

TITLE:

Comparative analyses of exogenous and endogenous antiherbivore elicitors enables a forward genetics approach to identify maize gene candidates mediating sensitivity to Herbivore Associated Molecular Patterns (HAMP)

RUNNING HEAD: Genes mediating maize HAMP sensitivity

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SIGNIFICANCE STATEMENT

Expression analyses and forward genetics approaches identify a maize locus and receptor kinase gene candidate associated with response sensitivity to Fatty Acid-Amide Conjugate (FAC) elicitors widely present in insect oral secretions. Despite broadly overlapping, Herbivore Associated Molecular Pattern (HAMPs) responses can be decoupled from Danger Associated Molecular Pattern (DAMP) responses, providing defined resources to investigate early signaling events underlying plant responses to diverse lepidoptera herbivores.

ABSTRACT (250)

Crop damage by herbivorous insects remains a significant contributor to annual yield reductions. Following attack, maize (*Zea mays*) responds to Herbivore Associated Molecular Patterns (HAMPs) and Damage Associated Molecular Patterns (DAMPs), activating dynamic direct and indirect antiherbivore defense responses. To define underlying signaling processes, comparative analyses between Plant Elicitor Peptide (Pep) DAMPs and fatty acid-amino acid conjugate (FAC) HAMPs were conducted. RNA-seq based early transcriptional changes following Pep and FAC treatments revealed quantitative differences in the strength of response yet a high degree of qualitative similarity, providing evidence for shared signaling pathways. In further comparisons of FAC and Pep responses across diverse maize inbred lines, we identified Mo17 as part of a small subset of lines displaying selective FAC insensitivity. Genetic mapping for FAC sensitivity using the Intermated B73 x Mo17 population identified a single locus on chromosome 4 associated with FAC sensitivity. Pursuit of multiple fine-mapping approaches further narrowed the locus to 19 candidate genes. The top candidate gene identified, termed FAC Sensitivity (*ZmFACS*), encodes a leucine-rich repeat receptor-like kinase (LRR-RLK) that belongs to the same family as a rice receptor gene previously associated with the activation of induced responses to diverse Lepidoptera. Consistent with reduced sensitivity, *ZmFACS* expression was significantly lower in Mo17 as compared to B73. Transient heterologous expression of *ZmFACS* in *Nicotiana benthamiana* resulted in a significantly increased FAC-elicited response. Together, our results provide useful resources for studying early elicitor-induced antiherbivore responses in maize and approaches to discover gene candidates underlying HAMP sensitivity in grain crops.

INTRODUCTION

Crop stress driven by insect pests and disease can cause 50% losses of total annual yield, with increased severity of environmental stresses expected to exacerbate the future losses (Chakraborty and Newton, 2011). Among the more damaging insect pests are lepidoptera in the family Noctuidae which include many *Spodoptera* species (Parra *et al.*, 2021). For example, fall armyworm (FAW; *Spodoptera frugiperda*) is highly polyphagous pest that attacks over 350 host plants across 76 plant families (Montezano *et al.*, 2018). Despite success as a generalist, FAW exhibit measurable specialization on grain crops like maize (*Zea mays*), driving defoliation, seedling loss and introduction of fungal pathogens that contaminate grain with mycotoxins (Overton *et al.*, 2021). Together with native *Heliothis* spp. and *Helicoverpa* spp crop pests in the Americas, FAW has been partially controlled by the transgenic of stacking *Bacillus thuringiensis* (Bt) cytotoxin-encoding genes in many crops (Shehryar *et al.*, 2020). However, despite robust protective Bt crop traits, evidence in Brazil is emerging that FAW is evolving resistance to Bt-mediated crop protection (Horikoshi *et al.*, 2016). Furthermore, FAW is a formidable global invasive pest. In 2016 FAW was detected in West Africa and has now spread across the entire continent causing billions of dollars in annual losses (Day *et al.*, 2017). Entering India in 2018, FAW has rapidly proliferated throughout Asia and is now found in Australia (Overton *et al.*, 2021). Given the global challenge posed by Noctuid pests and potential breakdown of current control measures, new knowledge of plant resistance mechanisms and control strategies are essential to reduce crop losses (Douglas, 2018).

As a major crop species attacked by diverse lepidoptera including FAW and *Spodoptera exigua*, maize has been a leading research model for understanding plant responses to herbivory. For example, in what is now appreciated as a common phenomenon, indirect plant defense responses against lepidopteran pests were first described in maize (Turlings *et al.*, 1990). The elicited production of volatile organic chemicals (VOC) emitted from leaves following *Spodoptera* herbivory can attract parasitoid wasps such as *Cotesia marginiventris*, enhance parasitization rates and aid in plant protection (Turlings and Erb, 2018; Turlings *et al.*, 1990). Herbivore-elicited volatiles are produced by young maize leaves, and while there is qualitative and quantitative variation among inbred lines, the blend is largely dominated by monoterpenes, homoterpenes, sesquiterpenes, fatty acid-derived green leafy volatiles (GLVs), indole and methyl anthranilate (Degen *et al.*, 2004). In addition to attracting parasitoids, indole and GLV components of the volatile blend act in interplant communication, priming defense responses in undamaged neighboring plants (Engelberth *et al.*, 2004; Erb *et al.*, 2015). Volatiles also mediate indirect maize defenses belowground; herbivory by Western corn rootworm (WCR; *Diabrotica virgifera*) larvae triggers

emission of the sesquiterpene E- β -caryophyllene, attracting entomophagous nematodes that prey on the larvae (Rasmann *et al.*, 2005). In addition to volatiles, maize produces complex blends of directly protective chemicals and defensive proteins that vary with tissue, developmental stage and genetic backgrounds. Toxic and antifeedant chemicals include the insecticidal silk-localized C-glycosyl flavone maysin (Waiss *et al.*, 1979; Casas *et al.*, 2016), benzoxazinoid toxins such as dihydroxy-7-methoxy-1,4-benzoxazin-3-one glucoside (DIMBOA-Glc), 2-hydroxy-4,7-dimethoxy-1,4-benzoxazin-3-one glucoside (HDMBOA-Glc) (Oikawa *et al.*, 2004; Maag *et al.*, 2016; Wouters *et al.*, 2016) and antifeedant diterpenoids (Schmelz *et al.*, 2011). While some maize defenses such as DIMBOA-Glc are constitutively present in young seedlings, the production of many defenses is upregulated by herbivory, enabling added protection against attack while minimizing production of costly defenses in the absence of herbivory (Erb, 2018; Fürstenberg-Hägg *et al.*, 2013; Mithöfer and Boland, 2012). Inducible responses can vary across genotypes and environments; however, the presence or absence of antiherbivore defenses is a major factor determining insect resistance (Chen *et al.*, 2009; Smith *et al.*, 2012).

Maize has also been a model for identification of molecules from insects that trigger protective responses. Maize bioassays for elicited VOC production using activity-guided fractionation efforts enabled discoveries of the first defined biomolecules from insect oral secretions (OS) that act as defense elicitors, termed herbivore-associated molecular patterns (HAMPs) (Alborn *et al.*, 1997; Felton and Tumlinson, 2008). Originally isolated from *Spodoptera exigua*, fatty acid-amino acid conjugates (FACs) are a family of molecules based on the conjugation of plant derived linolenic acid to either glutamine or glutamate in the insect gut (Alborn *et al.*, 1997; Pare *et al.*, 1998; Yoshinaga *et al.*, 2008; Lait *et al.*, 2003; Halitschke *et al.*, 2001). Among the naturally occurring FACs, 17-hydroxy *N*-linolenoyl L-glutamine (volicitin) and *N*-linolenoyl L-glutamine (Gln-18:3) are commonly the most highly abundant and potent elicitors of foliar volatile emissions (Mori and Yoshinaga, 2011; Schmelz *et al.*, 2009; Turlings *et al.*, 2000; Yoshinaga *et al.*, 2010; Yoshinaga *et al.*, 2008). FACs occur in diverse insects and play a nutritional role by increasing nitrogen assimilation efficiency in midgut tissues (Mori and Yoshinaga, 2011; Yoshinaga *et al.*, 2008; Yoshinaga *et al.*, 2007). FACs are potent defense elicitors in diverse plants, including maize, rice, soybean, *Medicago* and many solanaceous species (Grissett *et al.*, 2020; Turlings *et al.*, 2000; Shinya *et al.*, 2016; Wu and Baldwin, 2009; Gomez *et al.*, 2005; Schmelz *et al.*, 2009). Although FACs have been the dominant HAMP studied in maize, additional insect-associated molecules also promote maize defenses. Disulfoxy fatty acids, termed caeliferins, present in the American bird grasshopper (*Schistocerca americana*) and microbes associated with the insect digestive tract elicit antiherbivore defense responses in maize, while

chitinases in *Helicoverpa zea* frass suppress antiherbivore defense responses (Ray *et al.*, 2015; J., Wang *et al.*, 2018; Ray *et al.*, 2016; Alborn *et al.*, 2007). Finally, plant hormones contained in both larval oral secretions and frass modulate maize immunity as well (Dafoe *et al.*, 2013; Acevedo *et al.*, 2019). While these diverse insect-associated molecules contribute to elicitation of maize defenses, maize is insensitive to numerous other oral cues from chewing insects such as Lepidopteran-produced inceptin, *Helicoverpa*-associated glucose oxidase and ATPases, and β -glucosidase from *Pieris brassicae* (Schmelz *et al.*, 2007; J., Wang *et al.*, 2018; Mattiacci *et al.*, 1995; Tian *et al.*, 2012).

