

Note

Key Impact of *Vgt1* on Flowering Time Adaptation in Maize: Evidence From Association Mapping and Ecogeographical Information

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ABSTRACT

An association study conducted on 375 maize inbred lines indicates a strong relationship between *Vgt1* polymorphisms and flowering time, extending former quantitative trait loci (QTL) mapping results. Analysis of allele frequencies in a landrace collection supports a key role of *Vgt1* in maize altitudinal adaptation.

AMONG cereals, maize (*Zea mays* L.) is exceptional in its geographic distribution. Maize originated in tropical central Mexico and was spread during pre-Columbian times to cold temperate regions, with latitudes ranging from ~40°S in Chile up to ~45°N in Canada. This diffusion has been made possible via adaptation of flowering time to local environmental conditions, thus enabling the plant to reach the mature state within a shorter growing season. Linkage analyses have shown that flowering time variation involves numerous chromosomal regions, some of them displaying a major effect and often in diverse populations (CHARDON *et al.* 2004). One of these major quantitative trait loci (QTL), namely *Vegetative to generative transition 1* (*Vgt1*), located in bin 8.05, has been recently cloned using a map-based approach and assigned to a ~2-kb noncoding region that regulates an *APETALA2*-like gene (*ZmRap2.7*) located ~70 kb downstream (SALVI *et al.* 2007). The same study showed that three polymorphisms within *Vgt1* were strongly associated with flowering time across a panel of 95 inbred lines. Reproducibility of association studies has proven to be a challenging issue in human genetics (NEWTON-CHEH and HIRSCHHORN 2005; GORROUCHURN *et al.* 2007) but is not yet well-documented in plants. One exception is the relationship between the *Dwarf8* gene and flowering time in maize addressed by ANDERSEN *et al.* (2005) and CAMUS-KULANDAIVELU *et al.* (2006), follow-

ing the pioneering study by THORNSBERRY *et al.* (2001). These investigations showed that for traits and polymorphisms among highly differentiated genetic groups, sampling and evaluation of population structure are critical issues and can lead to contrasting results.

Because *Vgt1* has been positionally cloned, it offers a unique opportunity to assess association mapping reliability in maize. The objectives of this study were therefore (i) to evaluate the reproducibility of the association between flowering time variation and molecular polymorphisms in the *Vgt1* region using a large set of inbred lines specifically selected to include early materials and (ii) to investigate to what extent *Vgt1* could have contributed to maize adaptation to temperate climates and possibly other adaptative constraints, by investigating a large and geographically dispersed set of open-pollinated varieties from America and Europe.

Association mapping and linkage disequilibrium in the *Vgt1* region: We identified 269 polymorphisms across 10 amplicons encompassing a region of 2.2 Mbp around the *Vgt1* locus. The most significant associations with flowering-time variation evaluated under long days were obtained in *Vgt1* itself: among the 81 tests performed using a mixed model, 67 gave a *P*-value < 1.10⁻² (Figure 1). Three of the four most strongly associated polymorphisms corresponded to the ones highlighted by SALVI *et al.* (2007), including a miniature inverted-repeat transposable element (MITE) insertion. Additionally, a previously uncharacterized 2-bp indel, termed *CGindel587*, displayed the lowest *P*-value (*P* = 2.10⁻¹⁴) and explained 4% of the phenotypic variation not accounted for by

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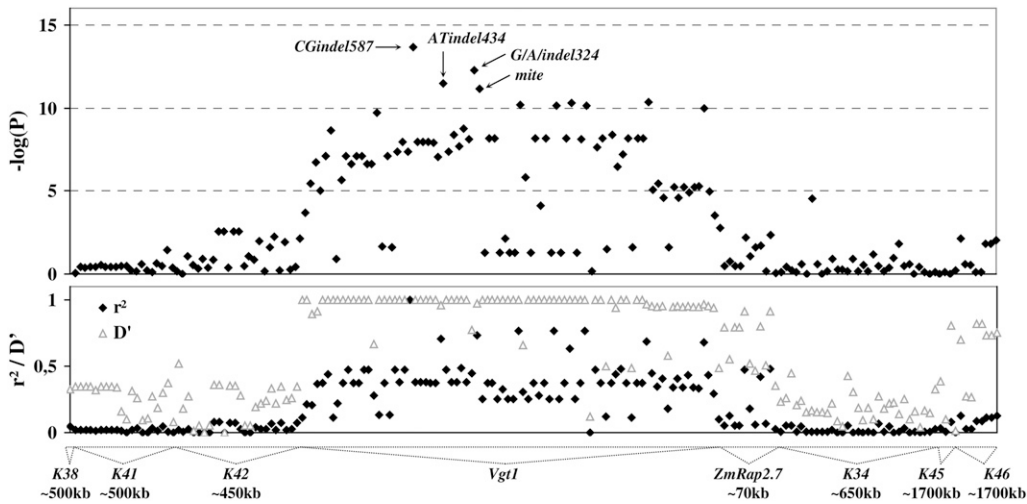


