

Aluminum tolerance in maize is associated with higher *MATE1* gene copy number

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Genome structure variation, including copy number variation and presence/absence variation, comprises a large extent of maize genetic diversity; however, its effect on phenotypes remains largely unexplored. Here, we describe how copy number variation underlies a rare allele that contributes to maize aluminum (Al) tolerance. Al toxicity is the primary limitation for crop production on acid soils, which make up 50% of the world's potentially arable lands. In a recombinant inbred line mapping population, copy number variation of the Al tolerance gene multidrug and toxic compound extrusion 1 (*MATE1*) is the basis for the quantitative trait locus of largest effect on phenotypic variation. This expansion in *MATE1* copy number is associated with higher *MATE1* expression, which in turn results in superior Al tolerance. The three *MATE1* copies are identical and are part of a tandem triplication. Only three maize inbred lines carrying the three-copy allele were identified from maize and teosinte diversity panels, indicating that copy number variation for *MATE1* is a rare, and quite likely recent, event. These maize lines with higher *MATE1* copy number are also Al-tolerant, have high *MATE1* expression, and originate from regions of highly acidic soils. Our findings show a role for copy number variation in the adaptation of maize to acidic soils in the tropics and suggest that genome structural changes may be a rapid evolutionary response to new environments.

abiotic stress | selection

Aluminum toxicity severely limits plant growth on highly acidic soils (pH <5). Under these conditions, the rhizotoxic Al species Al³⁺ is solubilized, inhibiting root growth and function (1), and leaving plants more vulnerable to drought and mineral nutrient deficiencies. Approximately 30% of the earth's total land area consists of highly acid soils, and as much as 50% of the world's potentially arable lands are acidic (2). As large areas of acid soils in the tropics and subtropics are critical food-producing regions, Al toxicity constitutes a food security threat exceeded only by drought with regard to abiotic limitations on crop production (2).

A major physiological mechanism of plant aluminum tolerance consists of the Al-activated release of organic acids from the root apex, the site of Al toxicity (1). Organic acid anions such as malate, citrate, and oxalate chelate Al³⁺ in the rhizosphere forming stable, nontoxic complexes (3). This “exclusion” mechanism limits Al³⁺ uptake by the roots, minimizing harmful interactions with Al-sensitive sites in the root apical apoplast and/or inside root cells, and has been correlated with differential Al tolerance in a large number of monocot and dicot species (1). Transporters from the multidrug and toxic compound extrusion (*MATE*) family have been shown to mediate the release of citrate, and contribute to Al tolerance in a number of plant species (4–9).

Maize (*Zea mays*) is a major food crop throughout the tropics and subtropics, where acid soils are prevalent. Therefore, maize breeding programs in these regions have focused intensely on Al tolerance. The development of Al-tolerant varieties of maize and other crops has been crucial for the agricultural development of

previously unproductive regions such as the Brazilian Cerrado, an area of 205 million hectares (10). Al tolerance in maize is a quantitative trait (11–13), and quantitative trait loci (QTLs) associated with the trait have been identified. Five QTLs explaining 60% of the variance in Al tolerance were detected in a mapping population generated from a cross between a highly Al-tolerant tropical inbred line commonly used as a tolerance donor in breeding programs, Cateto AI237, and the Al-sensitive inbred line L53 (14).

We recently reported on the characterization of the maize Al tolerance gene *MATE1* (15), originally identified in a microarray study (16) as the most up-regulated gene in root tips of an Al-tolerant maize line under Al stress. *MATE1* was mapped to the telomeric region of chromosome 6, colocalizing with the QTL of largest effect on Al tolerance identified by Ninamango-Cárdenas et al. (14), explaining 16.2% of the phenotypic variance. *MATE1* shares significant identity to its homologs in sorghum and *Arabidopsis*, SbMATE (52%) and AtMATE (64%). *MATE1* mediates citrate efflux, as observed in [¹⁴C]citrate efflux studies in *Xenopus laevis* oocytes. Expressing *MATE1* in transgenic *Arabidopsis* conferred a significant increase in Al tolerance and root citrate exudation in response to Al. In maize, *MATE1* expression is concentrated in the root apex and is strongly up-regulated by Al. *MATE1* expression is significantly higher in the Al-tolerant parent AI237 (and in Al-tolerant C100-6, used in the original microarray study) than in Al-sensitive L53, both in the absence or presence of Al (15).

Our recent findings suggested that functional, structural, or regulatory variation in *MATE1* may underlie the large-effect Al tolerance QTL previously detected on chromosome 6. In the present study, we show that copy number variation (CNV) in the *MATE1* locus is associated with both gene expression and phenotypic differences within our recombinant inbred line (RIL) population. We also demonstrate that structural variation in this locus is a rare and probably recent occurrence, and discuss its potential implications for maize adaptation to acid soils.

