threshold for percent methylation for the low methylated regions was set to 20%. MethylSeekR [31] defines unmethylated regions (UMRs) and low methylated regions (LMRs); in this study, we combined both regions into one class, LUMRs (Low and UnMethylated Regions). Any identified regions with more than or equal to 20% DNA methylation using bwtool [48] were further filtered out. For enhancer identification, regions with both low CG and low CHG methylation, which were identified using BEDtools [102], were called as LUMRs. The methylation frequency at every mCG, mCHG and mCHH position was extracted for further analysis.

**Characterisation of each data set**

Genomic regions were defined as follows: genic regions, exons and transposable elements (TEs) were annotated according to the reference annotation. The annotated exons include the untranslated regions (UTRs). The entire genome, except for the genic regions, were called intergenic regions. Introns were genic regions excluding exons. Promoters were defined as the sequence 1 kb upstream and 200 bp downstream of TSSs. Flanking regions were defined as sequences 4 kb upstream from promoter regions and 5 kb downstream from the TTSs. Distal regions were intergenic regions that were not classified above. Uniquely mappable regions in the whole genome were identified using Uniqueome [42] for theoretical read lengths of 93 bp, which was the longest read possible for the ISAS uniqueome aligner (http://www.imagenix.com) to handle and closest to the actual read length (100 bp), allowing 2 mismatches. The ISAS uniqueome aligner performs all-against-all sequence alignment with a given read length (93 bp in this case), and deduces percent uniqueness for each nucleotide position based on the percentage of reads mapped to this position that are uniquely mapping at this location. In this study, uniquely mappable regions showed 90% or higher uniqueness. The number of uniquely-mappable base pairs within each genomic region was counted using BEDtools [102] and plotted using plotrix package [112] in R [113].
The total lengths of each genomic region in Mbs and the numbers of features (DHS, H3K9ac and LUMR) overlapping with the defined genomic regions were counted using BEDtools [102] and plotted using R [113].

For correlations between gene expression levels and DNase hypersensitivity or H3K9ac enrichment, first the genes were binned based on their expression levels in RPKM from the lowest (bin 0) to the highest (bin 6). Bin 0 contains all the genes with no and lower than 1 RPKM expression. The other 6 bins were defined so that each bin contained exactly the same number of genes. The average intensities of DNase hypersensitivity and H3K9ac enrichment in RPM over genic regions were calculated using bwtool [48] and plotted using R [113].

**Data integration**

**Candidate identification**

The enhancer prediction in this study was focused on active enhancers. The DNase I hypersensitivity, H3K9ac enrichment and LUMR data were integrated. All LUMRs that overlap with DHSs and H3K9ac (Figure 2), excluding the ones overlapping with genes and promoter regions, and the numbers of candidates overlapping with TEs and CNSs were selected and counted using BEDtools [102]. The CNS coordinate data was extracted from published rice v6 vs maize v2 data [66], and the coordinates were converted from v2 to v4 using Assembly Converter available on Ensembl Plants [40].

**TE enrichment analysis:**

TE annotations are available at the Gramene database:
ftp://ftp.gramene.org/pub/gramene/CURRENT_RELEASE/data/gff3/zea_mays/repeat_annotation/ [67].

TE families have been named according to the guidelines described at the Maize transposable element (TE) database:
http://maizetedb.org/cgi-bin/cgiwrap/maize/TE_show_family.cgi?do_table=1.

To prepare the annotation file, nested TE insertions were resolved using RTrackLayer [114] in R [113]. Bedtools intersect [102] was then used to find overlaps between enhancer candidate coordinates and TE coordinates. Enhancers candidates that were at least 80% contained within a single TE were selected for further analysis. To create a baseline for the number of TEs that could contain an intergenic enhancer candidate, the full list of TEs was filtered for elements not
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contained within introns and that are longer than 635 bp, long enough that the enhancer candidates at the 20th percentile by length could overlap 80% of a TE. The filtered TE set was used as the baseline for number of elements within families containing enhancer candidates (Additional file 4: Dataset 3) and for comparisons between TEs with and without enhancer candidates. Conserved sequence motifs were identified using four de novo motif discovery tools, HOMER, MEME, DREME and RSAT plants [68–71] and enhancer candidates, the TE family RLG00010 and randomly selected intergenic sequences of the corresponding size were scanned for the identified motifs using FIMO [77].

