

# The maize heterotrimeric G-protein $\beta$ subunit controls shoot meristem development and immune responses.

Qingyu Wu<sup>a,b,1</sup>, Fang Xu<sup>a,c,1</sup>, Lei Liu<sup>a</sup>, Si Nian Char<sup>d</sup>, Yezhang Ding<sup>e</sup>, Eric Schmelz<sup>e</sup>, Bing Yang<sup>d,f</sup>, David Jackson<sup>a,2</sup>

<sup>a</sup>Cold Spring Harbor Laboratory, Cold Spring Harbor, NY 11724, USA

<sup>b</sup>Institute of Agricultural Resources and Regional Planning, Chinese Academy of Agricultural Sciences, Beijing 100081, China

<sup>c</sup>School of Life Sciences, Shandong University, Qingdao 266237, China

<sup>d</sup>Division of Plant Sciences, University of Missouri, Columbia, MO 65211, USA

<sup>e</sup>Division of Biological Science, University of California, San Diego, CA 92093, USA

<sup>f</sup>Donald Danforth Plant Science Center, St. Louis, MO 63132, USA

<sup>1</sup>These authors contribute equally to this work.

<sup>2</sup>To whom correspondence may be addressed. Email: [jacksond@cshl.edu](mailto:jacksond@cshl.edu)

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## Abstract

Heterotrimeric G-proteins are important transducers of receptor signaling, functioning in plants with CLAVATA receptors in control of shoot meristem size, and with pathogen associated molecular pattern (PAMP) receptors in basal immunity. However, whether specific members of the heterotrimeric complex potentiate crosstalk between development and defense, and the extent to which these functions are conserved across species, has not been addressed. Here we used CRISPR/Cas9 to knockout the maize  $G\beta$  subunit gene, and found that the mutants were lethal, differing from *Arabidopsis*, where homologous mutants have normal growth and fertility. We show that lethality is not caused by a specific developmental arrest, but by autoimmunity. We used a genetic diversity screen to suppress the lethal  $g\beta$  phenotype, and also identified a new maize  $G\beta$  allele with weak autoimmune responses but strong development phenotypes. Using these tools, we show that  $G\beta$  controls meristem size in maize, acting epistatically with  $G\alpha$ , suggesting that  $G\beta$  and  $G\alpha$  function in a common signaling complex. Furthermore, we used an association study to show that natural variation in  $G\beta$  influences maize kernel row number, an important agronomic trait. Our results demonstrate the dual role of  $G\beta$  in immunity and development in a cereal crop, and suggest that it functions in crosstalk between these competing signaling networks. Therefore, modification of  $G\beta$  has the potential to optimize the tradeoff between growth and defense signaling to improve agronomic production.

## Significance

Cereal crops, such as maize provide our major food and feed. Crop productivity has been significantly improved by selection of favorable architecture and development alleles, however crops are constantly under attack from pathogens, which severely limits yield due to a defense-growth tradeoff. Therefore, it is critical to identify key signaling regulators that control both developmental and immune signaling, to provide basic knowledge to maximize productivity. This work shows that the maize G protein  $\beta$  subunit regulates both meristem development and immune signaling, and suggests that manipulation of this gene has the potential to optimize the tradeoff between yield and disease resistance to improve crop yields.

## Introduction

Shoots are derived from meristems, pool of self-renewing stem cells that initiate new organs from their daughter cells (1). The development of the shoot apical meristem (SAM) is controlled by the CLAVATA (CLV)-WUSCHEL (WUS) feedback signaling pathway (1). This pathway includes a secreted peptide, CLV3, its leucine-rich repeat receptor-like kinase (LRR-RLK) CLV1, and a homeodomain transcription factor WUS, which promotes CLV gene expression and stem cell fate (2-5). CLV1 binds and perceives CLV3 peptide, leading to WUS repression (2-5). A second LRR protein, CLV2, is a receptor-like protein that controls meristem size in parallel to CLV1 (4, 6). The CLV-WUS feedback loop was discovered in the model species *Arabidopsis*, but is conserved widely, including in cereal crops. Through characterizing maize *fasciated ear* (*fea*) mutants with enlarged inflorescence meristems (IMs), the *THICK TASSEL DWARF1* (*TD1*), *FASCIATED EAR2* (*FEA2*) and *ZmCLAVATA3/EMBRYO SURROUNDING REGION-RELATED7* (*ZmCLE7*) genes were identified as orthologs of *CLV1*, *CLV2*, and *CLV3* respectively (7-11). In addition to the conventional CLV1 receptor, the LRR receptor-like protein FASCIATED EAR3 (FEA3) represses WUS from below, and perceives a distinct CLE peptide, *ZmFON2-LIKE CLE PROTEIN1* (*ZmFCP1*) that is expressed in differentiating cells (10). Therefore, distinct CLV receptors perceive small CLE peptides to maintain the balance of meristem proliferation and differentiation. However, the downstream signaling events from these receptors are not well understood.

Heterotrimeric G-proteins, consisting of  $G\alpha$ ,  $G\beta$ , and  $G\gamma$  subunits, transduce signals downstream of receptors (12). In the standard animal model, a GDP-bound  $G\alpha$  associates with a  $G\beta\gamma$  dimer and a 7-pass transmembrane (7-TM) G-protein coupled receptor (GPCR) in its inactive state. Upon ligand perception, the GPCR promotes GDP release and binding of GTP by  $G\alpha$ , activating the G-proteins and promoting interaction with downstream effectors (12). However, G-protein signaling in plants appears to be fundamentally different, and whether plants have 7-TM GPCRs is still under debate (13-15). In contrast, emerging evidence suggests that heterotrimeric G-proteins in plants interact with single-pass transmembrane receptors (16-19). For example, the maize G-protein  $\alpha$  subunit COMPACT PLANT2 (CT2) interacts with the CLV2 ortholog FEA2 to control shoot meristem development, and *ct2* mutants have enlarged SAMs and fasciated ears (16). Similarly, the *Arabidopsis* G-protein  $\beta$  subunit (AGB1) interacts

with another CLV-like receptor, RECEPTOR-LIKE PROTEIN KINASE2 (RPK2) to control Arabidopsis SAM development, and Arabidopsis *agb1* mutant SAMs are larger (16, 18).