FIGURE 1.—Association of DNA polymorphisms with days to pollen shed (DPS) and extent of linkage disequilibrium in the vicinity of the *Vgt1* region. Statistical significance is expressed as $-\log(P)$; LD is given as both r^2 and D' with *CGindel587*. Approximate positions of amplicons are given relative to *Vgt1* according to the physical map (<http://www.maizesequence.org>). *ATindel434*, *G/A/indel324*, and *mite* correspond to the polymorphisms highlighted by SALVI *et al.* (2007). *CGindel587* has been termed following the same nomen-

clature on the basis of the clustal alignment of N28 and its early derivative C22-4 (SALVI *et al.* 2007). The mapping population was composed of 375 inbred lines representative of the American and European diversity, with a wide range of flowering times, as previously described by CAMUS-KULANDAIVELU *et al.* (2006). Association mapping analyses were performed using TASSEL version 2.0.1 (BRADBURY *et al.* 2007). Polymorphisms with low allele frequency ($<5\%$) were removed, leaving 181 of the 269 polymorphisms for analysis (see supplemental Materials and Methods for further details). Data presented here were obtained using a mixed linear model (MLM) approach accounting for both population structure and relatedness among individuals (YU *et al.* 2006). Estimation of population structure and Loiselle kinship coefficient (LOISELLE *et al.* 1995) was performed using Structure (PRITCHARD *et al.* 2000) and SPAGeDi (HARDY and VEKEMANS 2002), respectively, using 55 tri- to hexa-nucleotide SSRs.

population structure with the mixed model. This percentage increased to 17% when a general linear model (GLM) was applied. To further investigate which of these polymorphisms could underlie the *Vgt1* effect, we performed association tests with *CGindel587*, *ATindel434*, *G/A/indel324*, or *mite* as a cofactor. We observed that *CGindel587* remained significant ($P < 1.10^{-3}$) when one of the other polymorphisms was included in the model, whereas the reverse was not true. Consistently, a haplotype-based analysis combining *mite* and *CGindel587* led to a P -value of 4.10^{-13} , higher than that of *CGindel587* alone. Finally, one can note that including *Dwarf8* in the model did not affect the significance levels and that no interaction was found between *Dwarf8* and *Vgt1*.

Outside *Vgt1* itself, among the 11 polymorphisms considered in *ZmRap2.7*, 4 were significantly associated with flowering time at the $P = 0.05$ threshold (among which 2 were significant at the $P = 0.01$ threshold). Here again, the P -values were no longer significant when *CGindel587* was included in the analysis as a cofactor.

Finally, nine significant associations ($P < 0.01$) were also found in amplicons located several hundreds of kilobases upstream or downstream of the *Vgt1/ZmRap2.7* region; seven of them remained significant at a $P = 0.05$ threshold when *CGindel587* was included in the analysis. Considering the false discovery rate calculated with Q -value software (STOREY 2002) at a 5% level, five of these polymorphisms appeared as true positives, although it cannot be precluded that these significant tests may be due to a residual effect of population structure not fully accounted for by the mixed model we used, as discussed by YU *et al.* (2006). The level of linkage disequilibrium

(LD) was high ($r^2 > 0.7$) over a distance of 1 kb at the *Vgt1* locus, and r^2 values >0.4 were observed between *ZmRap2.7* polymorphisms and *Vgt1* (Figure 1 and supplemental Figure 1). A modeling of LD, following the approach of VEYRIERAS *et al.* (2007), enabled us to distinguish between four ancestral haplotypes at *Vgt1*, which have undergone limited recombination (supplemental Figure 2). The most frequent haplotype combines all the early alleles at loci that display the strongest associations with earliness, including the MITE (absent in other ancestral haplotypes) and *CGindel587*. Another ancestral haplotype is characterized by the late allele at *ATindel434*, *G/A/indel324*, and *mite* and displays segregation at *CGindel587*, with 24 and 17 individuals bearing the early and late allele, respectively.