Maize signaling promoted by HAMPs is mediated and amplified by an array of endogenous signals (Schmelz *et al.*, 2003; Huffaker *et al.*, 2013; Poretsky *et al.*, 2020; Schmelz, 2015). Generally, mechanical wounding of plant tissue leads to the release of damage-associated molecular patterns (DAMPs) including oligogalacturonic acid, extracellular ATP, and peptides similar to systemin and Plant Elicitor Peptides (Peps) (Huffaker *et al.*, 2006; Orozco-Cardenas and Ryan, 1999; Pearce *et al.*, 1991; Tanaka *et al.*, 2014). In addition to HAMPs, DAMPs further amplify wounding-mediated production of phytohormones including jasmonates (JA) and ethylene (ET) to regulate herbivore-associated defense responses (Diezel *et al.*, 2009; Erb *et al.*, 2012; Schmelz *et al.*, 2003; Shinya *et al.*, 2018). Additionally, rapid signaling cascades involving glutamate receptor-like (GLR) proteins, MAP kinase (MAPK) cascades, Ca^{2+} influxes and bursts of reactive oxygen species (ROS) collectively contribute to the propagation of immune signaling both spatially and temporally (Erb and Reymond, 2019). As amplifiers of maize immune signaling, maize encodes 13 ZmPeps contained in 6 precursor protein genes (Huffaker *et al.*, 2011; Huffaker *et al.*, 2013; Poretsky *et al.*, 2020). Peps in rice (OsPeps) have also been demonstrated to protect against herbivores, and act synergistically with HAMPs to generate stronger responses (Shinya *et al.*, 2018). While the potency of ZmPep family members vary in the magnitude of elicited responses, the promotion of JA and ET production, VOC emission, and accumulation of transcripts encoding proteinase inhibitors and other defense proteins is shared (Poretsky *et al.*, 2020). Within the ZmPep family, ZmPep3 is the most potent DAMP signal (Huffaker *et al.*, 2013; Poretsky *et al.*, 2020). ZmPeps are recognized by 2 leucine-rich repeat (LRR) receptor like kinases (RLK), ZmPEPR1 and ZmPEPR2, and plants with *Zmpepr* mutations produce fewer volatiles and are less capable of generating a protective response against *Spodoptera* larvae after ZmPep treatment (Poretsky *et al.*, 2020). This is consistent with other studies demonstrating that impairments to wound and DAMP signaling commonly result in reduced herbivore resistance, and support functional roles for interconnected signaling pathways (Onkokesung *et al.*, 2010; Orozco-Cardenas *et al.*, 1993; Poretsky *et al.*, 2020; Thaler *et al.*, 2002; L., Wang *et al.*, 2018).

Direct comparisons in maize of exogenous HAMPs and endogenous DAMPs, such as Gln-18:3 and ZmPep3, revealed striking overlap in the elicitation of defenses and protective responses against *Spodoptera* herbivores (Huffaker *et al.*, 2013). Given the highly similar activation of defenses, our current research effort seeks to better understand early maize responses to ZmPep3 and Gln-18:3 and comprehensively assess transcriptome wide overlap and response divergence. Transcriptional profiling demonstrates that while ZmPep3 is a more potent signal, both ZmPep3 and Gln-18:3 promote highly similar reprogramming responses at 2 hours, largely represented by transcripts encoding signaling proteins. We identified 312 maize genes rapidly responding to both HAMP and DAMP signals that globally display over 87% overlap. Towards the goal of uncoupling highly similar HAMP and DAMP responses, characterization ZmPep3 and Gln-18:3 sensitivity across diverse maize lines revealed defined maize inbred lines specifically insensitive to Gln-18:3. In a parallel, unbiased and focused approach to candidate gene discovery, we then used forward genetics to better understand FAC response sensitivity. Association mapping using the Intermated B73 x Mo17 (IBM) Recombinant Inbred Line (RIL) population enabled the identification of a single locus specifically associated with response sensitivity to Gln-18:3, but not ZmPep3. Fine-mapping and characterization of this locus led to identification of an *LRR-RLK* gene, termed FAC Sensitivity (*ZmFACS*), as the top candidate consistent with contributions to Gln-18:3 response sensitivity. Using 2 diverse approaches, our work expands the current knowledge of maize genes involved in early signaling responses to defined HAMPs and DAMPs. Furthermore, we provide a long-sought path to uncoupling linked HAMP and DAMP responses in plants.

RESULTS

Comparison of early HAMP- and DAMP-elicited transcriptional changes in maize

Comparative analyses of canonical defense responses against herbivores in maize and rice have revealed considerable connections between HAMP- and DAMP-elicited responses, consistent with shared signaling pathways (Huffaker *et al.*, 2013; Shinya *et al.*, 2018). To confirm that Gln-18:3 and ZmPep3 elicit antiherbivore defenses in the maize B73 inbred similar to previous observations hybrid sweet-corn (Huffaker *et al.*, 2013), B73 seedlings were treated with water, ZmPep3 and Gln-18:3, and VOC emission was measured 16 hours later. In support of earlier findings, both ZmPep3 and Gln-18:3 treatments result in significantly higher levels of VOC emission compared to water-treated samples (Fig. 1A). At equal concentrations ZmPep3 generates stronger induced volatile production than Gln-18:3 (Fig. 1A). To

comprehensively compare early responses to Peps and FACs, B73 leaves were treated with water, ZmPep3 and Gln-18:3 and RNA-seq-based transcriptomes were generated for the 2-hour time point (Table S1). Analyses of the number of differentially expressed genes (DEGs) as compared to water-treated controls showed a similar pattern to that observed for elicitor-induced VOC emission, namely a greater number of ZmPep3 elicited DEGs (1703) compared to Gln-18:3 elicited DEGs (358) (Fig. 1B, Table S2). While a quantitative difference in ZmPep3 and Gln-18:3 DEGs exists, Euler diagram analyses of the overlap between the combined up- and down-regulated DEGs reveals that Gln-18:3 responses display a high degree of overlap with ZmPep3 responses (Fig. 1C, Table S2). Specifically, 87% of all Gln-18:3 DEGs were also differentially expressed following ZmPep3 treatment. In the context of up-regulated DEGs, Gln-18:3 elicited transcripts displayed 92.4% overlap with ZmPep3 responses (Fig. S1A). Further supporting largely shared processes, none of the upregulated DEGs in either treatment were downregulated in the other treatment, and vice versa (Fig. S1A).

To consider biological processes differentially regulated following ZmPep3 and Gln-18:3 treatment, both the maize Phytozome Gene Ontology (GO) and maize MAPMAN annotations were used (The Gene Ontology Consortium, 2019; Goodstein *et al.*, 2012; Thimm *et al.*, 2004). To visualize large-scale differences in the mean expression values of numerous enriched GO terms, a heatmap representing the mean expression values of the DEGs associated with each term was generated (Fig. 1D, Table S3). Among the GO terms enriched in the upregulated DEGs were terms associated with defense signaling and defense responses, including phytohormone and kinase signaling, specialized metabolism, response to wounding and defense to insects, that had the highest mean expression in the ZmPep3 treated leaves followed by the Gln-18:3 treated leaves (Fig. 1D, Table S3). Among the GO terms enriched in the downregulated DEGs were terms associated with growth and development, including red light signaling, gibberellic acid (GA) signaling and regulation of leaf morphogenesis, that had the lowest mean expression in the ZmPep3 treated leaves followed by Gln-18:3 treatment (Fig. 1D, Table S3). In contrast to GO terms, the maize MAPMAN bin annotation provides more specific gene groups and (Thimm *et al.*, 2004). Enrichment analysis revealed a diverse set of MAPMAN bins associated with ZmPep3- and Gln-18:3-upregulated genes, including Pathogen Associated Molecular Pattern (PAMP)-triggered immunity, MAPKs, indole biosynthesis, Guard cell S-type anion channels (SLAC), PAMP-Induced Secreted Peptide (PIP) and PIP-LIKE (PIPL) signaling L-lectin receptors and transcription factors (TFs) in the myeloblastosis (MYB), WRKY, basic Helix–Loop–Helix (bHLH), Dehydration Responsive Element Binding (DREB) and TIFY families (Table S4). Enriched MAPMAN bins associated with ZmPep3- and Gln-18:3-downregulated genes included

PHYTOCHROME B, cell cycle and abscisic acid (ABA) signaling, biosynthesis of gibberellic acid (GA) and brassinosteroids (BR), and TFs from the MYB, C2H2, bHLH and Trihelix families (Table S4). Of the 51 enriched bins for ZmPep3-upregulated genes, only 13 contained more than 10 DEGs, which were associated with 6 different TF families, L- and G-lectin receptor families, and enzymes with peptidase, oxidoreductases, acyltransferases, glycosyltransferase and phosphotransferase activity (Fig. 1E, Table S4). Additionally, heatmap visualization of the ranked FPKM expression data of all ZmPep3-upregulated genes in these enriched bins showed that the majority of ZmPep3 upregulated DEGs also exhibited higher mean expression following Gln-18:3 treatment compared to the water treatment (Fig. 1E). Similarly, a global comparison of all DEGs using the ranked FPKM expression data showed that for over 95% of ZmPep3 DEGs displayed an intermediate Gln-18:3 ranked mean expression, between ZmPep3 and water, despite a nearly 5-fold greater number of ZmPep3 DEGs compared to Gln-18:3 (Table S5). Overall trends in early transcriptional responses following ZmPep3 and Gln-18:3 treatment support highly similar regulation.