In addition to their developmental functions, heterotrimeric G-proteins also positively regulate plant immunity. For example, AGB1 and EXTRA-LARGE GTP BINDING PROTEIN2 (XLG2), a non-canonical  $G\alpha$  in Arabidopsis, interact with the immune receptor FLAGELLIN SENSITIVE2 (FLS2), as well as with its downstream kinase BOTRYTIS-INDUCED KINASE1 (BIK1), which is stabilized by this interaction (19), thus immunity is compromised in *xlg* or  $g\beta$  mutants (19, 20). RNAi suppression of the rice  $G\beta$  gene *RGB1* causes browning of internodes and ectopic cell death in roots, phenotypes associated with immune defects (21, 22). However, the functions of monocot  $G\beta$  genes in development have never been dissected, because CRISPR/Cas9-derived *rgb1* null mutants die soon after germination (23, 24).

Here, we report that CRISPR/Cas9 induced knockouts of maize  $G\beta$  (*ZmGB1*) are seedling lethal, distinct from Arabidopsis but similar to rice. We found that lethality was due to autoimmunity, rather than a developmental arrest. We rescued lethality by introgressing *Zmgb1* CRISPR (*Zmgb1<sup>CR</sup>*) mutants into a suppressive genetic background, and found that the mutants had larger SAMs and fasciated inflorescences. We also identified a viable allele of *ZmGB1* by map-based cloning of a fasciated ear mutant, *fea\*183*, which preferentially alleviated immune phenotypes. Our study dissects the dual functions of  $G\beta$  in shoot meristem development and immune responses, suggesting that modulation of G protein signaling has the potential to optimize the tradeoff between yield and disease resistance in crop plants.

## Results

### **Knocking out *ZmGB1* using CRISPR/Cas9 caused lethality due to autoimmunity**

Maize  $G\alpha$  (CT2) and Arabidopsis  $G\alpha$  and  $\beta$  subunits control meristem development (16, 18, 25). However, the role of  $G\beta$  in meristem regulation in the grasses remains obscure, because rice  $G\beta$  knockouts are lethal, leading to the proposal that it is essential for growth (23, 24). To study the function of maize  $G\beta$ , we used CRISPR/Cas9 to generate multiple alleles, including 1-bp and 136-bp deletions with premature stop codons predicted to result in null alleles (Fig. 1A). Homozygous *Zmgb1<sup>CR</sup>* mutants germinated normally, but arrested and turned yellow then brown, and died at an early seedling stage (Fig. 1B). The necrotic appearance of *Zmgb1<sup>CR</sup>* mutants,

along with the known role of *AGB1* in *Arabidopsis* immune responses (19, 20), prompted us to survey immune markers. We first checked for cell death by staining with trypan blue. *Zmgb1*<sup>CR</sup> mutants were heavily stained compared to wild-type, suggesting that they were undergoing cell death (Fig. 1C). In support of this, 3,3'-diaminobenzidine (DAB) staining showed that H<sub>2</sub>O<sub>2</sub>, another marker for immune responses, accumulated in the mutants (Fig. 1D). We also checked the expression of two immune marker genes, *PATHOGENESIS-RELATED PROTEIN1 (PR1)* and *PR5*, and found that both were significantly higher in *Zmgb1*<sup>CR</sup> mutants (Fig. 1E), as were levels of the defense hormone salicylic acid (Fig. 1F). Similar necrotic phenotypes were found in mutants grown in sterile culture, which together with the up-regulation of immune markers suggests that *Zmgb1*<sup>CR</sup> mutants died because of an autoimmune response.

To confirm that the phenotypes were due to mutation of *ZmGB1*, and not to an off-target effect of CRISPR/Cas9, we made a translational fusion of the *ZmGB1* genomic sequence with YELLOW FLUORESCENT PROTEIN (YFP)-STREPTAVIDIN-BINDING PEPTIDE (SBP) at its N-terminus, under the control of its native promoter and terminator. This construct was transformed into maize and backcrossed twice to *Zmgb1*<sup>CR</sup> heterozygotes in the B73 background. The YFP-SBP-ZmGB1 transgene was able to complement the lethal phenotypes of *Zmgb1*<sup>CR</sup> mutants (SI Appendix Table S1). Imaging revealed YFP-SBP-ZmGB1 localization to the plasma membrane (Fig. 1G), as expected (26), and confirmed by co-localization with FM4-64 after plasmolysis (Fig. 1G). Consistent with its anticipated role in shoot development, *ZmGB1* was expressed throughout the SAM and inflorescence meristems (Fig. 1G).

Having confirmed the *Zmgb1* phenotypes, we asked why the phenotypes of *Arabidopsis Gβ* mutants (reduced immune response, but overall normal growth and fertility) are weaker than in maize. To ask if this was due to the differences in the Gβ protein, or in immune signaling pathways, we expressed maize *ZmGB1* in *Arabidopsis*, driven by the native *AGB1* promoter. The *ZmGB1* transgene fully rescued the developmental and immune defects of *agb1* mutants (SI Appendix Fig. S1), suggesting that Gβ function is conserved between maize and *Arabidopsis*, and that the contrasting immune phenotypes are due to differences in immune signaling pathways.

### ***Zmgb1* lethality can be suppressed**

The early lethality of *Zmgb1<sup>CR</sup>* plants precluded us from observing their meristem phenotypes. Autoimmune phenotypes are common for proteins that are “guardees”, protected (or guarded) by RESISTANCE (R) proteins (27). Since *R* genes are highly polymorphic across accessions, we attempted to suppress *Zmgb1<sup>CR</sup>* autoimmunity by crossing viable heterozygotes to each of the 25 nested association mapping (NAM) maize diversity lines (28), and screened for suppression in the F2s. Indeed, we found that the lethality of *Zmgb1<sup>CR</sup>* could be partially suppressed after crossing to a tropical maize line, CML103. The suppressed *Zmgb1<sup>CR</sup>* mutants were dwarfed with wider stems, similar to the maize *ct2* (*Gα*) mutants (Fig. 2A) (16), and some of the plants survived to flowering (Fig. 2B). Consistent with this growth recovery, the induction of *PR1* and *PR5* immune marker genes was reduced in the suppressed *Zmgb1<sup>CR</sup>* mutants (SI Appendix Fig. S2), confirming that autoimmunity was also suppressed. We took advantage of these lethality-suppressed *Zmgb1<sup>CR</sup>* mutants to study development of their meristems. The mutants had significantly larger SAMs compared to wild-type sibs (Fig. 2C and D) and fasciated IMs (Fig. 2E), indicating that *ZmGB1* controls both SAM and IM development in maize.