Congruence with QTL mapping results: Flowering time has been the focus of several linkage studies or is often scored as a secondary trait. We investigated the congruence between QTL and association mapping at the *Vgt1* locus on the basis of eight studies found in the literature (Table 1). Considering the significance levels reported above, we focused on the genotype at *mite* and *CGindel587*.

Vgt1 was identified by VLADUTU *et al.* (1999) in a cross between N28 and its early near-isogenic line, E20 (with Gaspé Flint allele at *Vgt1*). This study highlighted a large effect of *Vgt1* (LOD ≈ 17). Two other studies revealed a QTL of large effect at *Vgt1* in two different crosses: F838 \times F286 (BARRIERE *et al.* 2005) and B73 \times Mo17 (BEAVIS *et al.* 1994). The presence of a QTL at *Vgt1* in these three crosses is consistent with the genotype observed both at *mite* and *CGindel587*, the early parents bearing the early

TABLE 1
Comparison of QTL mapping studies with genotypes observed in parental lines

Study ^a	Cross	Trait ^d	QTL at <i>Vgt1</i> locus	<i>R</i> ² (%) ^e	LOD ^e	Additive effect (days) ^e /late parent		
						<i>mite</i> ^f	<i>CGindel587</i> ^f	
VLADUTU <i>et al.</i> (1999) ^b	N28 × E20	DPS	Detected	NA	~15	2.82/N28	+/-	+/-
BARRIERE <i>et al.</i> (2005)	F838 × F286	SD	Detected	28.8	17.8	2.3/F838	+/-	+/-
BEAVIS <i>et al.</i> (1994)	B73 × Mo17	DPS	Detected	31	11.2	NA/B73	+/-	+/-
BLANC <i>et al.</i> (2006)	DE × F283 × F9005 × F810	SD	Detected	5	NA	0.4/F9005	-/-/-/-	-/-/-/-
AUSTIN and LEE (1996)	H99 × Mo17	SD/DPS	Not detected	—	—	—	-/-	-/-
POUPARD <i>et al.</i> (2001)	F2 × MBS847	SD	Not detected	—	—	—	-/+	-/-
MECHIN <i>et al.</i> (2001)	F2 × MBS847	SD	Not detected	—	—	—	-/+	-/-
BOUCHEZ <i>et al.</i> (2002) ^c	F2 × MBS847 ^e	SD	Not detected	—	—	—	-/+	-/-

^a Other studies reported a QTL for days to flowering in the *Vgt1* region (e.g., ABLER *et al.* 1991; KOESTER *et al.* 1993; TUBEROSA *et al.* 1997; JIANG *et al.* 1999) but, due to incomplete information, they cannot be included in our comparison.

^b Mean value of two environments.

^c In BOUCHEZ *et al.* (2002) the iodent line (Io) was MBS847 (A. CHARCOSSET, personal communication).

^d Days to pollen shed (DPS); silking date (SD).

^e NA, not available.

^f +, late allele; -, early allele, in parental line1/parental line2/...

alleles at these loci. Two studies conducted by BLANC *et al.* (2006) and AUSTIN and LEE (1996) involved parental lines that were identical at both *mite* and *CGindel587*, which is consistent with no QTL of major effect being detected at *Vgt1*. Finally, no QTL has been reported in bin 8.05 in three distinct mapping populations involving F2 and MBS847 (MECHIN *et al.* 2001; POUPARD *et al.* 2001; BOUCHEZ *et al.* 2002). These inbred lines differ at *mite* but share the same allele at *CGindel587*.