Results

Cloning and sequencing the *MATE1* ORFs from the parents of the mapping population (AI237, Al-tolerant, and L53, Al-sensitive)

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previously revealed six nucleotide differences between their coding regions, of which only two resulted in amino acid substitutions (R171C and G527V) (15). We performed further functional investigations to establish whether these two structural modifications resulted in differences in the transporter's basic functional properties (Fig. S14). For this purpose, the proteins encoded by each of the two *MATE1* alleles were functionally characterized in *Xenopus laevis* oocytes (Fig. S1 B–D). Oocytes expressing either allele showed similar time- and pH-dependent electrogenic activities (i.e., negative currents), which were significantly larger than those of control cells (Fig. S1 B and C). We also examined substrate recognition by testing the effect of exogenous citrate or malate added to the bath media (Fig. S1D). Addition of citrate or malate increased *MATE1*-mediated inward currents, with the changes in current magnitude being dependent on the extracellular concentration of organic acid supplied. Again, no differences between oocytes injected with either allele were observed under these conditions.

We did not identify any significant structural or functional differences of the *MATE1* proteins encoded by the parental alleles in our analysis that could explain the difference in Al tolerance. However, we previously showed that *MATE1* expression is higher in Al237 than in L53, both in the absence and presence of Al (15). These results suggest that the causal polymorphism underlying the QTL must somehow affect this gene's expression levels. Therefore, we looked for sequence polymorphisms in potential *cis* regulatory regions, by cloning an 8-kbp fragment upstream of the *MATE1* ORFs from both parental lines (Fig. S2). The sequences revealed extremely low levels of polymorphism (99.9% identity) between the two lines, suggesting that sequence polymorphism in *cis* regulatory regions are also not likely to explain the Al tolerance QTL.

CNV in *MATE1*. To assess whether a structural polymorphism is associated with the expression differences observed between the parental lines, we quantified gene copy number using quantitative real-time PCR (qPCR). The method was validated using sequences with known copy number in the reference maize line B73 and indicated that, although B73 and L53 have a single copy of

MATE1, three copies are found in the Al237 genome (Fig. S3). The location of the *MATE1* gene copies on maize chromosomes of inbred lines Al237 and L53 was investigated using fluorescent in situ hybridization (FISH). We localized *MATE1* on the short arm of chromosome 6 in both lines, distal to the nucleolar organizing region (Fig. 1), in agreement with our mapping results (15). Minor signals were also consistently detected on 1L, 7S, and 9L. Weak or infrequent signals were observed on 2L, 3L, 7L, 8L (Al237 only), and 10L. The sequence locations of partial gene transpositions of *MATE1* located on chromosomes 2, 7, and 10 (Fig. S3) are consistent with the minor FISH signals observed on these chromosomes; however, their precise locations cannot be determined at this FISH resolution level. Nevertheless, these results indicate that the *MATE1* copies in Al237 must be located near each other on the short arm of chromosome 6, and that no gene copies undetected by qPCR exist at other genomic locations. In addition, although only one signal per chromatid was observed in all L53 images ($n = 17$), in over 70% of Al237 images (15 of 21), partially overlapping *MATE1* signals were observed on a single chromatid (Fig. 1C). These results are consistent with multiple copies of the gene being present at that location.

***MATE1* CNV and Al Tolerance.** Taking into account that the parents of the population used to map Al tolerance QTL vary in both *MATE1* copy number and *MATE1* expression levels, we set out to investigate the relationship between these genotypic differences and the Al tolerance phenotype. First, we determined the distribution of *MATE1* copy number in the Al237×L53 RILs using qPCR (Fig. 2A and Dataset S14). The proportion of progeny with one versus three gene copies conformed to a 1:1 segregation ratio: 57 RILs have a single *MATE1* copy, and 53 RILs have three copies (P value of χ^2 test = 0.7029). These results are in agreement with what was observed using FISH and suggest that the paralog copies are in tight genetic linkage. Then, we examined the relationship between CNV in the *MATE1* locus and phenotypic variation for Al tolerance in the Al237×L53 RILs. *MATE1* copy number is positively correlated with Al tolerance in the RIL population ($P = 5.46 \times 10^{-7}$, $R^2 = 0.20$), such that individuals with