### **A newly identified fasciated ear mutant, *fea\*183*, encodes a viable allele of *ZmGB1*.**

Concurrently, we identified a viable recessive allele of *Zmgb1* by map-based cloning of *fea\*183*, a fasciated ear mutant from an ethyl methanesulfonate (EMS)-mutagenesis screen. *fea\*183* mutants were semi-dwarfed, and had shorter, wider leaves with prominent lesions (Fig. 3A). They also had striking inflorescence defects, including fasciated ears and compact tassels (Fig. 3B,C and SI Appendix Fig. S3A), reminiscent of *ct2* mutants (16). We analyzed developing ear and tassel primordia using scanning electron microscopy (SEM), and found their inflorescence meristems were significantly enlarged (Fig. 3B, SI Appendix Fig. S3B). In addition to IM defects, *fea\*183* mutants had larger shoot apical meristems (Fig. 3D and E). The mutants also had obvious cell death and up-regulation of *PR* genes, suggesting an autoimmune phenotype, albeit much weaker than *Zmgb1<sup>CR</sup>* mutants (SI Appendix Fig. S3C and D). Bulked segregant analysis and map-based cloning delineated the *fea\*183* mutation between 257.3MB and 258.9 MB on chromosome 1 (Fig. 3F, SI Appendix Fig. S4A). Whole genome sequencing identified a single non-synonymous mutation within this region, a G-to-A substitution in the fourth exon of *ZmGB1*, leading to a change in the 277<sup>th</sup> amino acid from aspartic acid to asparagine in one of the WD40 domains (SI Appendix Fig. S4B). This residue is fully conserved across a wide range

of species including *S. cerevisiae*, *C. elegans*, *H. sapiens*, and *Arabidopsis*, implying its essential role in G $\beta$  function (SI Appendix Fig. S4C). We next confirmed that *fea\*183* encoded an allele of *ZmGB1*, by crossing with *Zmgb1<sup>CR</sup>* heterozygous plants. In the F1, about half of the plants had enlarged IMs and dwarfism, similarly to *fea\*183* mutants (Fig. 3G and H), indicating a failure to complement, and that *FEA\*183* encodes the maize G protein  $\beta$  subunit. Thus, we renamed *fea\*183* as *Zmgb1<sup>fea\*183</sup>*.

We next asked how the D<sup>277</sup>N mutation affected *Zmgb1<sup>fea\*183</sup>* function, by comparing it with human G $\beta$ , HsGB1, and guided by a structure of the human G protein complex (29). The D<sup>277</sup> residue in *Zmgb1<sup>fea\*183</sup>* aligned to D<sup>254</sup> in HsGB1 (29) (Fig. 4A), which lies at the interface of G $\beta$  and G $\gamma$  (Fig. 4B). We thus asked whether this residue was required to form the heterotrimeric complex, using a yeast-3-hybrid (Y3H) experiment (30). We found that unlike the wild-type protein, the *Zmgb1<sup>fea\*183</sup>* protein could not form a complex with a maize G $\gamma$  subunit (ZmRGG2) and G $\alpha$ /CT2, or with any of the XLG proteins (Fig. 4C), indicating that *Zmgb1<sup>fea\*183</sup>* was unable to form a heterotrimeric complex, and suggesting it resembles a null allele.

### ZmGB1 functions in the CLAVATA pathway

To further decipher the role of ZmGB1, we made double mutants using the *Zmgb1<sup>fea\*183</sup>* allele with other meristem regulatory genes, including *fea2*, *ct2*, and *fea3* (8, 10, 16), and measured meristem size in segregating populations. The SAMs and ear inflorescence meristems of *Zmgb1;fea2* double mutants were not obviously different from those of *fea2* single mutant, indicating that *fea2* is epistatic to *Zmgb1*, and suggesting that they act in a common pathway (Fig. 5A,B,C). Similarly, IMs of *Zmgb1;ct2* double mutants were no more fasciated than either single mutant, suggesting that ZmGB1 and CT2/G $\alpha$  function together in regulating IM development (Fig. 5D). However, vegetative SAMs of *Zmgb1;ct2* double mutants were more severely affected than the single mutant, presumably because CT2 acts redundantly with ZmXLGs during vegetative development (30) (Fig. 5E,F). Finally, *Zmgb1;fea3* double mutants had significantly larger SAMs and more strongly fasciated IMs than either single mutant (Fig. 5G,H,I), indicating an additive genetic effect and that *Zmgb1* and *fea3* act in different pathways in both SAM and IM regulation, in line with previous observations (31). In summary, our data suggest that ZmGB1 functions together with CT2/G $\alpha$  in inflorescence development, downstream of the FEA2 CLAVATA receptor.

### ***ZmGB1* associates with maize kernel row number (KRN)**

KRN is an important agronomic trait that directly contributes to yield (10, 32, 33). Natural or induced variation in *FEA2* or *FEA3* is associated with KRN, and manipulation of *CT2* also enhances KRN (10, 30, 32). We therefore asked if *ZmGB1* also associates with this yield trait by conducting a candidate gene association study using a maize association panel of 368 diverse inbred lines (34). Indeed, we found that five SNPs in the first and third exons of *ZmGB1* significantly associate with maize KRN (Fig. 6A). These five KRN associated SNPs can form four kinds of haplotypes among the 368 lines and two of them (Hap3 and Hap4) have significantly more kernel rows than the other two (Fig. 6B). For example, the Hap4 has 1.5 to 2.5 more kernel rows comparing with Hap2 and Hap1, respectively (Fig. 6B). However, the frequencies of favorable Hap3 and Hap4 in the association panel are only 2.17% and 4.07%, implying that the favorable *ZmGB1* alleles haven't been selected during maize breeding. Therefore, our results suggesting that natural variation in *ZmGB1* influences IM size and KRN, with the potential to benefit maize yields.