Heatmap plot

For DNase I hypersensitivity and H3K9ac enrichment, RPM signal tracks were generated from pooled data during peak calling using MACS2 [47]. DNase I hypersensitivity, H3K9ac enrichment and transcript coverage data, Methylation frequency data and TE annotation data (0 = absence, 1 = presence of TEs) were converted to BigWig files using wigToBigWig tool [115]. For the DNA methylation data, methylation frequency over 100 bp fixed-windows were calculated using bwtools [48].

The data on DNase I hypersensitivity, H3K9ac enrichment, mCG, mCHG and mCHH levels, TE presence and transcript levels were extracted for each DHS and its 1 kb flanking regions in our candidate list using bwtool [48]. The DHSs were clustered based on H3K9ac enrichment with k-means clustering, re-ordered, and all the data sets were plotted according to the order defined based on H3K9ac k-means clustering using the gplots package [116] in R [113]. For the heatmap profile at TSSs, 429 DHSs mapped at TSSs in V2-IST and 1,400 in husk were randomly selected and heatmaps were generated in the same manner as for the DHSs in the candidates. To make the heatmaps comparable, for DHSs at TSSs, the same number of DHSs were selected as the number of DHSs in candidates in the two tissues.

Genomic feature profiling at DHSs in enhancer candidates and TSSs

To understand the behaviour of H3K9ac enrichment and DNA methylation around DHSs at our potential candidate regions, average profiles were generated. Firstly, all the intergenic DHSs were taken. For each DHS, H3K9ac enrichment values 300 bp up- and downstream were extracted using bwtool [48] and the end with higher H3K9ac enrichment was defined as 3' end of the DHS. Using bwtool [48], the average RPM for DNase I hypersensitivity and H3K9ac
enrichment and methylation frequencies at CG, CHG and CHH were calculated at the intergenic DHSs and their flanking regions. The values were plotted using R [113]. Average profiles for TSSs were generated in the similar manner except the DHSs were oriented based on their gene strand. For generating average plots for each category, the DHSs were first binned by the categories and average values were calculated for each bin.

**Candidate ranking**

Once enhancer candidates were identified, they were ranked according to their presumed tissue-specificity. We assumed that the tissue-specificity of an enhancer is correlated to its DNase I hypersensitivity and H3K9ac enrichment. Therefore, the tissue-specificity of each candidate was determined using the largest differences in DNase I digestion sensitivity and H3K9ac enrichment between the two tissues (Figure 2). For each candidate, for both the DNase hypersensitivity and H3K9ac enrichment separately, the intensity differences in the candidate region were calculated from the signal tracks and the largest values were taken as the difference using bwtool [48]. The candidates were then ranked based on the differences in DNase I hypersensitivity and H3K9ac enrichment independently, and the DNase I and H3K9ac rankings were summed for each candidate. Then, the enhancer candidates were re-ranked based on the sum. The V and H numbers provided in the additional files 2 and 3 show the final ranking after the summation. For p-value calculation, two (or three for H3K9ac husk data as it had three replicates) lists of numbers (1 to 398 for V2-IST, 1 to 1320 for husk tissue, the same number as the numbers of candidates) were generated. Random combinations of two (or three) numbers were summed and re-ranked according to the sum for 1,000 times to create lists of theoretical summation scores for each ranking. The frequency occurrence of the value less than or equal to the real data in the theoretical score list was computed and provided as p-values.

**Linking enhancer candidates to potential target genes**

Enhancer candidates were linked to putative target genes based on the defined tissue-specificity of candidates and expression data of nearby genes. The assumption was that an enhancer targets its closest up- or downstream gene. Firstly, gene expression levels and the statistical significance of their differential expression data from Cuffdiff [86] were linked to the gene coordinate data. The closest up- and downstream genes were identified for each candidate using BEDtools [102]. For tissue-specific candidates, significantly differentially expressed genes
were identified first, then the tissues in which the genes were expressed higher were identified. When the tissue-specific gene expression levels matched with the tissue-specificity of the candidate, the gene(s) was linked to the candidate. For example, if one of the candidates was determined as V2-IST-specific and the upstream gene had higher expression in V2-IST than in husk, we concluded that the candidate most probably regulates its upstream gene. For shared candidates, adjacent genes being expressed in both tissues were associated.