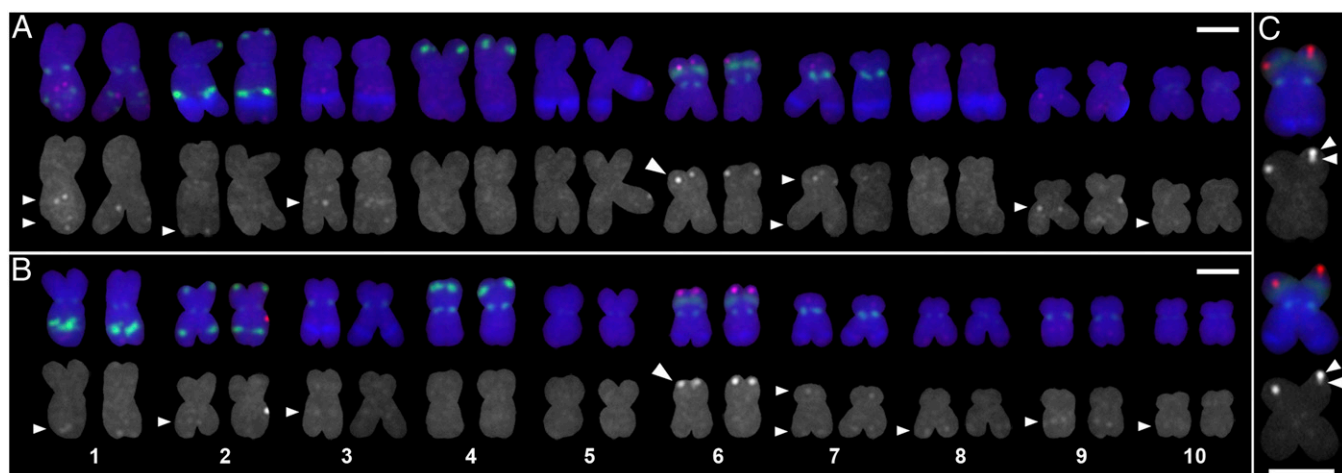


Fig. 1. Localization of *MATE1* in maize lines L53 and Al237 using FISH. Homologous metaphase chromosomes from root meristem cells are aligned as pairs. (A) L53, Al-sensitive, one copy of *MATE1*. (B) Al237, Al-tolerant, three copies of *MATE1*. A large arrowhead indicates the location of *MATE1* on 6S; small arrowheads indicate minor signals. (C) Multiple *MATE1* FISH signals are observed on Al237 chromosome 6S. Multicolor image and grayscale version of the respective red channel are shown. The arrowheads indicate regions with two *MATE1* signals per chromatid. The *MATE1* probe is labeled with Texas Red. Green fluorescein isothiocyanate-labeled probes include centromere C repeat sequences (found on all chromosomes but copy number highly variable), a TAG microsatellite repeat (chromosome arms 1L, 2S, 2L, and 4S), and the ribosomal gene cluster (6S at the nucleolar organizing region). Blue DAPI counterstain was used to visualize heterochromatic 180-bp knob regions (line-specific locations). The second karyotype shown in each panel is the grayscale version of the *MATE1* probe (red) channel in the respective multicolor karyotype. Adjustment of individual color channels was performed on original images. (Scale bars, 3 μ m.)

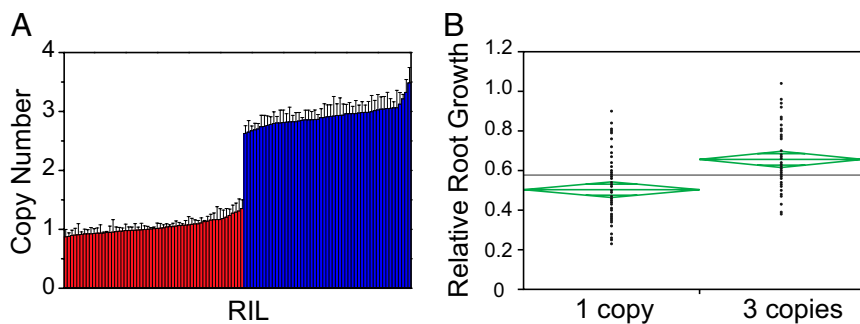


Fig. 2. *MATE1* CNV in Al237xL53 RILs correlates with aluminum tolerance. (A) *MATE1* copy number in the RIL population derived from a cross between Al237 (Al-tolerant, three copies of *MATE1*) and L53 (Al-sensitive, one copy of *MATE1*), determined by qPCR. RILs are arranged based on copy number in ascending order (from one copy to three copies). The red bars denote single-copy RILs ($n = 57$); the blue bars denote RILs with three copies of *MATE1* ($n = 53$). Error bars indicate 1 SD. (B) One-way ANOVA of RRG (standard phenotypic index of Al tolerance) by *MATE1* copy number in the Al237xL53 RIL population. Mean diamonds represent 95% confidence intervals, and mid-bars in diamonds represent the group mean. The gray line represents the overall mean.

three copies have a significantly higher mean Al tolerance than those inheriting a single copy (Fig. 2B).