## **Discussion**

Heterotrimeric G-proteins are important signal transducers that control many biological processes across a wide range of species (12, 35). They also control many important agronomic traits in cereals (16, 23, 30, 36-39), and understanding G-protein signaling requires a study of each subunit. Rice *G $\beta$*  CRISPR null mutants undergo an early developmental arrest and death, but the underlying mechanism was unclear (23, 24). Here, we show that maize *G $\beta$*  null alleles are also lethal, and this is due to autoimmunity, not to specific developmental defects. We suppressed the *G $\beta$*  lethal phenotype in the CML103 tropical maize genetic background, and identified a viable EMS allele, allowing developmental analysis *G $\beta$*  meristems. Using the suppressed CRISPR null and the viable *Zmgb1<sup>fea\*183</sup>* alleles, we show that *G $\beta$*  controls shoot meristem development. Our results suggest that *G $\beta$*  interacts with different downstream effectors to independently function in immune and development signaling.

An important question is why only monocot *G $\beta$*  mutants, such as in rice or maize, but not *Arabidopsis* mutants, develop autoimmunity? Intriguingly, the *Arabidopsis* *G $\beta$*  mutant *agb1* has

reduced immune response, contrasting with the autoimmune phenotype in rice or maize (19, 20). Expression of maize  $G\beta$  fully complemented the immune defects of *Arabidopsis agb1* mutants (SI Appendix Fig. S1), suggesting that  $G\beta$  protein function is conserved, and that the contrasting phenotypes are due to differences in immune signaling pathways. Plants have a two-tiered immune system. First, Pathogen-Associated Molecular Pattern (PAMP) receptors recognize conserved microbial elicitors and induce Pattern Triggered Immunity (PTI) (40-42). To overcome PTI, pathogens have evolved effectors that they secrete into plant cells to interfere with PAMP signaling, and in turn, plants evolved  $R$  genes to activate the stronger “Effector Triggered Immunity” (ETI), which often results in programmed cell death (43-46). Some  $R$  proteins guard native plant proteins, known as ‘guardees’ that are targeted by pathogen effectors. Thus, mutation of a ‘guardee’ may mimic the presence of a pathogen, and activate the guarding  $R$  protein, resulting in an autoimmune phenotype (47). It is therefore reasonable to speculate that grass  $G\beta$  proteins function as immune guardees. Supporting this hypothesis,  $G\beta$  has 7 WD-40 domains, and forms a propeller structure, similar to some other effector targets (48). Our hypothesis explains why the immune phenotypes of  $Zmgb1^{fea*183}$  mutants are weaker, because presumably that allele accumulates some, albeit mutant,  $G\beta$  protein.  $R$  genes are highly polymorphic across accessions, and our results suggest that  $G\beta$  is guarded in monocots rice and maize, but not *Arabidopsis* (48). To test this hypothesis, further studies are needed to identify the gene(s) that are responsible for the suppression of  $Zmgb1$  lethality in CML103.

Our genetic analyses suggest that ZmGB1 works in a common pathway with FEA2 and CT2/ $G\alpha$ , but independent from FEA3. *fea2* was epistatic to both *ct2* ( $G\alpha$ ) and *Zmgb1* in inflorescence meristem fasciation, suggesting that both G protein subunits function together downstream of the FEA2 receptor. However, *ct2/G\alpha* and *Zmgb1* phenotypes were additive in the SAM, which could be explained by redundancy with the non-canonical  $G\alpha$  proteins, or XLGs, in the SAM (30). However, *Zmxlg* triple mutants are also lethal (30), preventing us from making higher order mutants in maize. Identification of a viable genetic background for  $G\alpha$  higher order mutants would help to address this question.

Geneticists and breeders have used QTL and genome wide association (GWA) analysis to identify genes involved in yield traits. Several yield QTL that correspond to heterotrimeric G proteins or CLV-WUS genes have been cloned in rice, maize, and tomato (33, 37, 49-51). For example, *FEA2* is a QTL responsible for variation in maize kernel row number (KRN) (32), and

one of the rice  $G\gamma$  genes, *GS3* is a QTL for grain length, weight, and thickness (39), while another rice  $G\gamma$  gene, *DEP1*, is a QTL for rice grain yield and nitrogen use efficiency (37, 38). These studies indicate that G proteins and other meristem regulators have the potential to benefit yield traits. In this study, we found that *ZmGB1* also associated significantly with KRN under multiple environments, suggesting that it also contribute to quantitative variation in KRN, and engineering of weak *ZmGB1* alleles may be an effective approach to improve yield traits (10, 32, 52).

Improvement of crop productivity involved selection of favorable architecture and development alleles. Despite these striking innovations, crops are constantly under attack from pathogens. However, turning on defense signaling often causes a reduction in growth and yield (53, 54). This defense-growth tradeoff results from defense signaling being intertwined with physiological networks regulating plant fitness (53). It is therefore critical to understand developmental and immune signaling crosstalk to provide basic knowledge to maximize productivity. Our study shows that *ZmGB1* is a critical regulator in both meristem development and immunity; therefore, this gene has the potential to optimize defense-development tradeoffs to improve agronomic production.

## Materials and Methods

The *Zmgb1<sup>CR</sup>* alleles were created using CRISPR/Cas9, and the *Zmgb1<sup>fea\*183</sup>* allele was from an EMS mutagenesis screen by Gerry Neuffer. Complete details regarding materials, experimental methods, and data analyses are provided in the SI Appendix.