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Availability of data and material
The DNase-seq data, H3K9ac ChIP-seq data and RNA-seq data from B73 V2-IST and husk tissue were deposited in the NCBI Gene Expression Omnibus (GEO) (http://www.ncbi.nlm.nih.gov/geo/) under accession number GSE94291, GSE94251 and GSE94252.
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Authors' contributions
R.O., J.Z., B.W., H.C.J.H., F.T. and M.S. designed the research, J.Z., B.W. cultivated the plants and performed the experiments, R.O. and J.Z. performed genome-wide data analysis, J.J.W. provided general help in NGS data analysis, J.G. helped with the BS-seq data analysis, S.N.A and N.M.S performed the TE analysis, R.O. and H.C.J.H. performed statistical analysis, C.H. contributed to the candidate ranking method and R.O., J.Z., B.W., H.C.J.H., F.T. and M.S. wrote the paper.
Figure S1. The tissues used in this study. (A) Inner stem tissue (at the right) isolated from V2 seedling (left) and (B) soft husk leaves (at the right) isolated from maize husk (left).
Figure S2. Reproducibility of RNA-seq data. Gene expression levels (in read counts) were plotted for all genes. Read counts were normalised by the total number of reads in the respective replicates. (A) V2-IST samples, (B) Husk samples. Pearson correlation coefficients are given in each scatter plot.
Figure S3. Randomised distributions of features over genomic regions within the uniquely mappable part of the genome. Distributions of (A, B) DHSs, (C, D) H3K9ac (E) LUMRs and (F, G) enhancer candidates over the different genomic regions. The numbers below the pie charts are the total numbers of the indicated features in the genome. Inner ring: intergenic (red) and genic (blue); outer ring: intergenic TEs (yellow), distal (sand), and flanking (peach) regions, promoters (dark green), exons (light green), and introns including intronic TEs (mint) as depicted in Figure 3A.
Figure S4. Characteristics of genomic features and distribution of gene expression levels. (A) Size distributions (in kilobase pairs) of chromatin and DNA features as indicated on x-axis categories. Left panel shows full ranges including all outliers, right panel a zoom into the data. Boxes include second and third quartile, the median is indicated by a horizontal black line. Whiskers extend the interquartile range by 1.5, values beyond are plotted as outliers. (B) Cumulative fraction of genes contributing to total expression quantified as sum of normalized (RPKM) RNA-seq reads in V2-IST (blue) and husk (red).
Figure S5. Examples of (A and B) enhancers that contain TEs and (C and D) TEs that contain an enhancer. Jbrowse tracks show from the top: AGPv4 TE (ochre) [1] and candidate enhancer (turquoise) annotation (H stands for husk candidate), DNase-seq (blue) and H3K9ac (green) coverage, and peak position (indicated as bars) in V2-IST and in husk, mCG, mCHG and mCHH levels [2] and Uniqueness as mappability in percentage. (A, B) H411 and H1153 containing DTH00410 and DTH00409, respectively, both belong to the TIR TE order. (C, D) RLG00010 family members containing enhancer candidates H864 and H545. Note that tracks for accessibility and H3K9ac coverage represent pooled replicates. Peaks at those tracks might not be present in each replicate, affecting enhancer candidate prediction.
Figure S6. Characteristics of enhancer-overlapping TEs. Comparison of (A) the lengths of all TEs and per major TE order. (B) Distribution of distance between TEs and their closest flanking genes. (C) Relative locations of enhancer candidates within TEs per major order. (D, E) Comparison of RLG00010 family members with and without enhancer candidate for (D) distances to their closest flanking genes and (E) TE lengths. Boxes include second and third quartile, the median is indicated by a black line. Whiskers extend the interquartile range by 1.5, values beyond are plotted as outliers.