The results above strongly suggest that greater gene copy number is the basis for higher *MATE1* expression levels in the RILs, which in turn results in increased Al tolerance. Thus, we also examined how *MATE1* expression relates directly to gene copy number in the Al237xL53 population. Before designing gene expression assays, we performed extensive Sanger sequencing of the entire coding region of *MATE1* from Al237 genomic DNA to identify SNPs or indels between the three gene copies. Strikingly, no polymorphisms were identified anywhere in the sequence: the exons, introns, 5'- and 3'-UTRs of the three *MATE1* copies in Al237 are 100% identical. Therefore, any measurement of *MATE1* expression will capture the combined expression levels of the three copies. With this in mind, we used qPCR to determine root tip *MATE1* expression in the Al237xL53 population, in plants grown under control and Al stress conditions (Fig. 3A and Dataset S1A).

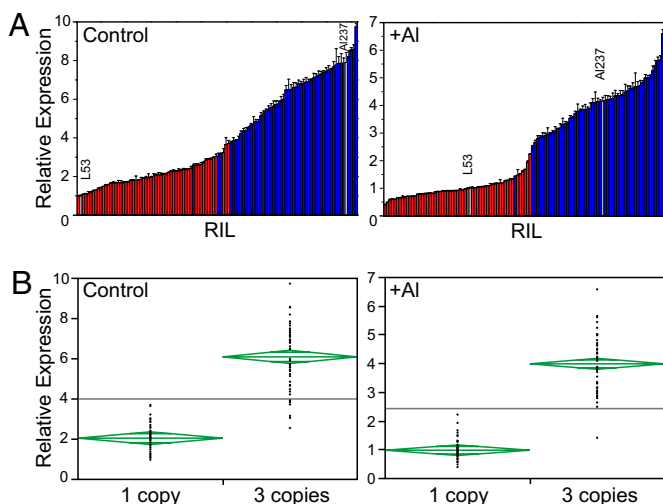


Fig. 3. *MATE1* expression in the Al237xL53 RIL population correlates with gene copy number. (A) *MATE1* relative expression in root tips quantified via qPCR from maize seedlings grown under control conditions (Left), or treated with 39 μM Al^{3+} activity (Right) for 24 h. In each graph, maize lines are arranged based on *MATE1* relative expression in ascending order. The bars are color-coded based on *MATE1* copy number: the red bars denote one-copy RILs, and the blue bars denote three-copy RILs. The gray bars denote the parents of the population, Al237 (Al-tolerant, three copies of *MATE1*) and L53 (Al-sensitive, one copy of *MATE1*). Samples were calibrated against L53 (relative expression = 1) within each dataset. Error bars indicate 1 SD. (B) One-way ANOVA of *MATE1* relative expression by *MATE1* copy number in the Al237xL53 RIL population, in control and Al stress conditions. Mean diamonds represent 95% confidence intervals, and the mid-bars in diamonds represent the group mean. The gray line represents the overall mean.

MATE1 expression in the RILs varies continuously under both growth conditions. In control conditions, the parents represent the extremes of the distribution (L53, low expression; Al237, high expression). Under Al stress, there is a greater number of RILs in which we observed *MATE1* expression lower than L53 or higher than Al237, indicating that some degree of transgressive segregation exists for *MATE1* expression under Al. Because *MATE1* expression is up-regulated under Al stress, variation in *trans*-acting elements involved in the regulation of this response is likely to account for this effect. *MATE1* copy number is also strongly correlated with *MATE1* expression (Fig. 3B), such that RILs carrying three copies of the gene have significantly higher expression levels than RILs carrying one copy, both in the absence ($P = 1.48 \times 10^{-33}$, $R^2 = 0.74$) and in the presence ($P = 1.23 \times 10^{-42}$, $R^2 = 0.82$) of Al. Thus, differences in *MATE1* expression between RILs can be largely explained by differences in gene copy number, both under control and Al stress conditions. This observation is also supported by the fact that, within the RIL population, *MATE1* expression levels under Al stress are highly correlated with those under control conditions ($R^2 = 0.64$; Fig. S4), so that RILs with higher expression in the absence of Al tend to also display higher expression when exposed to the stress.