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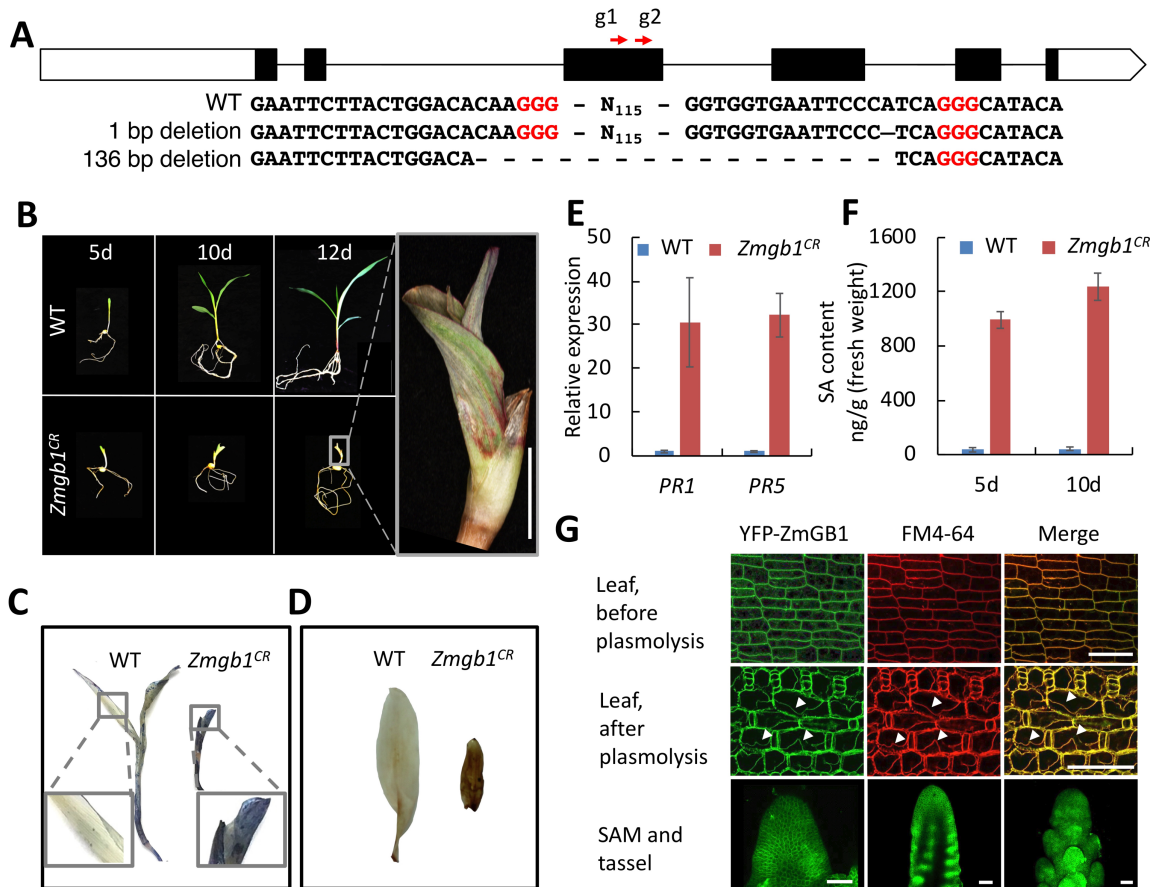
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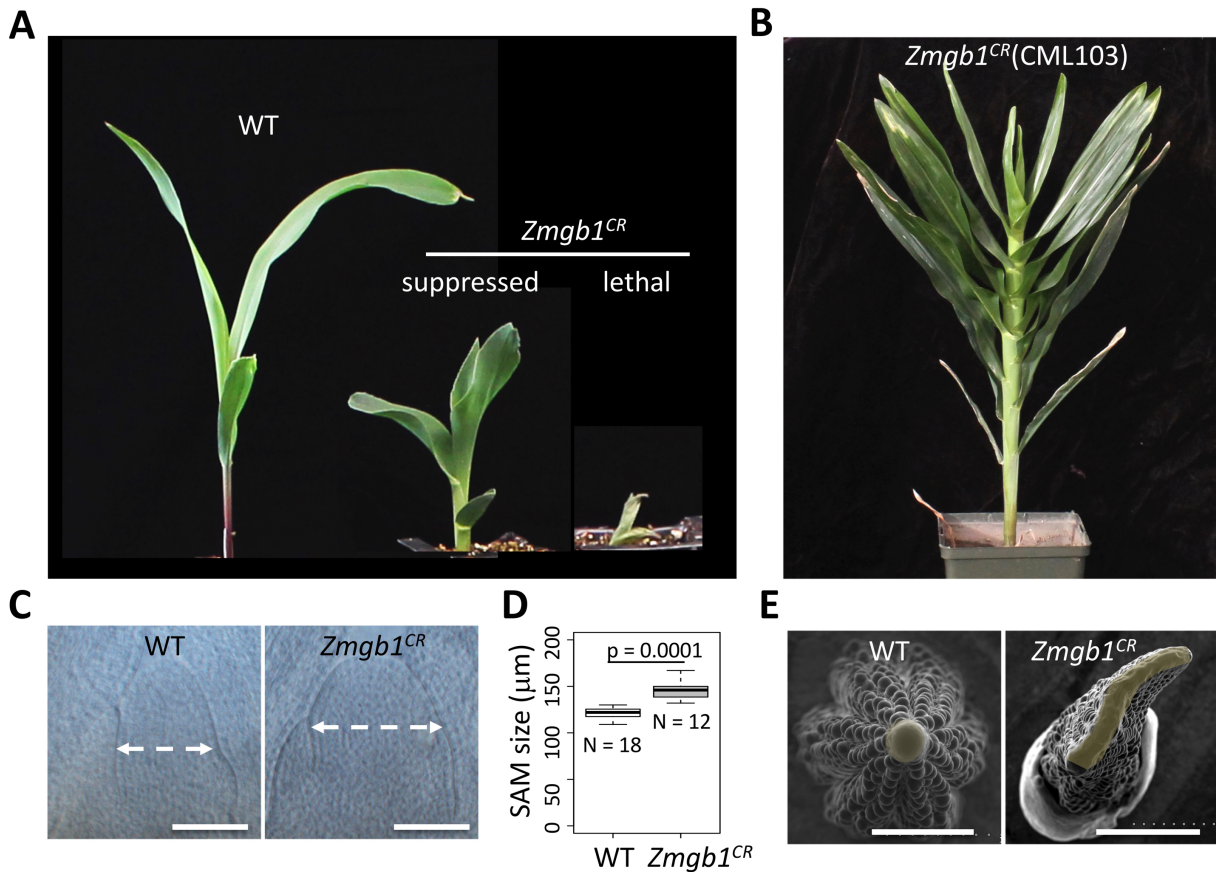
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## Figures