Identification of differentially expressed genes after 2 hours of HAMP and DAMP treatment

A comparative analysis of the ZmPep3 and Gln-18:3 DEGs was used to probe transcriptional changes in gene groups and pathways with putative roles in regulation of antiherbivore defenses. MAPMAN pathway annotation and gene descriptions were used to group genes with shared functions (Thimm *et al.*, 2004). Groups included genes involved in signal transduction across membranes, MAPK signaling, phytohormone biosynthesis and signaling, transcription factors, cytoplasmic signaling, and antiherbivore defenses (Fig. 2, Table S6). After identifying the ZmPep3 DEGs in selected groups, a heatmap was generated illustrating the fold change values of the DEGs and including the number of DEGs compared to the total number of genes in each group (Fig. 2, Table S6). Genes that were also significantly differentially regulated following Gln-18:3 treatment were marked using thatched, cyan-colored boxes (Fig. 2, Table S6). Together this assembled landscape of signaling-related genes includes leucine-rich repeat receptor-like kinases (LRR-RLKs) and LRR-receptor-like proteins (LRR-RLPs), lectin receptors, glutamate-like receptors (GLR), respiratory burst oxidase homologs (RBOH), calcium-dependent protein kinases (CDPK), receptor-like cytoplasmic kinases (RLCK), genes involved in ubiquitylation, MAPK-associated genes and phytohormone biosynthetic genes including JA, ET and other pathways. Among the genes involved in signaling, the lectin receptors had the highest proportion of DEGs compared to the group size (30%) while the LRR-RLK/RLP and ubiquitylation groups had among the highest proportion of DEGs split between the number of upregulated and downregulated DEGs (50% and 30% were upregulated, respectively). For bins involved in phytohormone biosynthesis and signaling, genes associated with JA and ET bins were upregulated,

whereas the majority of those in GA, auxin, cytokinin (CK) and ABA groups were downregulated. A large proportion of WRKY and TIFY TF groups were upregulated compared to the size of the group (32% and 52%, respectively), while the DEGs in the bHLH and MYB TF groups were split between upregulated and downregulated genes (61% and 76% were upregulated, respectively). Groups of genes with either established or putative roles in antiherbivore defenses include genes involved in indole, terpenoid, flavonoid, and coumaroyl-CoA biosynthetic pathways, as well as trypsin inhibitors, glucosidases, peroxidases and proteases. Among defense-related genes, groups associated with indole and coumaroyl-CoA biosynthetic pathways had among the highest proportion of ZmPep3 DEGs compared to the size of the group (62% and 30%, respectively). The TIFY TF group and the coumaroyl-CoA biosynthetic pathway group displayed high proportions of genes that were also upregulated by Gln-18:3 in comparison to ZmPep3 DEGs (61% and 62%, respectively) (Fig. 2).

Assessment of genetic variation in sensitivity to ZmPep3 and Gln-18:3

Large-scale overlap in early transcriptional regulation following ZmPep3 and Gln-18:3 treatments supports the hypothesis that defined HAMPs and DAMPs activate maize antiherbivore defenses through highly similar signaling pathways. Previous work on herbivore-induced VOCs revealed significant variation among genetically diverse maize lines (Degen *et al.*, 2004). To narrow the number of candidate genes that have the potential to uncouple ZmPep3 and Gln-18:3 responses, maize inbred lines with reference genomes were analyzed for differential responses to the two elicitors. We hypothesized that the observed response phenotype variation associated with herbivore-induced VOCs could be due to genetic impairments in either HAMP responsiveness, DAMP responsiveness or shared downstream signaling components. To identify genetic variability in either Pep- or FAC-elicited responses 27 maize inbred lines were screened for ZmPep3- and Gln-18:3-elicited volatile sesquiterpene production. This set included B73, Mo17 and W22 inbred lines, and parent lines of the Nested Association Mapping (NAM) population, which collectively represent over 90% of maize genetic diversity (McMullen *et al.*, 2009). CML52 could not be included due to insufficient seed germination. Observed B73 responses were typical of most examined inbred lines and emitted significantly more sesquiterpene volatiles after both ZmPep3 and Gln-18:3 treatments compared to water-treatments alone (Fig. 3A). Nineteen of the 27 lines responded similarly with increased sesquiterpene emissions to both treatments (Fig. 3A). Four inbred lines, namely HP301, CML333, MS71 and Ky21, failed to emit statistically significantly sesquiterpene volatile production in response to either ZmPep3 or Gln-18:3, suggesting the existence of potential mutations in shared signaling pathways (Fig. 3B). Importantly, four inbred lines, namely CML103, NC350, CML69 and Mo17, emitted

significant sesquiterpene volatiles after ZmPep3 treatment but produced no response after Gln-18:3 treatment (Fig. 3C). The final response phenotype (Fig. 3C) is consistent with selective FAC insensitivity and the existence of a genetic basis that underlies signaling nodes able to uncouple FAC- and Pep-elicited responses.

Differential responses in B73 and Mo17 enable forward genetics and the establishment of an FAC sensitivity-associated locus

As an inbred line specifically insensitive to Gln-18:3, Mo17 was selected for further study due to the availability of established genetic resources with recombinant inbred lines (RILs), Near Isogenic Lines (NILs) and high-density genotypic markers (Eichten *et al.*, 2011; Lee *et al.*, 2002; Romay *et al.*, 2013). To confirm that Mo17 is specifically Gln-18:3 insensitive, total VOCs were measured after treatment with water, ZmPep3 and Gln-18:3. As observed in the prior experiment, VOC emission was significantly increased after treatment with ZmPep3 in both B73 and Mo17, but only B73 emitted significantly more VOC after Gln-18:3 treatment (Fig. 4A). Because elicitor-induced VOC emission is a relatively late response measured 16 hours after treatment, analyzed differences in Mo17 responses occur long after application of the signals. To assess whether Mo17 displays insensitivity to Gln-18:3 during earlier signaling events, both B73 and Mo17 were treated for two hours with water, ZmPep3 and Gln-18:3 and analyzed for ET emission. Consistent with VOC emission, B73 emitted significantly more ET following both ZmPep3 and Gln-18:3 treatments, while Mo17 emitted significantly more ET following ZmPep3 treatment but not following Gln-18:3 (Fig. 1B). Using both early and late markers for signal activation, our results support the hypothesis that Pep responsiveness in Mo17 is decoupled from FAC responsiveness and that genetic variation in FAC-specific signaling components exist.

Based on differential FAC sensitivity between B73 and Mo17, the Intermated B73-Mo17 (IBM) Recombinant Inbred Line (RIL) population (Lee *et al.*, 2002) was used for genetic mapping of FAC sensitivity. Using 222 IBM-RILs, ZmPep3- and Gln-18:3-induced leaf VOCs were analyzed and compared to water-treated control plants. Association mapping was conducted based on the fold-change values of ZmPep3- and Gln-18:3-induced VOC emission as compared to water-treated leaves (Table S7). The single-nucleotide polymorphism (SNP)-based genetic marker map (B73 RefGen_V2) of the IBM-RILs (www.panzea.org, July 2012 Imputed All Zea GBS final build) was used for association mapping (Ding *et al.*, 2020). The general linear model (GLM) procedure in TASSEL 5.0 was used to calculate the statistical significance of SNP associations using the fold-change values of ZmPep3- and Gln-18:3-induced VOCs as

traits (Romay *et al.*, 2013; Bradbury *et al.*, 2007). Association mapping revealed a single locus on chromosome 4 as significantly associated with total VOC emission specifically elicited by Gln-18:3 and not by ZmPep3 (Fig. 4C). The FAC sensitivity locus coordinates are based on adjusted *P* values ($P < 0.05$; Bonferroni correction for multiple comparisons) and was defined as the region between B73 RefGen_V2 SNPs S4_237390439 and S4_238691014 (1.9Mbp), which correspond to the B73 RefGen_V4 region between the genes *Zm00001d053820* and *Zm00001d053932*, containing 81 genes (Fig. 4C, Table S8). Box-plot visualization of the ZmPep3 and Gln-18:3 VOC fold-change data split according to allele identity at the most highly associated SNP (S4_237322925) and confirmed that the FAC sensitivity locus was obtained due to Gln-18:3 sensitivity in the presence of the B73 allele and Gln-18:3 insensitivity in the presence of the Mo17 allele (Fig. 4D). To ensure that the Gln-18:3 specific association mapping result (Fig. 4C) was not driven by variation in a single VOC biosynthetic pathway, we investigated (*Z*)-3-hexenyl acetate, (*E*)-4,8-dimethyl-1,3,7-nonatriene (DMNT), and *E*- β -farnesene as separate mapping traits and obtained identical results consistent with a Mo17 lesion impacting early signal propagation (Fig. S2).

Near Isogenic Lines (NILs) narrow the maize chromosome 4 locus associated with FAC sensitivity

A second independent genetic population derived from B73 and Mo17 was used to provide additional support for the FAC Sensitivity locus. The B73 x Mo17 NILs contain small introgression regions from one of the parent genomes into the other nearly uniform genetic background (Eichten *et al.*, 2011). Two NILs, b050 and b154, were identified as containing an introgression of the Mo17 genome into the B73 background within the FAC sensitivity locus, and were screened for ZmPep3- and Gln-18:3-elicited VOC production. Both NILs emitted significantly more VOCs after treatment with ZmPep3, but only b154, emitted significantly more VOCs after treatment with Gln-18:3 (Fig. 5A). These results further support the IBM-RIL derived FAC sensitivity locus (Fig. 4C) and importantly narrow the consideration of candidate genes by reducing the locus to 1.5 Mbps containing 54 genes based on the B73 RefGen_V4 annotation between 242.2 Mbp and 243.7 Mbp (Fig. 5B, Table S8).

Fine-mapping the FAC sensitivity locus using a newly generated IBM-RIL marker map

IBM-RILs provide high resolution resources to investigate the genetic basis of traits associated with development and disease, in part due to five generations of intermating before selfing to create a high resolution RIL mapping population (Lee *et al.*, 2002; Liu *et al.*, 2020). Given considerable investments in the IBM-RIL population, improved resolution genetic marker maps are possible using genotype-by-sequencing (GBS) approaches to identify additional sites of recombination and associated SNPs from DNA