Genetic Factors Controlling *MATE1* Expression. We investigated the genetic factors controlling *MATE1* expression via expression QTL (eQTL) mapping in the Al237xL53 population. For this purpose, a high density of markers was generated via genotyping-by-sequencing (GBS), a restriction enzyme-based method for constructing reduced representation genomic libraries for genotyping (17). Using this new marker set, we repeated the mapping of Al tolerance QTL and also mapped eQTL controlling *MATE1* expression (Fig. 4). eQTL mapping was performed independently for *MATE1* expression both in the presence and in the absence of Al. The mapping of loci controlling Al tolerance in the Al237xL53 population confirmed our previous results showing that a genomic region on the distal end of chromosome 6, colocalizing with the *MATE1* locus, represents a major Al tolerance QTL. Mapping of *MATE1* expression identified a single genomic region controlling the trait, coincident with the location of the gene, under both control and Al stress conditions. These results indicate that *MATE1* expression is controlled mainly in *cis*. *Trans* factors involved in the up-regulation of *MATE1* expression under Al stress may exist, but their smaller effect is overshadowed by the large effects in *cis*; thus, they were not detected by eQTL mapping. Data from near-isogenic lines (NILs) also corroborate these results. NILs generated by introgressing the three-copy *MATE1* allele from Al237 into the Al-sensitive L53 background are nearly twice as tolerant as L53 (averaging 50% root growth inhibition, versus 80% in L53; Dataset S1B). The level of *MATE1* expression in root tips of these NILs is comparable to those of Al237, indicating that no

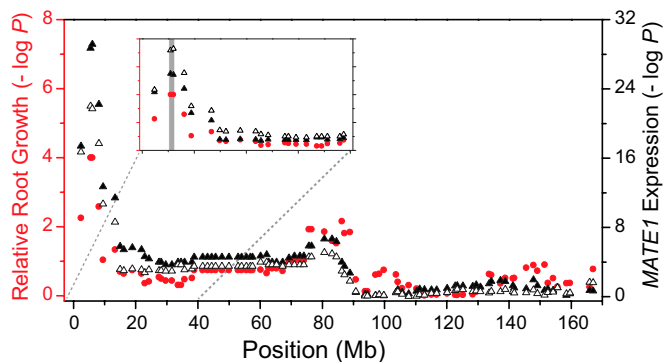


Fig. 4. A major aluminum tolerance QTL and eQTL for *MATE1* expression colocalize with *MATE1*. A GLM analysis identified GBS-generated SNP markers that are associated with Al tolerance (expressed as RRG; red circles, y axis on Left) and with *MATE1* expression (open triangles: expression under control conditions; solid triangles: expression under Al treatment; y axis on Right). The level of statistical association for each polymorphism is expressed as $-\log P$. x axis: physical position on maize chromosome 6. *Inset* is a magnification of a 40-Mb window on the top of chromosome 6. A gray bar indicates the location of the *MATE1* locus.

other loci acting in *trans* are necessary to retain the high levels of expression observed in the Al-tolerant parent, Al237.

Genomic Organization of the *MATE1* Locus. To examine the genomic structure of the *MATE1* locus and determine how the three gene copies are organized, a genomic BAC library from the Al-tolerant parent Al237 was constructed. A BAC library pooling and screening strategy was developed that did not require individual BACs to be arrayed (*SI Materials and Methods*). A positive BAC clone identified from the library screening was shown to carry three copies of *MATE1* and was fully sequenced via single-molecule real-time sequencing technology, using a combination of small- and large-insert sequencing libraries (600 bp and 8 kb; *SI Materials and Methods*). Although the long, single-molecule reads facilitate the assembly of repetitive DNA regions, the typical error rates generated by this technology are high. Therefore, the sequences from the small-insert library were used to error-correct the long reads generated by sequencing of the large-insert library, using a hybrid approach (18). The error rate of the long reads was reduced to less than 1%, as estimated based on the known vector sequence. After error correction, the BAC sequence was assembled manually from the corrected long reads to circumvent the limitation of existing assemblers in generating contigs from long repetitive sequences, with supporting data from long-range PCR and from depth-of-coverage analysis of the error-corrected long reads aligned along the BAC sequence (*SI Materials and Methods*) (Fig. S5).

Sequencing the genomic BAC clone revealed that the three *MATE1* copies are part of a tandem triplication, with each repeat unit totaling nearly 30 kbp in size (Fig. 5). Except for *MATE1-1*, -2, and -3, no other maize genes are predicted within the BAC sequence. Instead, each copy of *MATE1* is flanked by various transposon insertions, mainly upstream of each ORF. The three 30-kbp repeats share high sequence identity with each other. The second repeat region differs from the first and third repeats, in that a class I long terminal repeat (LTR) Gypsy retrotransposon is inserted just upstream of *MATE1-2*. This insertion is not present in the upstream sequence cloned from L53 (Fig. S2). Compared with its syntenic region, the BAC sequence shares no substantial sequence identity to the B73 reference sequence except within the *MATE1* coding region. This is not surprising, given the degree of heterogeneity in genome organization observed in maize. In fact, extensive noncolinearity in intergenic regions has been observed between different inbred lines for other loci (19–22).