**Figure S7. Example of data on *tb1* enhancer.** From the top: AGPv4 genome annotation [1], candidate annotation from our prediction (H stands for husk candidate), DNase-seq and H3K9ac ChIP-seq read coverage and data on peak position (indicated as bars) in V2-IST and in husk, mCG, mCHG and mCHH levels [2] and unique mappability in percentage. The number under the gene name indicates gene expression levels (V2-IST/husk). The dark blue bar in the gene annotation track indicates the previously annotated *tb1* enhancer. The vertical red box indicates the enhancer candidate identified in this study. Note that tracks for accessibility and H3K9ac coverage represent pooled replicates. Peaks at those tracks might not be present in each replicate, affecting enhancer candidate prediction.
Figure S8. Asymmetric H3K9ac enrichment at candidate DHSs. Distributions of H3K9ac enrichment 300 bp up- and downstream from the boundaries of DHSs in 431 V2-IST and 1,437 husk enhancer candidates (A, B) before and (C, D) after orientating the candidates based on the H3K9ac enrichment values 300 bp up- and downstream from the DHS boundaries; the higher H3K9ac values were plotted on the x-axis, the lower on the y-axis. In C and D, the number of data points below the line were 241 in V2-IST and 841 in husk, indicating asymmetrical H3K9ac enrichment.
Figure S9. Average profiles of the enhancer candidates in V2-IST and husk for each category. Average signal intensities of DNase I hypersensitivity, H3K9ac enrichment in RPM and DNA methylation levels in methylation frequency [2] at DHSs and their 1 kb flanking regions. The categories, identified in heatmaps by clustering DHSs on H3K9ac enrichment data using a k-means (k=4) algorithm (Figure 5), were numbered from 1 to 4 from the top of the figure to the bottom. The DHSs were oriented based on the flanking signals of H3K9ac enrichment (higher H3K9ac enrichment on the 3' side). The various sizes of DHSs have been scaled to an equal size. The numbers between brackets represent the number of enhancer candidates in the category and tissue indicated.
Figure S10. Heatmaps of chromatin, DNA and transcript features at TSSs. DNase I hypersensitivity, H3K9ac enrichment, mCG, mCHG and mCHH levels [2], presence of TE [1], and transcript levels at and around (+/- 1 kb) DHSs at TSSs (431 and 1,437 in V2-IST and husk, respectively). The selected regions were oriented according to the 5’ to 3’ orientation of the genes. The various sizes of DHSs have been scaled to an equal size. The colour scales are in RPM for DNase I hypersensitivity, H3K9ac enrichment and transcript levels, and in methylation frequency (0-1) for DNA methylation. For TE, red and white shows the presence or absence of TE, respectively. The DHSs were clustered on the H3K9ac enrichment data using a k-means (k=3) clustering algorithm.
Figure S11. Average profiles of randomly selected TSSs in V2-IST and husk for each category. 431 V2-IST and 1,437 husk TSSs overlapping with DHSs and H3K9ac were randomly selected. The DHSs at the TSSs were oriented according to the 5’ to 3’ orientation of the genes. (A, B) Average signal intensities of DNase I hypersensitivity, H3K9ac enrichment in RPM and DNA methylation levels in methylation frequency [2] at DHSs and their 1 kb flanking regions. (C-H) The three categories, identified in heatmaps by clustering DHSs on H3K9ac enrichment data using a k-means (k=3) algorithm (Figure S10), were indicated.
Figure S12. Comparison of expression levels between genes and enhancer candidates in V2-IST and husk. Boxplot showing expression levels (RPKM) for genes and enhancer candidates in V2-IST and husk tissue. Boxes include second and third quartile, the median is indicated by a black line. Whiskers extend the interquartile range by 1.5, values beyond are plotted as outliers.
Figure S13. Heatmaps and average profiles of H3K9ac and H3K4me3 at (A, B, D) candidates and (A, C, E) TSSs. (A) The H3K9ac clustered data in V2-IST and husk tissue are the same data as shown in Figure 5 and S10. In addition, H3K4me3 enrichment data [3] at the same loci was plotted in the same manner as for H3K9ac. (B-E) Average H3K9ac and H3K4me3 enrichment signal intensities are shown.
Figure S14. ChIP-qPCR data of H3K9ac and H3K4me1 enrichment at enhancer candidate regions. Genome browser view and levels of H3K9ac and H3K4me1 enrichment at (A) enhancer candidate H108, (B) b1 enhancer (H167), (C) DICE (V4/H1318), and (D) tb1 enhancer (H1233). The genome browser view shows H3K9ac enrichment as detected by ChIP-seq at each candidate for both V2-IST and husk. The sequences examined by qPCR are indicated by coloured boxes. Red and yellow and black boxes indicate the summits (P) and slopes of H3K9ac peak regions (S), respectively; black boxes indicate regions without H3K9ac enrichment (O). Below the top panels, the levels of enrichment for H3K9ac and H3K4me1 in V2-IST (green bars) and husk (purple bars) at the regions of interest relative to those at the actin locus are shown. Error bars indicate the standard error for three biological replicates.
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