sequence data (Romay *et al.*, 2013). To expand this direction, we obtained and analyzed the raw RNA-seq data (PRJNA179160) of a subset of 105 of the 302 IBM-RILs from an existing study on the regulation of gene expression in two-week old seedlings (Li *et al.*, 2013). Raw data was aligned to the B73 RefGen_V4 genome and used for variant calling. The resulting SNPs were filtered followed by aggregation of all SNPs to a B73 and Mo17 gene-based marker map containing a total of 10,043 marker genes (Fig. 6A, Table S9). Association analyses were performed using the Gln-18:3-induced VOC fold-change data, similar to the previous approach (Fig. 4C) using a subset of 86 IBM-RIL lines (Table S7) for which the refined marker data was generated. After using the Bonferroni-correction for multiple testing and selecting marker genes with a significant threshold of $P < 0.05$, the FAC-sensitivity locus was fine-mapped to a greatly narrowed locus between 242.2 Mbp and 242.8 Mbp (B73 RefGen_V4) containing only 19 genes (Fig. 6B, Table S8). To assess the accuracy of the gene marker map at the FAC sensitivity locus, the allele counts of all SNPs in the region were aggregated and grouped based on the called genotype. These results indicated that higher allele counts matched the predicted SNP genotype, while counts of the alternative alleles were generally close to zero (Fig. 6C). The SNPs that were called as heterozygous were similarly distributed in both the B73 and Mo17 allele counts (Fig. 6C). To verify the fine-mapping results, 3 IBM RILs for which a recombination event was detected within the fine-mapped region (B73 RefGen_V4: 242.2 Mbp and 242.8 Mbp) were tested for elicited VOC emission following treatment with water, ZmPep3 and Gln-18:3. In agreement with our refined IBM RIL-based fine-mapping results, while all lines had significant ZmPep3-induced VOC, significant Gln-18:3-induced VOC were only observed when the genotype at the fine-mapped region was B73 (Fig. 6D and 6E). Among the 19 predicted genes within narrowed FAC sensitivity locus (Table S8), multiple genes exist that theoretically have the potential to impact signaling and transcriptionally mediated responses. These include genes predicted to encode proteins with DNA- or RNA-binding activity (*Zm00001d053855*, *Zm00001d053860*), ATP- or GTP-binding activity (*Zm00001d053857*, *Zm00001d053858*, *Zm00001d053861*), RNA polymerase-related activity (*Zm00001d053872*, *Zm00001d053874*) and protein kinase superfamily proteins (*Zm00001d053853*, *Zm00001d053876*). Curiously, at the center the 19-gene locus exist 2 genes annotated as leucine-rich repeat receptor-like kinases (LRR-RLKs), namely *Zm00001d053866* and *Zm00001d053867*. The predicted *Zm00001d053866* sequence displays considerable physical overlap with the intact LRR-RLK encoding gene *Zm00001d053867* and thus is consistent with a gene prediction algorithm error resulting in misannotation of *Zm00001d053866*. Given the physical genetic position and common role in early signal transduction events mediated by LRR-RLK family members (Reymond, 2021), we selected the intact *Zm00001d053867*

gene, termed *FAC SENSITIVITY ASSOCIATED* (*ZmFACS*), as the top candidate for further assessment in mediating FAC sensitivity (Fig. 6F, Table S8).

***ZmFACS* belongs to the same gene family as the rice receptor *OsLRR-RLK1*, which positively regulates antiherbivore responses**

Of the gene candidates present in the FAC sensitivity locus, *ZmFACS* exists as a member of a receptor gene family that includes the rice gene *OsLRR-RLK1*, which was a candidate selected from a transcriptional profiling study and characterized as a positive regulator of rice antiherbivore responses to diverse lepidoptera attack and crude FAW OS (Fig. 7A) (Hu *et al.*, 2018; Zhou *et al.*, 2011). *ZmFACS* and *OsLRR-RLK1* belong to the LRR-Xb subfamily of LRR-RLKs which also includes the phytoosulfokine (PSKR1/2) and PSY1 receptors (PSY1R), with PSY1R being the more closely-related Arabidopsis LRR-RLK to *ZmFACS* and *OsLRR-RLK1* (Fig. 7A) (Shiu *et al.*, 2004; Hu *et al.*, 2018). *OsLRR-RLK1* transcript accumulation significantly increases approximately 2-fold in rice seedlings in response to FAW OS (Hu *et al.*, 2018; Zhou *et al.*, 2011). To understand whether *ZmFACS* is differentially expressed in maize upon FAC treatment, leaves of B73 and Mo17 plants were treated with water or Gln-18:3 and harvested after 0.5 and 1 hour. In B73, but not Mo17, *ZmFACS* transcripts displayed a significant two-fold increase in accumulation 1 hour after Gln-18:3 treatment (Fig. 7B). Importantly, basal B73 *ZmFACS* expression levels are approximately 200-fold higher compared to Mo17 *ZmFACS* levels over the analyzed treatments and time points (Fig. 7B). Given the differences in *ZmFACS* expression between the two inbred lines, genome sequences upstream and downstream of *ZmFACS* were compared. Comparison of the *ZmFACS* promoter sequences between B73 and Mo17 revealed large non-aligned regions, insertions of transposable element fragments in the DNA9 and Harbinger families as well as an expansion of hAT transposable element fragments (Fig. 7C)(Kohany *et al.*, 2006). In an effort to predict the existence of amino acid (AA) sequence differences potentially disrupting function, the aligned AA sequences of B73 *ZmFACS* and Mo17 *ZmFACS* were annotated through the identification of the signal peptides, LRRs, island domain, transmembrane domain, ATP binding site and the serine/threonine-protein kinase active site (Fig. S3)(Chen, 2021; Sievers *et al.*, 2011; Casas *et al.*, 2016; Kall *et al.*, 2007; Zhang *et al.*, 2020; Wang *et al.*, 2015; Mitchell *et al.*, 2019). Of the 13 total AA differences identified, none resulted in changes predicted to have large-scale impacts on protein function (Fig. S3). From these analyses, promoter sequence variation in Mo17 *ZmFACS* and dramatically reduced transcript abundance (Fig. 7) have greater potential to represent causal differences than the comparatively modest levels of observed AA variation in the predicted *ZmFACS* proteins (Fig. S3).

Heterologous expression of ZmFACS proteins enhance response sensitivity to Gln-18:3 in tobacco (*Nicotiana benthamiana*)

To empirically examine if ZmFACS can promote FAC-elicited response sensitivity, *Nicotiana benthamiana* leaves were used as a heterologous protein expression system. As shown previously, multiple Solanaceous species, including *N. benthamiana* are naturally sensitive to FACs and activate of antiherbivore defense responses upon FAC treatment (Grissett *et al.*, 2020; Xu *et al.*, 2015). Using ET emission to assess FAC-elicited responses, *N. benthamiana* leaves were infiltrated with *Agrobacterium tumefaciens* carrying constructs for the carboxyl terminal yellow fluorescent protein (YFP)-tagged fusions of ZmFACS (ZmFACS-YFP) and similarly tagged *Arabidopsis thaliana* ELONGATION FACTOR-TU RECEPTOR (AtEFR-YFP) as a control (Zipfel *et al.*, 2006). Two days after infiltration the plants were treated with either water, 1 μ M elf18 or 1 μ M Gln-18:3, and ET emission was measured after 2 h. Treatment of *N. benthamiana* leaves with elf18, triggered a significant increase in ET production following the expression of AtEFR-YFP but not ZmFACS-YFP (Fig. S4A). In contrast, treatment with Gln-18:3 triggered significant increases in ET emission in both AtEFR-YFP and ZmFACS-YFP expressing leaves (Fig. S4A). A positive Gln-18:3 result was anticipated given established FAC elicited responses in *N. benthamiana* (Grissett *et al.*, 2020). In an effort to leverage the experimental advantages of *N. benthamiana* as a rapid transient heterologous expression system, we considered alternative plant growth conditions that included lower light levels. In contrast to plants grown under normal light levels, ZmFACS-YFP expressing plants grown under diminished light intensity selectively triggered significant ET emission following Gln-18:3 treatment (Fig. S4B). We speculate that undescribed native *N. benthamiana* protein(s) mediating FAC sensitivity present under normal light conditions have reduced function in control plants (i.e. AtEFR-YFP expression) grown under diminished light. Protein levels of AtEFR-YFP and ZmFACS-YFP were probed and visualized by western blotting demonstrating successful heterologous expression of both (Fig. S4C). Our results are consistent with ZmFACS-YFP heterologous expression increasing *N. benthamiana* responsiveness to FACs (Fig. S4B). To understand the causal basis of the Gln:18:3-elicited VOC association analysis result, *N. benthamiana* plants grown under diminished light intensity were used to compare Gln-18:3 responsiveness to both B73 ZmFACS and Mo17 ZmFACS following heterologous expression. As a control, elf18 treatments triggered a significant increase in ET emission only in AtEFR-YFP expressing leaves (Fig. 8A). Likewise only treatments with Gln-18:3 elicited significant ET emission in B73 ZmFACS-YFP and Mo17 ZmFACS-YFP expressing leaves (Fig. 8A). Our results are consistent with the hypothesis that ZmFACS from both B73 and Mo17 are functionally capable of increasing FAC response sensitivity in *N. benthamiana* (Fig. 8A). Protein levels of

AtEFR-YFP, B73 ZmFACS-YFP and Mo17 ZmFACS-YFP were probed, visualized by western blotting, and displayed readily detectable levels of heterologous expression (Fig. 8B).

DISCUSSION

Plants respond to a complex series exogenous cues and endogenous signals during herbivory that include HAMPs, wound-induced jasmonates, DAMPs and complex amplification signals which collectively activate regulatory mechanisms promoting protective antiherbivore defenses (Erb and Reymond, 2019). For over 30 years maize has been a model system for the study of induced plant responses elicited by insect herbivory, crude insect OS and biochemically defined HAMPs present in insect OS (Turlings *et al.*, 1990; Turlings *et al.*, 1993; Alborn *et al.*, 1997; Pare *et al.*, 1998). Subsequent efforts have expanded to an array of plant-insect models; however, while diverse FAC family HAMPs are known to commonly activate plant defenses, core components involving FAC mediated signal transduction still remain undefined (Halitschke *et al.*, 2001; Schmelz *et al.*, 2009; Wu and Baldwin, 2009; Bonaventure *et al.*, 2011; Erb and Reymond, 2019). Previous research has revealed that like FACs, ZmPeps potently activate classical direct and indirect antiherbivore defenses in maize (Alborn *et al.*, 1997; Huffaker *et al.*, 2013; Poretsky *et al.*, 2020). The defensive biochemical outputs triggered by ZmPeps and FACs are increasingly well characterized; however, gene candidates for proximal mechanisms by which FAC outputs are activated remain less clear (Truitt *et al.*, 2004; Gilardoni *et al.*, 2011; Schmelz, 2015). Transcriptome analyses following biotic stress and treatments with defined elicitors provide increasingly comprehensive insights into the genetic regulation of plant physiological processes (Heidel and Baldwin, 2004; Gilardoni *et al.*, 2010; Zhou *et al.*, 2011; Tzin *et al.*, 2015; Tzin *et al.*, 2017; Poretsky *et al.*, 2020). In the context of microbial pathogens, characterization of transcriptional reprogramming following treatment with diverse defense elicitors identified a shared core of Arabidopsis immune response genes and highlighted conserved roles for glutamate receptor-like calcium-permeable channels in general stress responses (Bjornson *et al.*, 2021). In rice, transcriptional profiling following infestation with the caterpillar *Chilo suppressalis* identified WRKY and LRR-RLK genes involved in herbivore resistance (Hu *et al.*, 2015; Hu *et al.*, 2018). To define candidate maize regulatory genes and explore the degree of overlap between HAMP and DAMP signaling pathways, we employed both comparative transcriptional profiling and forward genetic analyses to consider genes underlying variation in plant responses to Gln-18:3 and ZmPep3.