Pervasiveness of *MATE1* CNV. Taking into account the effect of *MATE1* CNV on Al tolerance in the Al237×L53 population, we sought to probe the prevalence of this variation in a broad maize genetic pool. For this purpose, we surveyed a panel of 166 diverse maize inbred lines and 16 teosinte lines (23) for *MATE1* CNV (Dataset S1C). From a maize association panel (24) phenotyped for Al tolerance (Fig. 6A), we determined *MATE1* copy number via qPCR in lines with intermediate-to-high levels of Al tolerance [root growth inhibition below 50%, i.e., relative root growth (RRG) of >0.5]. We also included maize inbreds known as “founder” lines, which capture most of maize genetic diversity (25), regardless of their level of Al tolerance. The results of our survey show that *MATE1* CNV is rare: only two maize inbred lines with more than one copy of *MATE1* were identified (C100-6 and Il677a; three *MATE1* copies). Similar to the Al-tolerant parent of our population, Al237, these lines are Al-tolerant, ranking 6th and 29th (respectively) out of the 279 maize lines that were phenotyped for Al tolerance (Dataset S1C and Fig. 6A). Remarkably, maize lines with three *MATE1* copies can be traced to the same regional origin, the acid soils of the South American tropics. Al237 (formerly L1327) and C100-6 are Cateto inbreds originally from Brazil. The other maize inbred line found to carry three *MATE1* copies, Il677a, is a sweet corn inbred with a South American parent. This line (pedigree [(Bolivia 1035*Il44b)*Il422a]) is derived from a cross between Illinois sweet corn and a Bolivian line, donor of the sugary enhancer allele *se1* (26, 27). Last, we measured *MATE1* expression in root tips of these maize inbred lines carrying three *MATE1* copies in comparison with lines with only one copy of *MATE1* (Fig. 6B). Both under control and Al stress conditions, lines carrying three copies of *MATE1* (Al237, C100-6, and Il677a) have high *MATE1* expression compared with one-copy lines (L53 and B73). *MATE1* expression levels in C100-6 and Il677a are comparable with those of Al237 under both conditions. These

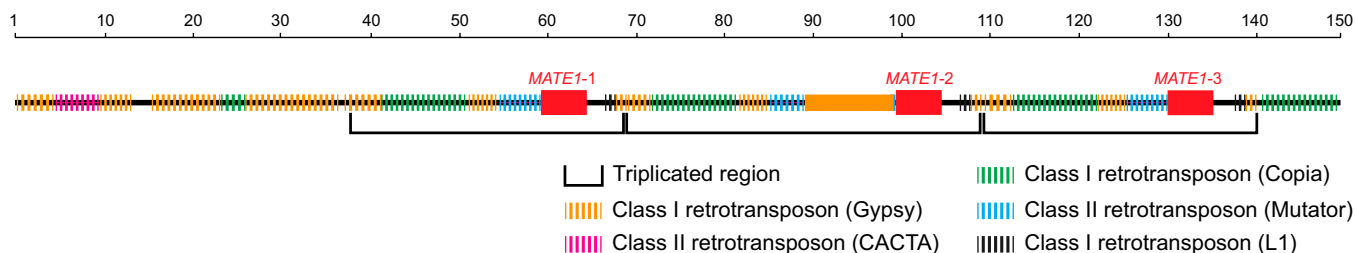


Fig. 5. Genomic organization of the *MATE1* locus shows three tandemly arrayed gene copies. Diagram showing Al237 genomic BAC ZMMCBa0006o22. The triplicated regions are indicated by brackets. The three copies of *MATE1* are indicated by solid red boxes. The dashed boxes of different colors indicate different families of retrotransposons (annotated using <http://maizetdb.org/~maize>). The unique Gypsy retrotransposon insertion upstream of *MATE1-2* is indicated by a solid orange box. Except for *MATE1-1*, -2, and -3, no other genes are predicted within the BAC sequence according to MAKER (45). The scale bar is in kilobase pairs.