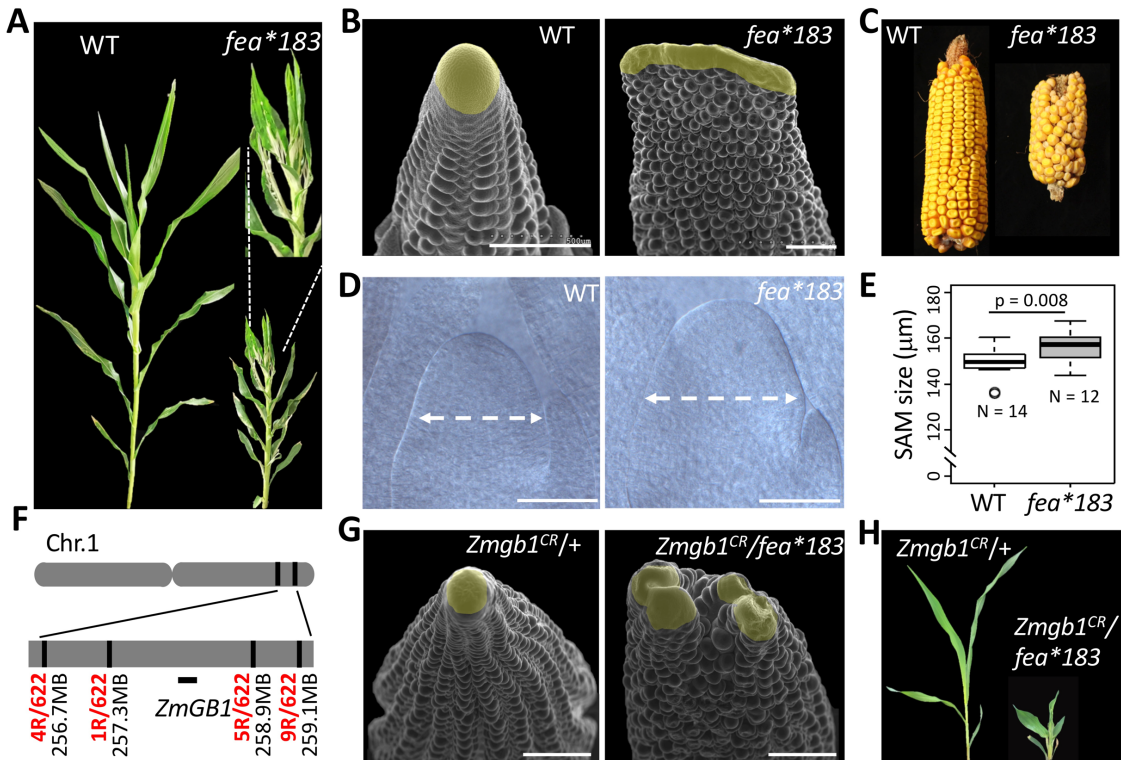


**Figure 1. CRISPR/Cas9 knockouts of *ZmGB1* led to autoimmune phenotypes.** (A) CRISPR/Cas9 editing of *ZmGB1* produced different frameshift alleles. White boxes indicate 5' and 3' UTRs; black boxes indicate exons, and black lines indicate introns. The positions of guide RNAs are indicated by red arrows. (B) *Zmgb1<sup>CR</sup>* mutants were lethal at the seedling stage. The pictures were taken 5, 10, and 12-d post sowing seeds in soil. The upper panels are wild-type (WT) and the lower panels are *Zmgb1<sup>CR</sup>* mutants. Scale bar = 1 cm. Trypan blue (C) and DAB (D) staining of WT and *Zmgb1<sup>CR</sup>* mutants showed increased staining in the mutants. (E) *PR1* and *PR5* expression were up-regulated in the *Zmgb1<sup>CR</sup>* mutants, and both 5-d and 10-d-old *Zmgb1<sup>CR</sup>* mutants accumulated significantly more salicylic acid (SA) (F). For (E and F), *p* value = 0.0001; Student's *t* test; *N* = 3. (G) YFP-SBP-ZmGB1 localizes to membranes in shoot meristems. Upper panels, leaf cells expressing YFP-SBP-ZmGB1 (green), counterstained with FM4-64 (red), both visible as a thin line around the cell. Middle panels, following plasmolysis, YFP-SBP-ZmGB1

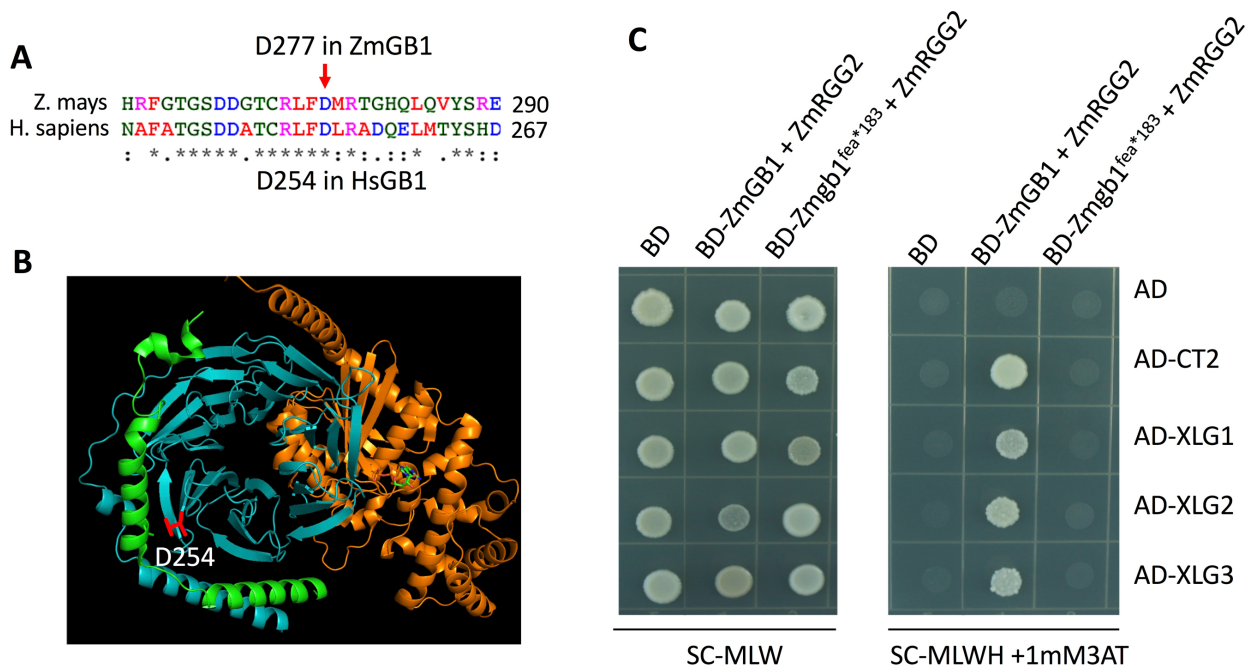
(arrows) remained co-localized with FM 4-64. In the lower panels, YFP-SBP-ZmGB1 expression was found through SAM and tassel inflorescence primordia. Scale bars = 50  $\mu$ m.