Our comparative analyses of early-elicited transcriptional changes revealed a core set of 312 DEG genes common to defined HAMP and DAMP immune responses (Fig. 1, Fig. 2; Table S2). Gene expression changes at 2 hours revealed significant quantitative differences yet a high degree of qualitative similarity between Gln-18:3 and ZmPep3 responses. While ZmPep3 treatment resulted in > 4-fold more DEGs than Gln-18:3, over 92% of positively regulated transcripts elicited by Gln-18:3 were shared by positive ZmPep3 responses (Fig. S1; Table S2). Moreover, 95% of the genes that were differentially regulated by ZmPep3 displayed greater average levels in Gln-18:3-treated samples compared to the water-treated controls (Table S5). Collectively our results support a high degree of overlap between ZmPep3 and Gln-18:3 mediated transcriptional responses in maize. From the shared DEGs, diverse regulators of ZmPep3- and Gln-18:3-induced maize responses have been identified that predominate in categories associated with signal transduction and transcriptional reprogramming (Fig. 2). A core set of 44 transcription factor genes rapidly responded to ZmPep3 and Gln-18:3, including transcription factors enriched in WRKY, TIFY, bHLH, MYB, ERF and DREB families (Table S6, Fig. 1C). While core transcription factors in the bHLH and WRKY families have demonstrated roles during herbivory (Schweizer *et al.*, 2013; Hu *et al.*, 2015; Li *et al.*, 2015), clearly a large array of transcription factors underlay extensive transcriptional changes in defense proteins and specialized metabolites following herbivory (Tzin *et al.*, 2017; Poretsky *et al.*, 2020; Erb and Reymond, 2019). Within transcription factor families, the greatest DAMP-associated DEG enrichment was observed in the *TIFY* family (Fig. 2; Table S6) which have established roles in the regulation of jasmonate signaling (Chung and Howe, 2009). A classical feature of Gln-18:3 and ZmPep3 elicitation in maize is the rapid accumulation of jasmonates that occur 2 hours after treatment (Schmelz *et al.*, 2003; Huffaker *et al.*, 2013). Family enrichment and elevated *TIFY* expression is consistent with strongly activated jasmonate signaling mediated by ZmPep3- and Gln-18:3 (Erb and Reymond, 2019; Poretsky *et al.*, 2020). Additionally, the expression of genes encoding numerous RLKs/RLPs/Lectin receptors, MAPK/MAP2K/MAP3K and calcium-dependent protein kinases (CDPK) were elevated, indicating specific networks likely to participate in phosphorylation cascades controlling response outputs (Fig. 2; Table S6). Transcripts for several maize orthologues of Arabidopsis genes encoding core components of wound and reactive oxygen species (ROS)-mediated signaling pathways also accumulated, for example nicotinamide adenine dinucleotide phosphate (NADPH) oxidase family members termed Respiratory Burst Oxidase Homologs (RBOH)- and glutamate receptor like-encoding (GLR) genes (Torres *et al.*, 2002; Mousavi *et al.*, 2013). Many of these transcriptionally-regulated candidate genes are similar to genes mediating antiherbivore defense signaling in other species (Howe and Jander, 2008; Arimura *et al.*, 2005; Maffei *et al.*, 2007a; Maffei *et al.*, 2007b). Our transcriptomic study pinpoints discrete members within maize gene families for further

characterization as predicted regulators of antiherbivore defense responses. These results also suggest that while ZmPep3 is a more potent elicitor in maize, Gln-18:3 and ZmPep3 share highly similar signaling components for antiherbivore response activation. Given the observed HAMP and DAMP overlap, the 1692 significant DEGs observed following ZmPep3 treatment represents a comprehensive view of early defense related changes at the transcriptome level. While a useful starting point, it would be unreasonable to predict that all genes of interest surrounding HAMP signaling will display significant transcriptional changes following elicitation at any one time point. Many candidate genes could potentially represent new nodes in either HAMP or DAMP signaling yet are not readily captured by transcriptomic data alone (Dressano *et al.*, 2020).

The earliest insights into herbivore, OS and HAMP specific maize responses involved the analyses of elicited VOC production as indirect defenses (Turlings *et al.*, 1990; Turlings *et al.*, 1993; Alborn *et al.*, 1997). When measured over time, foliar sesquiterpenes dominate the late term profiles of herbivore and HAMP-elicited maize VOCs (Turlings *et al.*, 1998; Schmelz *et al.*, 2003). Using both ZmPep3- and Gln-18:3-induced production of volatile sesquiterpenes, we screened for genetic variation in diverse maize inbred lines and found that a majority produced significant responses to both elicitors (Fig. 3). Four inbred lines, namely Ky21, HP301, Ms71 and CML333 did not emit statistically significant increases in sesquiterpenes to either treatment. Our results were generally consistent with a previous study observing low terpene emission in these same lines upon elicitation with the synthetic 6-substituted indanoyl isoleucine conjugate analog of JA-isoleucine (Richter *et al.*, 2016). The low VOC response to three different elicitors is consistent with either impaired defense signaling common to all three, or that elicited protective responses in these lines relies predominantly on nonvolatile defenses. Our study failed to detect a single inbred line responsive to Gln-18:3 yet specifically nonresponsive to ZmPep3. Collectively this supports the possible existence of defined maize inbreds, such as Ky21, with compromised signaling in shared pathways downstream of HAMPs, DAMPs and jasmonates (Fig. 3B) (Richter *et al.*, 2016). This is intriguing as many core signaling pathways are governed by gene family duplications and redundancies that create resiliency to single null mutations. For example, analyses of ZmPep receptor mutants, *Zmpepr1* and *Zmpepr2*, demonstrated that ZmPeps signal through both receptors (Poretsky *et al.*, 2020). Genetic redundancy displayed for both *ZmPROPEP* genes and *ZmPEPR* receptors is consistent with the known role of Peps/PEPRs across diverse plant species as core amplifiers of signaling elicited by multiple inputs (Huffaker *et al.*, 2006; Yamaguchi *et al.*, 2010; Liu *et al.*, 2013; Tintor *et al.*, 2013; Ross *et al.*, 2014; Shinya *et al.*, 2016; Poretsky *et al.*, 2020). In contrast, several maize inbred lines were found to be specifically insensitive to Gln-18:3 as assigned by

the elicited sesquiterpene volatile production assays (Fig. 3C). A reanalysis of Mo17 further demonstrated a selective deficiency to elicited ET production following Gln-18:3 treatment yet robust production after ZmPep3 treatment (Fig. 4B). Given that HAMP-elicited ET emission commonly occurs rapidly after elicitation and precedes significant VOC emission (Schmelz *et al.*, 2007; von Dahl *et al.*, 2007), we classified Mo17 as compromised in an early signaling node specifically acting downstream of Gln-18:3. A parallel forward genetics approach was then focused on to narrow the selection of candidate genes influencing HAMP signaling in maize.

To interpret selective plant responses, we used dual-input association analyses to ensure that the specific insensitivity of Mo17 to Gln-18:3 could be assessed in each IBM-RIL line compared to positive ZmPep3 responses. Using the fold-change in total elicited volatiles as a mapping trait, a single locus on chromosome 4 was identified to be significantly associated with sensitivity to Gln-18:3, but not to ZmPep3 (Fig. 4C). Further association analyses using individual volatile traits derived from different biosynthetic pathways demonstrated that Gln-18:3-induced production mapped to the same locus in each case and supported the hypothesis the underlying genetic difference was likely to effect signaling pathways rather than core VOC biosynthetic capacities (Fig. S2). Following multiple NIL- and RIL-based fine mapping approaches, enabled by comprehensive B73 and Mo17 community resources, we narrowed the FAC sensitivity locus to 19 genes (Fig. 5 and 6). At the physical center of the locus we identified the LRR-RLK signaling candidate gene *ZmFACS* (Fig. 6F). Surprisingly, *ZmFACS* exists in the *PSY1* receptor family which is shared by the rice gene *OsLRR-RLK1*, recently demonstrated to contribute to antiherbivore defenses (Hu *et al.*, 2018) (Fig. 7). Transcript analyses demonstrated that expression of *OsLRR-RLK1* was induced in rice during sustained striped stem-borer (SSB, *Chilo suppressalis*) herbivory as well as following FAW OS application. Gene silencing of *OsLRR-RLK1* resulted in rice plants displaying diminished JA and ET production and decreased MAPK3/6 activation following SSB infestation (Hu *et al.*, 2018). SSB-induced increases in expression of several rice WRKY transcription factor genes were delayed in *OsLRR-RLK1*-silenced plants and reduced production of defensive trypsin inhibitors coincided with decreased SSB resistance. In contrast, upon mechanical wounding alone *OsLRR-RLK1* silenced plants demonstrated no difference from wild type controls in JA/ET production, MAPK activation or defense gene expression (Hu *et al.*, 2018). The recent presented hypothesis is that *OsLRR-RLK1* is specifically involved in regulating these outputs in response to an unknown family of HAMPs which are not present during simple mechanical damage (Hu *et al.*, 2018). In the current effort, we employed multiple forward genetics approaches to identify *ZmFACS* as a gene candidate linked to maize insensitivity to a biochemically defined

HAMP, specifically Gln-18:3 (Pare *et al.*, 1998; Truitt *et al.*, 2004). Given the diverse approaches employed and convergent results obtained, it is likely that the ZmFACS and OsLRR-RLK1 have similar roles in transduction and propagation of specific HAMPs.