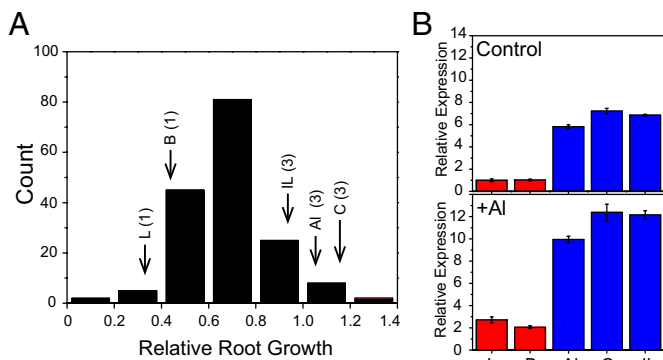


Fig. 6. Maize inbred lines carrying three *MATE1* copies also show high *MATE1* expression. (A) Distribution of Al tolerance (i.e., RRG) in maize diversity panel. The Al tolerance of lines in which *MATE1* expression was quantified are indicated (L = L53; B = B73; IL = IL677a; AI = AI237; C = C100-6). The number in parenthesis indicates *MATE1* copy number. (B) *MATE1* relative expression quantified via qPCR in root tips of maize inbred lines from a survey for *MATE1* CNV. Seedlings were grown under control conditions (Upper), or treated with 39 μM Al^{3+} activity (Lower) for 24 h. The red bars denote lines with one *MATE1* copy (L53 and B73, Al-sensitive), and the blue bars denote three-copy lines (AI237, C100-6, and IL677a, Al-tolerant). Error bars indicate 1 SD.

results are in agreement with the hypothesis that greater *MATE1* copy number leads to higher gene expression levels.

Discussion

In humans, copy number variants encompass more nucleotides and arise more frequently than SNPs, and to a larger extent have been shown to drive human evolution and diversity between individuals (28, 29). Genome structural variation was discovered to also be pervasive in plants (21, 29–31), challenging the notion that a plant genome can be understood through the reference sequence of a single genotype (32). One recent maize study identified thousands of structural variants between the reference genome of B73 and another North American inbred line, Mo17 (31). As the authors point out, the high level of structural variation and differences in genome content observed in maize are unprecedented among higher eukaryotes. In the recently released maize HapMap2 study, structural variation was probed through a global analysis of read-depth variants in over 100 maize lines. Results showed that more than 90% of the maize genome shows some degree of CNV between lines (33). These structural variations are enriched among genes related to stress and stimulus responses, suggesting a role in maize phenotypic diversity and adaptation. Similar associations were observed in soybean, where structural variants are enriched in genomic regions harboring genes involved in plant biotic defense (34).

Despite these results from large-scale studies, examples of the effect of structural variants on plant phenotypes are scarce. Rather than CNVs, instances described to date involve transposable element insertions that modify the gene's level and/or pattern of expression. This is the case for the maize domestication gene *tb1* (35), where a *Hopscotch* insertion in a regulatory region acts as an enhancer of gene expression. Another example involves another Al tolerance gene, *HcAACT1* from barley (36). In this case, a small CACTA-like transposon inserted 5 kb upstream of the ORF both enhances and alters the tissue localization of *HcAACT1* expression. The results presented here strongly suggest that CNV for *MATE1* is the genetic basis for a major Al tolerance QTL mapped in the AI237×L53 population, therefore providing a direct link between CNV and phenotype in maize. In plants, this is a landmark example of a QTL explained by CNV of the gene responsible for the molecular mechanism that is the QTL's mode of action.

The majority of CNVs described in maize are present in multiple inbred lines (33) and are thought to predate domestication, as suggested by their presence in the ancestor of maize, teosinte (29). CNV for *MATE1*, however, is rare among maize lines and possibly occurred after domestication, as it was not observed in teosinte. This suggests that de novo copy number variants are likely to still be arising in the maize genome, and may have significant roles in phenotypic variation and adaptation. Interestingly, only two alleles were found for the *MATE1* locus, and the three-copy allele appears at a very low frequency among maize lines. In addition, sequencing of the BAC clone containing the three copies of *MATE1* from maize line AI237 showed that the 30-kbp repeat regions share high sequence identity with each other. Taken together, these results suggest the triplication at the *MATE1* locus is a very recent event.

It is interesting to note that all three maize lines that carry the three-copy allele share the same geographical origin, in regions of acid soils of the South American tropics. AI237 and C100-6 are Cateto inbreds, the most widespread racial group in South America and the only native Brazilian race used in local maize hybrid breeding programs (37). Cateto constitutes a group of landraces, originally cultivated by the native Indians living in coastal areas from Argentina to the Guianas. Maize lines and hybrids derived from Cateto races have been identified as important sources of Al tolerance since the early 1980s (38, 39). Thus, Al tolerance can be considered an important adaptive trait carried by Cateto, which may have contributed to its overall acceptance, particularly in Brazil. In summary, maize lines carrying the three-copy allele of *MATE1* are all originally from a region where Al tolerance has a strong adaptive value, and where breeding for acid soil tolerance has been intense. The fact that the three-copy allele was fixed in these lines suggests that it may provide a specific advantage for growth on acid soils. Taken together, our data indicate that structural variation in the maize genome has contributed to local adaptation of this crop to the highly acidic soils of the tropics.