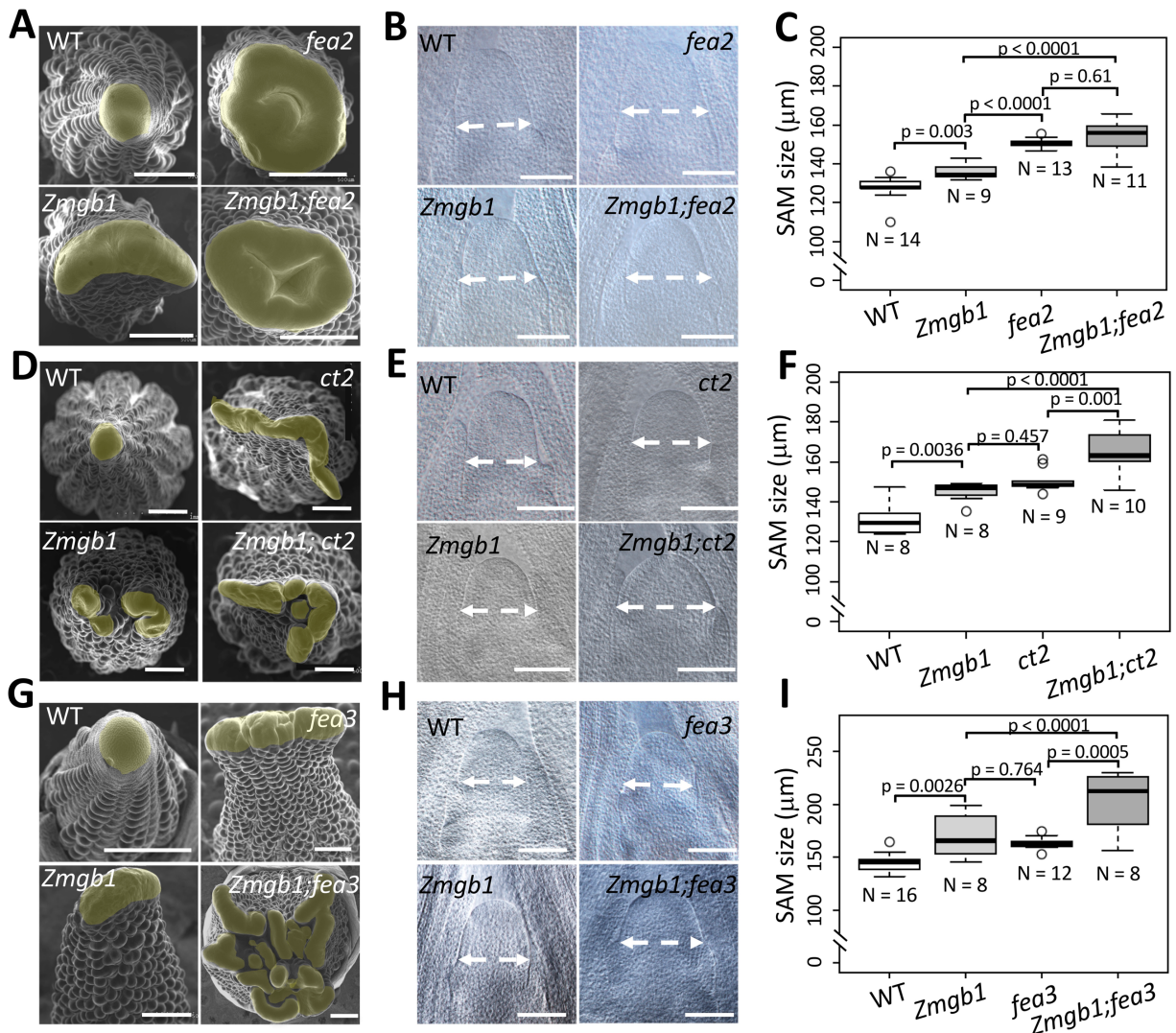
**Figure 2. The lethality of *Zmgb1<sup>CR</sup>* mutants was suppressed in the CML103 background.** (A) F2 progeny of a cross between *Zmgb1<sup>CR</sup>* heterozygotes and CML103 segregates for lethal and suppressed phenotypes. The pictures are of 7-d-old maize seedlings. (B) The suppressed *Zmgb1<sup>CR</sup>* plants in the CML103 background grew to the adult stage. (C) *Zmgb1<sup>CR</sup>* mutants had enlarged SAMs, quantification in (D)  $p$  value = 0.0001; Student's t-test,  $N=18$  for WT and 12 for *Zmgb1<sup>CR</sup>*. (E) Top-down view of WT and *Zmgb1<sup>CR</sup>* ear primordia by scanning electron microscopy; IMs are shaded in yellow. Scale bars = 100  $\mu\text{m}$  for (C) and 1 mm for (E).



**Figure 3. Characterization and mapping of the *fea\*183* mutant.** (A) *fea\*183* plants were semi-dwarfed with upright leaves and lesions (inset dotted lines). (B) SEM images showing that *fea\*183* mutant ear primordia had enlarged IMs, shaded in yellow. Scale bars = 500  $\mu\text{m}$ . (C) Representative mature cobs of WT and *fea\*183* showing fasciated ear phenotype. (D) Cleared SAM images of 12-d-old WT and *fea\*183* mutants. Scale bars = 100  $\mu\text{m}$ . (E) *Zmgb1<sup>CR</sup>* mutants had bigger SAMs;  $p$  value = 0.008, Student's t-test,  $N$  = 14 for WT and 12 for *fea\*183*. (F) Positional cloning of *fea\*183* mutants identified *ZmGB1* as the candidate gene. The vertical lines indicate the position of markers used. The number of recombinants at each position is listed in red. (G) *fea\*183* failed to complement *Zmgb1<sup>CR</sup>* in IM development; SEM images of ear primordia. Scale bars = 500  $\mu\text{m}$ . (H) *fea\*183* failed to complement *Zmgb1<sup>CR</sup>* seedling development, pictures are of 2-week-old seedlings.