Relationship between allele frequencies and geographical origin: Data obtained in the inbred line

panel suggested that *Vgt1* could have been subjected to differential selection, with *mite* and *CGindel587* allele frequencies varying from 0.3 in the late tropical group to 0.87 in the European and Northern Flint groups and with an intermediate frequency of 0.45 in Stiff Stalk and Corn Belt Dent groups. For ease of genotyping and considering its high LD with *CGindel587* when a broad genetic diversity is addressed, we used *mite* as a proxy for *CGindel587* and analyzed its frequency in a 256-landrace collection (Figure 2). This collection exhibits a latitudinal cline for flowering time (supplemental Figure 3)

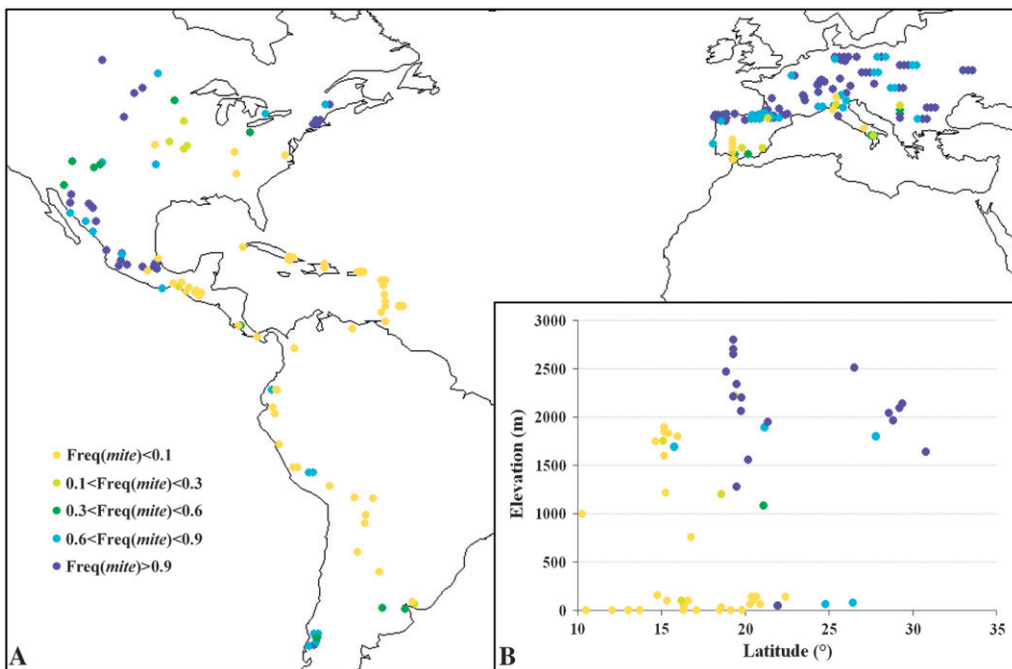


FIGURE 2.—Geographical distribution of *mite* frequency (A) for 256 European and American landraces and (B) for a subset of 77 landraces from Central America and the Caribbean considering both latitude and elevation. The genotypic analysis of landraces was focused on *mite* due to its strong association with flowering time and its high level of LD with *CGindel587*. Moreover it could be easily genotyped by size analysis on a standard agarose gel. Genotyping was performed on a bulk of 15 plants per population as previously described by DUBREUIL *et al.* (2006). Additional information is given in supplemental Materials and Methods.

with days to pollen shed (DPS) under long days, ranging from 612 growing degree days (GDD) for the PPS488 population (latitude 42.9°N) to 1567 GDD for ECUA881 (latitude 3°S).

MITE insertion frequency was generally high (>0.75) in European populations, except those originating from southern Spain and Italy. High frequencies were also observed in Northern Flint and northern Mexican materials. Populations originating from Corn Belt displayed variable frequencies (0–0.8). Caribbean populations exhibited low (<0.25) or null frequency. Populations originating from South America displayed a low frequency except those from southern Chile. We also observed a relationship between elevation and *mite* frequency (correlation coefficient = 0.71, $P < 0.0001$) for populations from Central America and the Caribbean. The frequency of the early allele reached a maximum in landraces collected in elevated sites and declined for landraces originating from lowlands (Figure 2).