Precisely how ZmFACS mediates maize response sensitivity to Gln-18:3 remains to be demonstrated. Heterologous expression of ZmFACS in *N. benthamiana* can enhance elicited ET emission after Gln-18:3 application, supporting a role in promoting FAC responses (Fig. 8). In dicot models, the Arabidopsis homologs of ZmFACS and OsLRR-RLK1, AtPSKR1 and AtPSY1R, have been implicated in regulating pathogen resistance and wound responsiveness through upregulation of JA signaling in addition to their canonical role as the PSK and PSY1 receptors, respectively (Igarashi *et al.*, 2012; Mosher *et al.*, 2013; Mosher and Kemmerling, 2013; Shen and Diener, 2013). Receptors may mediate signaling through direct ligand binding or through activating or repressing pathway functions (Han *et al.*, 2014; Hohmann *et al.*, 2017; Smakowska-Luzan *et al.*, 2018). In a legume model for plant-herbivore interactions, cowpea (*Vigna unguiculata*) was recently leveraged in forward-genetic mapping approaches using the inceptin family HAMP peptides to uncover the LRR-RLP inceptin receptor (INR) (Steinbrenner *et al.*, 2020). Physical interactions between inceptin, INR and members of the Somatic Embryogenesis Receptor Kinase (SERK) family co-receptor associations were supported by labeled-ligand binding assays. Importantly stable heterologous expression of INR in tobacco plants conferred both inceptin-induced responses and enhanced *Spodoptera* resistance (Steinbrenner *et al.*, 2020). Diverse receptors are increasingly implicated in regulating antiherbivore responses in multiple species through a variety of mechanisms. Reverse-genetic approaches were used to partly characterize receptors from tobacco (*Nicotiana attenuata*) and rice (*O. sativa*) that regulate antiherbivore responses. The *N. attenuata* co-receptor NaSERK3/BAK1 was examined due to established roles for Arabidopsis orthologues in flagellin signaling, and when silenced, resulted in reduced wound- and OS-induced JA accumulation, but not reduced MPK activity or herbivore resistance, suggesting that NaBAK1 acts downstream of OS perception (Yang *et al.*, 2011). Because NaSERK1 was not silenced, it is possible that the wound- and OS-induced MPK activation were not reduced due to the presence of redundant SERK co-receptors, and that full activation of induced antiherbivore responses requires both NaSERK1 and NaSERK3 (Yang *et al.*, 2011; Ma *et al.*, 2016). Herbivore Danger Signal (HDS)-ASSOCIATED RLK1 (HAK1) and HAK2 were identified through a sequence-mining approach to identify receptors transcriptionally responsive to fractionated *Spodoptera litura* oral secretions enriched in polysaccharides based on sequence homology to CERK receptors involved in recognition of chitin polysaccharides (Uemura *et al.*, 2020). Arabidopsis *hak1* insertional knockout lines demonstrated reduced

responses to an OS fraction containing polysaccharides, with the receptor-like cytoplasmic kinase (RLP) PBL27 identified as a HAK1-interacting coregulator. Similarly, the *N. attenuata* receptor gene, *LECTIN RECEPTOR KINASE1* (NaLRK1) was among the most highly transcriptionally upregulated genes following FAC treatment (Gilardoni *et al.*, 2010; Zhou *et al.*, 2011). Silencing of *NaLRK1* resulted in reduced accumulation of SA, but not JA, after OS treatment, suggesting a role in regulating a subset of responses downstream of OS perception (Gilardoni *et al.*, 2011).

Similar to most receptors previously identified as candidate regulators of antiherbivore defense responses, the precise molecular mechanism by which ZmFACS mediates sensitivity to Gln-18:3 remains to be determined. While heterologous expression of ZmFACS in *N. benthamiana* enhances response sensitivity to Gln-18:3, additional experiments are required to determine if ZmFACS is necessary or sufficient for FAC sensitivity in maize. ZmFACS knockdown or defined CRISPR/Cas9 *Zmfacs* mutants must be generated in B73 to fully prove that ZmFACS is required for FAC elicited defense responses. Stable heterologous expression of FACS in non-responding plant species could be accomplished to determine if the gene is sufficient to confer sensitivity to Gln-18:3. While FACs have not yet been identified as a component of *Chilo suppressalis* oral secretions, FAC family HAMPs occur in the OS of at least 19 examined lepidoptera species (Yoshinaga *et al.*, 2010). Thus comparative testing of FACs sensitivity in *OsLRR-RLK1* silenced rice lines could be informative. More broadly, the evaluation of comparative defense responses after FAC treatment and range of different elicitors, such as Peps or chitin, could further inform whether *OsLRR-RLK1* might selectively function in FAC signaling. In addition to these studies, experiments assessing whether ZmFACS directly binds Gln-18:3 as a ligand are of interest. Evidence for the existence of an FAC-binding protein was demonstrated using enriched plasma membrane preparations from hybrid maize (var. Delprim) and a [³H]-radiolabeled FAC (Truitt *et al.*, 2004). A critical assessment of whether ZmFACS or *OsLRR-RLK1* functions through direct interaction with FACs will prove challenging. Modern standards of proof are significant and recent critical analyses of the entire field of research states that not a single "bona fide HAMP/PRR pair has yet to be discovered" (Reymond, 2021). As the first challenge only modest radioisotope changes, such as [³H]-FAC, have been reported to retain FAC binding and biological activity. This is broadly consistent with lipid derived ligands that are comparatively intolerant to significant modification (Merkler and Leahy, 2018). The lack of free amide groups in Gln-18:3 make acridinium conjugate labeling similarly unfeasible (Steinbrenner *et al.*, 2020). Biophysical ligand-binding assays could overcome this challenge (Sandoval and Santiago, 2020; Sharma and Russinova, 2018); however, within seconds of contacting the wounded leaf surface FACs undergo rapid lipoxygenase mediated modifications

resulting in active and inactive oxygenated derivatives (VanDoorn *et al.*, 2010). Rapid plant-mediated modification of FACs creates an open question regarding the natural receptor ligands and could complicate the use of *in vitro* binding assays that lack modifying enzymes.

Our transcriptomic analyses of early maize responses to ZmPep3 and Gln-18:3 identified a targeted set of 312 shared transcriptionally co-regulated genes rich in signaling candidates. In an independent forward genetics approach we uncovered the candidate gene ZmFACS associated with maize FAC sensitivity. Most conceptually related studies have relied upon combinations of transcriptional profiling, candidate gene mutation and silencing approaches to highlight genes influencing herbivory signaling (Uemura *et al.*, 2020; Hu *et al.*, 2018, p.1; Hu *et al.*, 2015; Gilardoni *et al.*, 2011; Yang *et al.*, 2011, p.1). Of the diverse published efforts that examine FAC-elicited plant defense responses, the current study is the first utilize both transcriptome changes and unbiased forward genetics approaches to identify a gene candidate mediating FAC-elicited response sensitivity. Elicitor-induced antiherbivore signaling and defense activation have long been examined given the promise to naturally limit arthropod based crop damage, but applications are constrained by limited mechanistic knowledge (Karban and Baldwin, 1997; Kessler and Baldwin, 2002). As a rather unique model family of HAMPs, FACs have been closely examined by diverse research groups for a quarter of century and are established to both widely occur in lepidoptera herbivores and broadly elicit defenses responses in diverse monocot and dicot plants (Yoshinaga *et al.*, 2010; Halitschke *et al.*, 2001; Schmelz *et al.*, 2009; Grissett *et al.*, 2020; Shinya *et al.*, 2016). Lepidoptera pests in the genus *Spodoptera* drove early discoveries of HAMP-induced plant responses mediated by FACs and informed critical concepts of complex yet common plant-insect interactions more broadly (Turlings *et al.*, 1993; Alborn *et al.*, 1997; Turlings and Erb, 2018; Ling *et al.*, 2021). An improved molecular understanding of how HAMP responses mediated by FACs are activated to directly or indirectly suppress *Spodoptera* pests (Alborn *et al.*, 1997) has the continued potential to guide the development of improved crop resistance to herbivores.

MATERIALS AND METHODS

Plant materials and growth conditions

Maize (*Zea mays*) seeds for B73, Mo17, W22 and Nested Association Mapping (NAM) (McMullen *et al.*, 2009) parental line seeds and B73 x Mo17 near isogenic lines (NILs) were obtained from the Maize Genetics Cooperation Stock Center (Urbana, IL, USA). Maize seeds for the Intermated B73 x Mo17 (IBM)-

recombinant inbred lines (RILs) (Lee *et al.*, 2002) were provided by Dr. Peter Balint-Kurti (U.S. Department of Agriculture-Agricultural Research Service [USDA-ARS]). All maize lines used for association analyses are listed (Supplementary Table 7). Maize plants were grown in BM2 soil (Berger Mixes, Saint-Modeste, Quebec, Canada) inside a greenhouse (12-h light, minimum of 300 $\mu\text{mol m}^{-2} \text{s}^{-1}$, and 12-h dark) at 24°C/28°C (night/day) temperature cycle. Maize plants were supplemented with a 18-18-21 Tomato Plant Food fertilizer (Miracle-Gro) and grown for 21-25 days before use of leaves. The maize plants that were used for the combined RNA-seq and VOC analyses were grown inside a growth room (16-h light, at 150 $\mu\text{mol m}^{-2} \text{s}^{-1}$, and 8-h dark cycle) at 22°C. *Nicotiana benthamiana* plants were grown inside a growth room under two different conditions, normal light intensity (150 $\mu\text{mol m}^{-2} \text{s}^{-1}$) and reduced light intensity (80 $\mu\text{mol m}^{-2} \text{s}^{-1}$), as specified. For plants grown under reduced light intensity condition the light intensity for was further reduced (10 $\mu\text{mol m}^{-2} \text{s}^{-1}$) by disconnecting the overhead lights on the shelf in the growth room immediately following *Agrobacterium* infiltration and until elicitor treatment two days later. In both the normal light intensity and the reduced light intensity conditions, a Gro Lite WS lamp (Interlectric Corp.) was included as a supplementary light source and the plants experienced a 16-h light and 8-h dark cycle at 22°C. The plants were planted in BM2 soil and supplemented with a 20-20-20 General Purpose fertilizer (Jack's Professional, JR Peters, Inc., Allentown, PA).