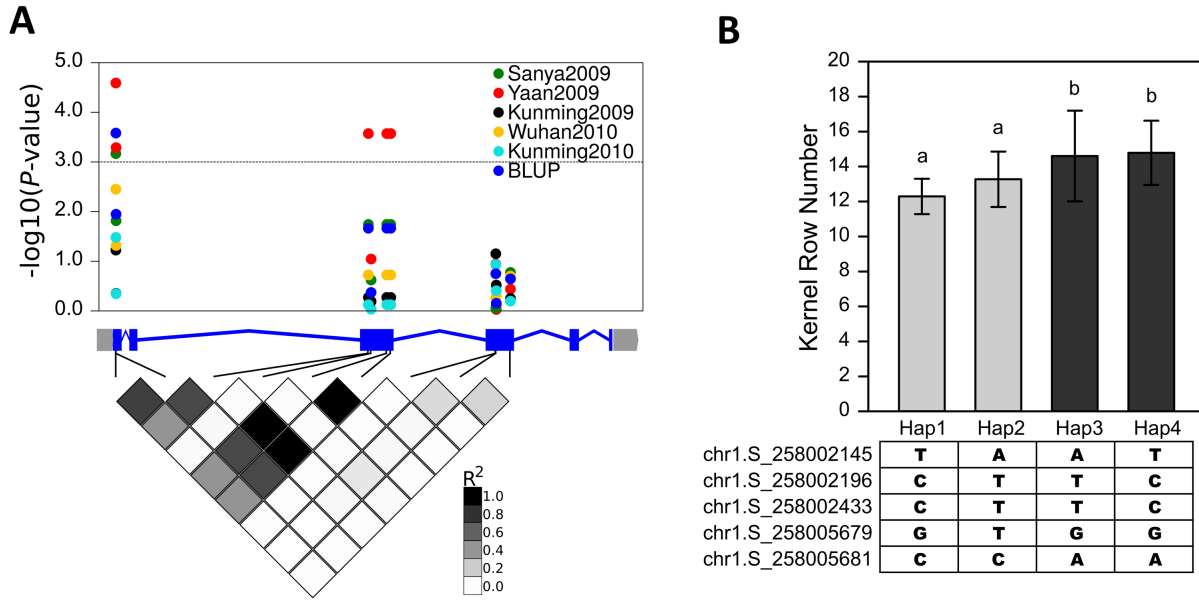


**Figure 4. *Zmgb1<sup>fea\*183</sup>* failed to form a protein complex with G $\alpha$  and G $\gamma$  subunits.** (A) The D277 residue mutated in *Zmgb1<sup>fea\*183</sup>* aligns to D254 in Human HsGB1. (B) D254 highlighted in red in HsGB1 is located at the G $\beta$  and G $\gamma$  interface. Viewed by PyMoL, with G $\alpha$  subunit in orange, G $\beta$  subunit in cyan and G $\gamma$  subunit in green. (C) ZmGB1 and the ZmRGG2 G $\gamma$  subunit formed complexes with G $\alpha$ / CT2 or XLGs in a Y3H assay, while *Zmgb1<sup>fea\*183</sup>* could not. ZmGB1 was fused with the BD domain and co-expressed with RGG2 using a pBridge construct (Clontech). G $\alpha$ / CT2 or individual XLG proteins were fused with the AD domain in the pGADT7 vector. Yeast growth on synthetic complete -Met -Trp -Leu (SC-MLW) medium confirmed transformation and cell viability. Interactions were assayed on SC-Met -Trp -Leu -His (SC-MLWH) medium supplemented with 1 mM 3-AT.



**Figure 5. ZmGB1 functions in a CLAVATA pathway.** (A) SEM images of WT, *Zmgb1*, *fea2* and *Zmgb1;fea2* ear primordia. The double mutants showed similar IMs as the *fea2* single mutant. (B) Representative SAM pictures from 16d-old WT, *Zmgb1*, *fea2* and *Zmgb1;fea2* plants. (C) SAM size quantification showed that the SAM size of the *Zmgb1;fea2* double mutants was indistinguishable from *fea2* single mutants. (D) SEM images of WT, *Zmgb1*, *ct2* and *Zmgb1;ct2* ear primordia. The double mutants showed similar IMs as the single mutants. (E) Representative SAM pictures of WT, *Zmgb1*, *ct2* and *Zmgb1;ct2* plants. (F) The SAM size of the *Zmgb1;ct2* double mutant was significantly increased compared to single mutants. (G) SEM images of WT, *Zmgb1*, *fea3* and *Zmgb1;fea3* ear primordia. The double mutants showed significantly enlarged IMs compared to the single mutants. (H) Representative SAM pictures of 16d-old WT, *Zmgb1*, *fea3* and *Zmgb1;fea3* plants. (I) SAM size of the *Zmgb1;fea3* double mutants was significantly

larger than the single mutants. ANOVA analysis was performed using the R program for (C, F, I). The *p* values, mean values and replicate numbers are indicated in the figures. Scale bar = 500  $\mu$ m for (A, D, G) and = 100  $\mu$ m for (B, E, H).



**Figure 6. The association analysis of *ZmGB1* with kernel row number.** (A) The dots show multiple coding SNPs that associate positively with KRN over multiple environments, and their BLUP (best linear unbiased prediction) data. 368 diverse inbred lines were used in the association analysis using the MLM + Q model. Shaded diamonds below gene model show the SNP linkage disequilibrium (LD) by pairwise  $R^2$ . (B) The haplotype analysis using the five KRN associated SNPs and the KRN (BLUP) of these haplotypes in the association panel, multiple comparisons  $P$ -Value  $< 0.05$ .