Discussion: Our study confirms the association between nucleotidic variation at *Vgt1* and flowering time initially found by SALVI *et al.* (2007). Association tests appear extremely robust for this locus, as illustrated by very high levels of significance observed (down to $P = 2.10^{-14}$ for *CGindel587*) while population structure, strongly related to both polymorphisms and flowering time [50% of the phenotypic variation according to CAMUS-KULANDAIVELU *et al.* (2006)], was accounted for. P -values lower by several orders of magnitude than those observed by SALVI *et al.* (2007) suggest a higher power of the present experiment, due to a larger population size and a more diversified material, including early-flowering lines from Northern Flint and European genetic groups. Furthermore, population structure seems to “absorb” the effect of *Vgt1* in the present panel: *CGindel587* explained 17 and 4% of the phenotypic variation not related to population structure with the GLM and mixed model, respectively, *vs.* 32% for the polymorphisms reported by SALVI *et al.* (2007). Despite difficulties inherent to the estimation of polymorphism effects in such situations, our results suggest that the *Vgt1* early allele shortens the plant cycle by ~100 GDD, *i.e.*, ~7 days under northern France conditions. We can also notice that significance levels at loci other than *CGindel587* were considerably reduced when including *CGindel587* as a cofactor. This indicates that associations found at other positions, including those located in the *ZmRap2.7* gene, are likely the result of linkage disequilibrium with *CGindel587*. It can be noted that $-\log(P)$ for these polymorphisms better correlates with r^2 statistics with *CGindel587* than with D' (0.9 and 0.74, respectively). The region indeed shows a high level of LD, with r^2 values up to 0.4 between *Vgt1* and *ZmRap2.7*, located 70 kb apart. Although infrequent in maize, such a level of LD has also been reported by PALAISA *et al.* (2004) in the region surrounding the *Y1* locus. This

suggests that LD-based genomewide scans at the density of one marker per 10 kb may be relevant in maize to identify regions of the genome with strong contributions to the traits of interest.

Modeling of haplotype structure suggests that four ancestral haplotypes contributed to the variation observed at *Vgt1*. The two ancestral haplotypes bearing the early allele at *CGindel587* display a high similarity in this subregion, suggesting a common origin and the possibility of a recombination event(s) assembling the early allele of *CGindel587* with late alleles at the other strongly associated loci.

The effect of *CGindel587* on early flowering is supported by both a highest significance level and a better consistency with results from QTL analyses. Indeed, our study also illustrates that comparison with QTL studies could be a beneficial way to validate or to pinpoint the weakness of association mapping results. No QTL for flowering time was observed in F2 × MBS847 crosses, although the MITE is inserted in F2 but not in MBS847. As a result, this mite transposon is not expected to be directly responsible for the *Vgt1* effect. The results are more congruent with genotype observed at *CGindel587* since the two lines share the same allele. The ultimate discrimination of the “causal polymorphism” will require further analyses but *CGindel587* appears as a good candidate. It can be noted that this 2-bp indel is neither located in a conserved noncoding sequence (CNS) according to SALVI *et al.* (2007) nor in a known regulatory domain. The underlying molecular mechanisms by which it could impact on flowering time thus needs to be elucidated.

When considering this polymorphism, it is worth noting that the high LD of *CGindel587* with *mite* makes *mite* a reliable proxy to estimate early allele frequency in the inbred line and population panels. These investigations show that early allele frequency at *Vgt1* was highly correlated with geographical origins. Early allele of *Vgt1* displayed a higher frequency in the Northern Flint and European groups. Frequencies were intermediate in the Corn Belt Dent genetic pool and lower in the late tropical pool. This further supports association results, showing that variation at *Vgt1* was selected in adaptation of maize to cool temperate climates. As compared to the *Dwarf8* early allele (CAMUS-KULANDAIVELU *et al.* 2006), the *Vgt1* early allele displays a higher frequency in tropical materials. This suggests that it was preexisting in this genetic pool, as corroborated by the detection of the MITE among a limited set of teosintes (data not shown).

Furthermore, the frequency of *Vgt1* alleles among tropical populations varies with the altitude of the collection site, the early allele being rare at low elevations. This, in addition to an excess of haplotypes and a positive and significant Tajima's D at the *Vgt1* locus in tropical inbred lines (results not shown) supports the hypothesis of diversifying selection following domesti-

cation of maize, with early and late materials adapted to high and low cultivation systems, respectively. Indeed, late materials display an advantage in warmer conditions of lowlands by preventing flowering at a stage where the plant would not have accumulated enough resources to ensure reproductive success. So, in addition to having played a key role in adaptation to a cool temperate climate, *Vgt1* may also have been involved in the differentiation of maize varieties according to elevation in tropical Central America.

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