Materials and Methods

Plant Growth and Treatment, DNA and RNA Isolations. Maize seeds were germinated and seedlings were grown in full nutrient solution at pH 4.0 in a growth chamber at 26 °C/24 °C (light/dark, 16/8 h), as previously described (40). Free Al^{3+} activities were calculated using the chemical speciation software GEOCHEM-EZ (41). Phenotyping for Al tolerance based on RRG was performed using RootReader2D software (www.plantmineralnutrition.net/rootreader.htm), as described (42). For RNA extraction, root tips (1 cm) were collected and flash-frozen in liquid N_2 , and then stored at -80 °C. Total RNA was isolated using the RNeasy Plant Mini Kit (Qiagen) according to the manufacturer's instructions and quantified spectrophotometrically in a Nanodrop 1000 (Thermo Scientific). Genomic DNA was isolated from young shoot tissue using the DNeasy Plant 96 and Mini kits (Qiagen), and quantified using PicoGreen dsDNA quantitation assay (Life Technologies); fluorescence was read in a DTX880 microplate reader (Beckman Coulter).

Gene Copy Number Determination via qPCR. Primers for qPCR were designed using Primer Express software (Applied Biosystems) with default parameters, or obtained from previously published material. Primer sequences and sources are listed in Table S1. Real-time PCR was performed with Power SYBR Green PCR mastermix (Applied Biosystems) and conducted in a 7900HT Sequence Detection System (Applied Biosystems) using a relative standard curve method. In each run, samples were assayed for both the gene-of-interest (target) and for actin, a gene with known copy number (endogenous control). A set of standards was built from a serial dilution of genomic DNA; standard curves were run for both assays (target and endogenous control), and samples were quantified against the curves using 6 ng of genomic DNA as template. Relative quantities (RQs) were calculated by dividing the target sample quantity by the endogenous control sample quantity, and then normalizing against a calibrator sample (indicated in the text). Therefore, RQ values represent gene copy number relative to the calibrator genotype. Three technical replicates of each sample were averaged per assay (i.e., target and endogenous control). Because RQs are the ratio of two means, SDs were calculated from the coefficients of variation (*cv*, defined as the

ratio between the SD s and the mean \bar{X}). The cv for RQ values is as follows:

$cv_{RQ} = \sqrt{cv_t^2 + cv_{ec}^2}$, where t is target and ec is endogenous control. Once cv_{RQ} is known, then SD is calculated by resolving $cv_{RQ} = s/\bar{X}$ for s .

Gene Expression Analysis via qPCR. First-strand cDNA was synthesized from total RNA using the High Capacity RNA-to-cDNA Kit (Applied Biosystems) according to the manufacturer's instructions. *MATE1* expression was determined using a custom-designed TaqMan assay (forward primer, 5'-CACCCGCTTAGCGTATTCT-3'; reverse primer, 5'-GCACCCGATCCTCATGAT-3'; and probe, 5'-TCTGAATGCGAGCCTCG-3'). A predesigned TaqMan assay for Eukaryotic 18S (Applied Biosystems) was used as the endogenous control. Real-time PCR was conducted in a 7900HT Sequence Detection System (Applied Biosystems). Relative expression was calculated using a relative standard curve method (see above), using standard curves built from a bulked cDNA pool. When determining *MATE1* expression in the RILs, each dataset (i.e., control and Al-treated) was generated independently. The Al-sensitive parent L53 was set as calibrator sample; therefore, RQ values represent expression relative to L53 = 1 within each dataset. Three technical replicates of each sample were averaged per assay; SDs were calculated as described above.

FISH. Procedures are as previously described (43). Details are described in *SI Materials and Methods*.

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GBS of the Al237xL53 Population. GBS was performed as previously described (17). Details are described in *SI Materials and Methods* and Fig. S6.

QTL and eQTL Mapping. The phenotypic index used for mapping Al tolerance QTL was RRG as previously described (15). For mapping *MATE1* expression (eQTL), two phenotypes were used: relative expression under control conditions and relative expression under Al stress. Mean RQ values were obtained as described above. Before single marker analysis, lines and SNP loci with more than 10% of missing data were excluded. Furthermore, markers with the least amount of missing data were selected within 2-Mbp windows, among 1,894 SNPs distributed along 165 Mbp of chromosome 6. Association between SNP markers generated by GBS and phenotypic data were evaluated using a general linear model (GLM) in TASSEL (44), and P values for the F tests were generated.

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