Elicitor treatment of plant leaves

The peptides elicitors ZmPep3 and elf18 were synthesized as 23mers and received as 95% purity (Sigma-Aldrich). The FAC *N*-linolenoyl L-glutamine (Gln-18:3) was synthesized as described (Alborn *et al.*, 2000) and obtained as a gift from Dr. Hans Alborn (USDA-ARS). All elicitors were diluted in water to the concentrations indicated. All maize treatments were performed on leaf 5 of 3-week-old plants. For the elicitation of volatiles, excised leaves were placed into individual 4 mL vials containing 1 mL of the treatment solutions for 16 hours of darkness prior to volatile collections. For RNA-seq analysis whole B73 leaves were excised, put in 4 mL vials containing 1 mL of the treatment solutions for 2 hours, after which 2 inches from the base of the leaves were cut and flash frozen in liquid N₂. For the analysis of transcript abundance by qRT-PCR, leaf tissue was treated by scratch application of 20 μL treatment solutions for the specified times prior to flash freezing tissue in liquid N₂. For measurement of ET production, maize leaves were excised and put in 4 mL vials containing 1 mL of the treatment solutions for 1 hour before being inserted into a sealed 7 mL vial for 1 hour of head space collection as described (Schmelz *et al.*, 2006). Measurement of ET production in *N. benthamiana* leaves was conducted using plants grown under two distinct growth conditions, normal light intensity and diminished light intensity. For the treatments, 2 of

the most recent fully expanded leaves of 4 week old plants grown under normal light intensity or 5 week old plants grown under diminished light intensity were infiltrated with *Agrobacterium tumefaciens* carrying constructs for the indicated YFP-fusion *LRR-RLK* genes two days before treatment. After two days the *N. benthamiana* leaves were infiltrated with the treatment solutions and 3 leaf disks were immediately cut with a 13 mm cork borer and sealed in 7 mL vial for 2 hours before headspace ET collection.

Measurement of plant volatiles

Collection of volatile organic compounds (VOCs) from maize leaves was conducted by enclosing the leaves in glass tubes under light for 30 minutes and collecting head-space volatiles on a 50 mg Porapak Q (80/100 mesh; Sigma Aldrich). VOCs were eluted using methylene chloride, with the addition of nonyl acetate as an internal standard and analyzed by gas chromatography (GC) and flame ionization detection (FID) as previously described (Schmelz *et al.*, 2001). Established *Spodoptera*-elicited maize volatiles (Turlings *et al.*, 1991) including (Z)-3-hexenyl acetate, linalool, (3E)-4,8-dimethyl-1,3,7-nonatriene (DMNT), β -caryophyllene, (E)- α -bergamotene, (E)- β -farnesene and E-nerolidol were identified by comparing their retention times with those of pure standards. ET production was measured by enclosing leaves in a 7 mL tube and collecting head-space volatiles for 2 hours. A 1 mL sample was then analyzed by GC-FID and quantified using an ET standard curve as described (Schmelz *et al.*, 2006).

RNA-seq preparation and transcriptome analyses

Total RNA was isolated with the NucleoSpin RNA plant kit (Clontech), treated with the Turbo DNA-free kit (Ambion) and quantified using Qubit 2.0 fluorometer (Life Technologies). Preparation of RNA-seq data was done by Novogen Corporation. Briefly, insert size was checked on an Agilent 2100 Bioanalyzer (Agilent Technologies), and sequencing of the 250–300-base-pair (bp) insert cDNA library was conducted using Illumina HiSeq platform PE150. RNA-seq raw reads are in the process of being deposited to NCBI Sequence Read Archive (SRA). Filtering of raw reads was done by discarding reads with adaptor contaminations, reads with more than 10% of uncertain nucleotides and reads with more than 50% low quality nucleotides (base quality < 20). TopHap2 was used to align reads to the Maize RefGen_V4 genome with default parameters and mismatch parameter set to 2 (Kim *et al.*, 2013). HTSeq was used to analyze gene expression levels in union mode (Anders *et al.*, 2015). DESeq was used for analysis of differential gene expression followed by calculation of *P*-values using the negative binomial distribution and adjusted using Benjamini–Hochberg procedure for False-Discovery Rate (FDR) (Anders and Huber, 2010). Genes with \log_2 Fold Change (FC) > 1 or < -1 and adjusted *P*-value < 0.05 were considered differentially expressed. GO

term enrichment analysis was conducted with the Phytozome GO term annotation for Maize B73 V4 using the hypergeometric test to identify enriched GO terms with Benjamini–Hochberg FDR adjusted P -value < 0.05 (Goodstein *et al.*, 2012). The Maize B73 V4 MAPMAN bin annotations were used for bin enrichment analysis using the hypergeometric test to identify bins with Benjamini–Hochberg FDR adjusted p -value < 0.05 (Thimm *et al.*, 2004).

Association mapping to identify a maize FAC sensitivity locus

To screen for the genetic variation in maize response sensitivity to FAC elicitation, we examined replicated leaves ($n=4$) of B73, Mo17, W22 and the NAM parent founders for elicited volatile emission following treatments with water, 1 μM Gln-18:3 and 1 μM ZmPep3. ZmPep3 was used as a positive control for plant responsiveness. Mo17 was among 4 maize inbred lines specifically insensitive to Gln-18:3. Based on differential Gln-18:3 response sensitivity in B73 and Mo17, leaves from IBM-RILs were analyzed for volatile emission after treatment with either water alone, 1 μM ZmPep3 or 1 μM Gln-18:3. Fold-change data for elicitor induced VOC emission was generated for 222 IBM RILs (Table S7). Association analyses were conducted in TASSEL 5.0 (Bradbury *et al.*, 2007) using the General Linear Model (GLM). Genotypic data from imputed IBM RIL SNP markers (July 2012 Imputed All Zea GBS final build; www.panzea.org) was filtered for a site minimum count of 10, minor allele frequencies of $> 10\%$ and major allele frequencies $< 90\%$ to generate 165,033 final SNP markers. Significantly associated SNPs were assigned following the Bonferroni P -value correction procedure using a cutoff of adjusted $P < 0.05$.

Generating a custom marker map based on SNPs called from RNA-seq data of 105 IBM RILs

Raw FASTQ files for B73, Mo17 and 105 IBM RILs were obtained from the NCBI (<https://www.ncbi.nlm.nih.gov/sra>) PRJNA179160 study accessions (Li *et al.*, 2013). The raw FASTQ files were first filtered using FASTP with default parameters (Chen *et al.*, 2018) and then aligned to the *Zea mays* B73 RegGen_4 genome obtained from plant ensemble version 44 using the STAR RNA-seq aligner with adjusted parameters (outFilterMultimapNmax 10, outFilterMismatchNoverLmax 0.04, outFilterIntronMotifs RemoveNoncanonicalUnannotated, alignIntronMax 6000) (Dobin *et al.*, 2013). The BCFTools mpileup and call functions were used for SNP calling in all annotated CDS gene regions using the filtering parameters of $\text{DP} \geq 10$ and $\text{QUAL}=999$ (Li, 2011). After SNPs were called, SNPs that had more than 10 lines with an allele read depth lower than 10 were removed. A final filtering step was conducted by removing all SNPs that disagreed with the SNPs called using the control B73 and the Mo17 inbred RNA-seq reference samples. Individual SNPs were assigned gene IDs based on their location and the genotypes

of genes with the associated SNPs were called based on aggregated most common SNP genotype in each gene. The genotypes of 10,043 gene-based markers were called by aggregating all the SNPs in each gene and selecting the most common allele.

RNA Isolation and measurement of transcript abundance by qRT-PCR

Total RNA was isolated with TRIzol (Invitrogen) as per the manufacturer's protocol and treated with DNase according to instructions (Life Technologies). M-MLV Reverse Transcriptase (Thermo Fisher Scientific) was used for cDNA synthesis with random decamer primers and 1 µg of RNA. For the qRT-PCR 1 µl of 2-fold diluted cDNA was added using SsoAdvancedtm Universal SYBR(R) Green Supermix (Bio-Rad) and a StepOneTM Real-Time PCR System (Applied Biosystems). Based on the threshold cycle (Ct) value, ΔC_t was calculated relative to 60S ribosomal protein L17 (RPL17). Transcript abundance was calculated relative to its corresponding untreated control.

Plasmid construction for transient assays and immunoblotting in *Nicotiana benthamiana*

ZmFACS (Zm00001d053867) was amplified from genomic B73 and Mo17 DNA and inserted into pENTR/D-TOPO vector (Invitrogen). Expression vectors of the corresponding *ZmFACS* entry vectors were generated through Gateway cloning into the destination vector pGWB441, to generate a C-terminal fusion with eYFP driven by the CaMV35S promoter (Nakagawa *et al.*, 2007). Expression vectors were transformed into *Agrobacterium tumefaciens* GV3101 (pMP90). *A. tumefaciens* carrying the expression vectors were infiltrated into *N. benthamiana* leaves at OD600 of 0.8 for transient expression. For immunoblotting, 50 mg of leaf tissue were ground in liquid N₂ and homogenized in 100 µL 2x SDS loading buffer (20% SDS was used in making the 5x SDS loading buffer, instead of 10%) for 5 minutes in 95°C. Western blotting was performed with α -GFP polyclonal primary antibodies (ThermoFisher) at 1:1000 dilution and α -rabbit secondary antibodies (SigmaAldrich) at 1:10000 dilution. SuperSignalTM West Pico PLUS (ThermoFisher) chemiluminescent substrate was used for protein detection.

Phylogenetic tree construction

The Phytozome blastn tool was used to identify coding sequences (CDS) related to FACS from *Zea mays*, *Oryza sativa* and *Arabidopsis thaliana* (Goodstein *et al.*, 2012). The CDS sequences of the putative genes were translated to amino acid sequences and the full sequences were aligned using MUSCLE with default parameters (Edgar, 2004). The best-fit model for the phylogenetic estimation was selected using ModelFinder (Kalyaanamoorthy *et al.*, 2017). A Maximum Likelihood (ML) phylogenetic tree was

constructed using IQ-TREE under the selected best-fit model, with 1000 bootstrap replications (Nguyen *et al.*, 2015). The tree was visualized and annotated using FigTree (<http://github.com/rambaut/figtree/>).

SUPPLEMENTARY DATA:

Figure S1: Comparison of overlapping ZmPep3- and Gln-18:3-induced DEGs

Figure S2: Genetic mapping sensitivity locus using the fold-change (FC) values of VOCs from three different biosynthetic pathways

Figure S3: Alignment of the encoded amino acid FACS sequences of B73 and Mo17

Figure S4: Native Gln-18:3 sensitivity in *N. benthamiana* is reduced in plants grown under diminished light intensity but heterologous expression of FACS increases responsiveness to Gln-18:3

Table S1: Early elicitor-induced transcriptomic data from treated B73 leaves

Table S2: Summary of elicitor-induced differential gene expression analysis

Table S3: Summary of elicitor-induced gene ontology (GO) term enrichment analysis

Table S4: Summary of elicitor-induced MAPMAN bin enrichment analysis

Table S5: Comparative analysis of elicitor-induced DEGs using ranked mean expression

Table S6: List of elicitor-induced DEGs used for the preparation of figure 2

Table S7: Elicitor-induced VOC data used for association mapping with the IBM mapping population

Table S8: Summary table of the FAC sensitivity locus derived from association mapping and fine mapping results

Table S9: The newly generated IBM RIL genetic marker map using RNA-seq data

Table S10: List of ZmFACS-related genes from *Zea mays*, *Oryza sativa* and *Arabidopsis thaliana* used to generate figure 7A

Table S11: List of primers used

AUTHOR CONTRIBUTIONS

AH conceived of research plans. EP, MR, NA, EAS and AH designed and performed experiments and analyzed the data. AS cloned maize receptor genes. KD prepared the samples for RNA-seq. EP, EAS and AH wrote the manuscript with input from all authors. AH agrees to serve as the author responsible for contact and ensures communication.

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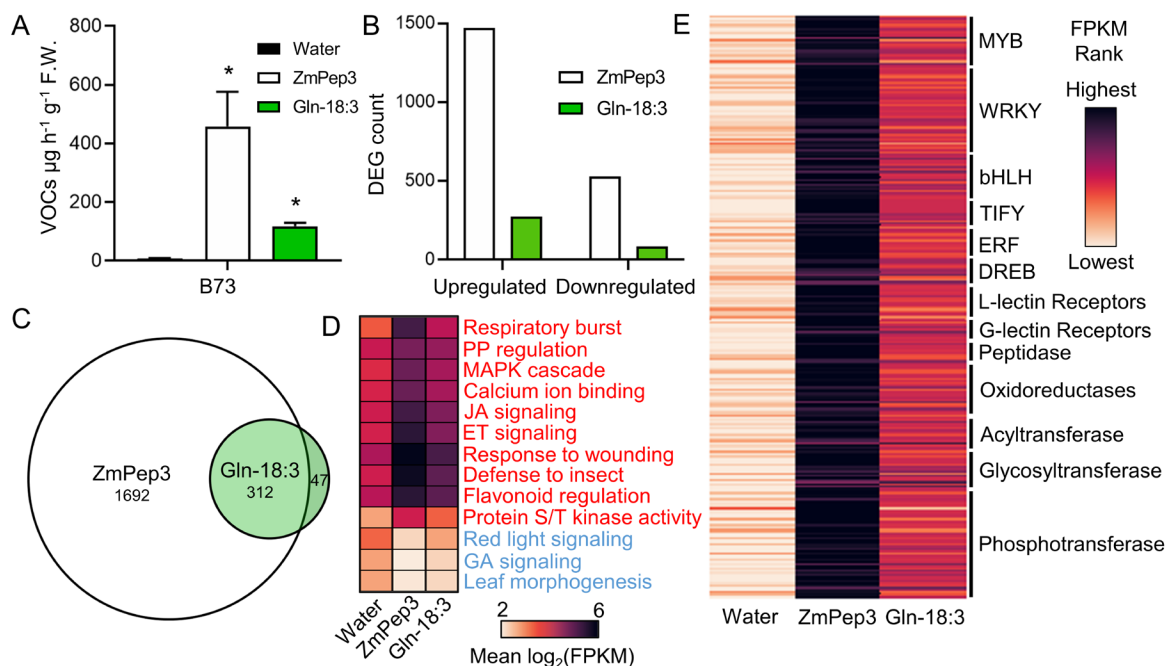


Figure 1. Comparative analysis of ZmPep3- and Gln-18:3-induced VOC emission and early transcriptional changes. **(A)** Analysis of total elicitor-induced VOC emission 16 hours after treatment of B73 leaves with water, 5 μM ZmPep3 or 5 μM Gln-18:3. Total VOC is represented by the sum of (Z)-3-hexenyl acetate, linalool, (3E)-4,8-dimethyl-1,3,7-nonatriene (DMNT), β -caryophyllene, (E)- α -bergamotene, (E)- β -farnesene and E-nerolidol. **(B)** Measurement of the number of differentially expressed genes (DEGs) following elicitor-induced 2-hour treatments measured by RNA-seq. **(C)** Euler diagram representing the overlap of the combined upregulated and downregulated DEGs. **(D)** Summary heatmap of the mean $\log_2(\text{FPKM})$ data of all ZmPep3- and Gln-18:3-induced DEGs in selected enriched GO terms. Enriched GO terms for upregulated DEGs are in a red font and downregulated DEGs are in a blue font. PP stands for phenylpropanoid **(E)** Summary heatmap of all ZmPep3 upregulated DEGs assigned to their enriched MAPMAN bins with more than 10 DEGs in respective bins. Row colors are based on the treatment mean of the ranked FPKM values for each gene. For all treatments shown $n=4$ and error bars represent SEM. Student t-tests (two-tailed distribution, unpaired), was used for detection of significant differences. Asterisks indicate significant difference with $P < 0.05$; ns, not significant.

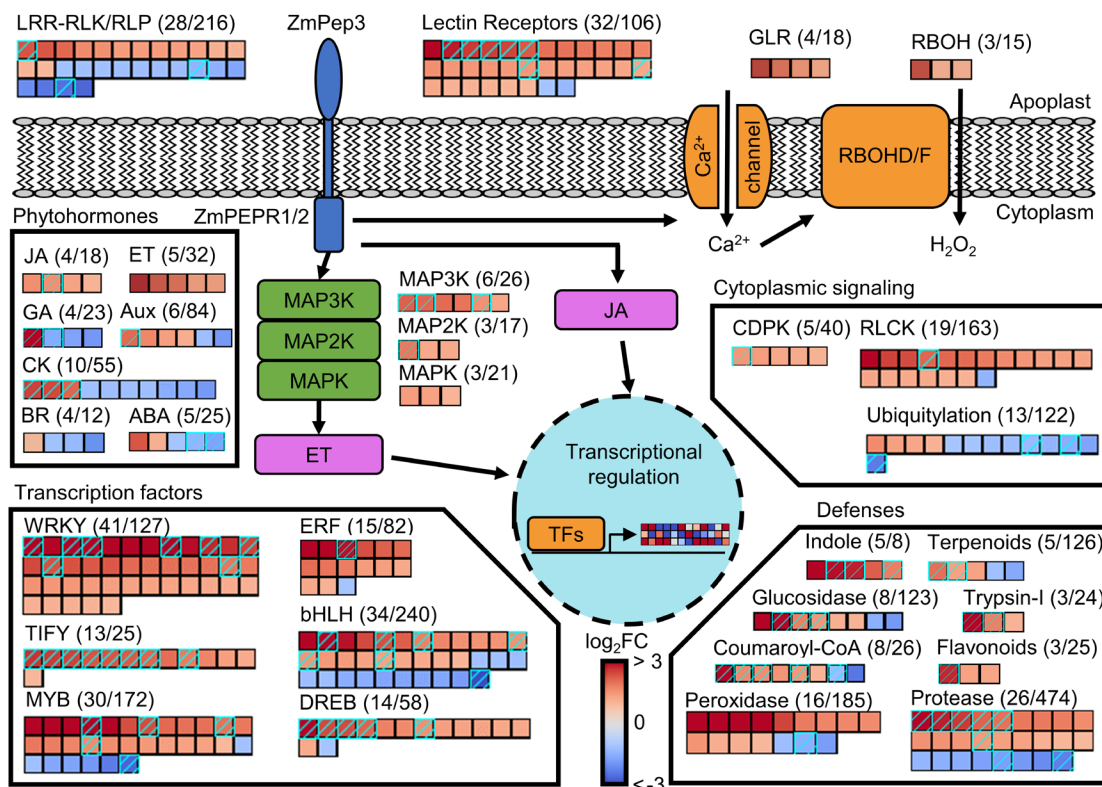


Figure 2. Early elicitor-induced transcriptional regulation of multiple components associated with antiherbivore responses. Heatmaps of gene groups represent \log_2 fold-change (FC) data from RNA-seq results of ZmPep3-induced differentially expressed genes and based the *Zea mays* V4 MAPMAN annotation. Hatched cyan boxes indicate genes that were also significantly differentially expressed following Gln-18:3 treatment. The selected gene groups were obtained either from the MAPMAN bin annotations or from searching selected keywords in the MAPMAN gene descriptions in the case of the glucosidases, peroxidases, proteases and trypsin-inhibitors (Trypsin-I) gene groups. Parenthesis next to each abbreviated group title are the number of ZmPep3 elicited DEGs compared to the size of the gene group. Abbreviations are as follows: leucine-rich repeat receptor like kinase/receptor like protein (LRR-RLK/RLP); glutamate-like receptors (GLR); respiratory burst oxidase homolog (RBOH); jasmonic acid (JA); ethylene (ET); gibberellic acid (GA); auxin (Aux); cytokinins (CK); brassinosteroids (BR); abscisic acid (ABA); mitogen activated protein kinase (MAPKs); calcium-dependent protein kinase (CDPK); receptor-like cytoplasmic kinase (RLCK); transcription factors (TFs); ethylene response factor (ERF); and Trypsin inhibitor (Trypsin-I). All associated B73 V4 gene IDs and transcript \log_2 FC data underlying this figure are detailed in Table S6.

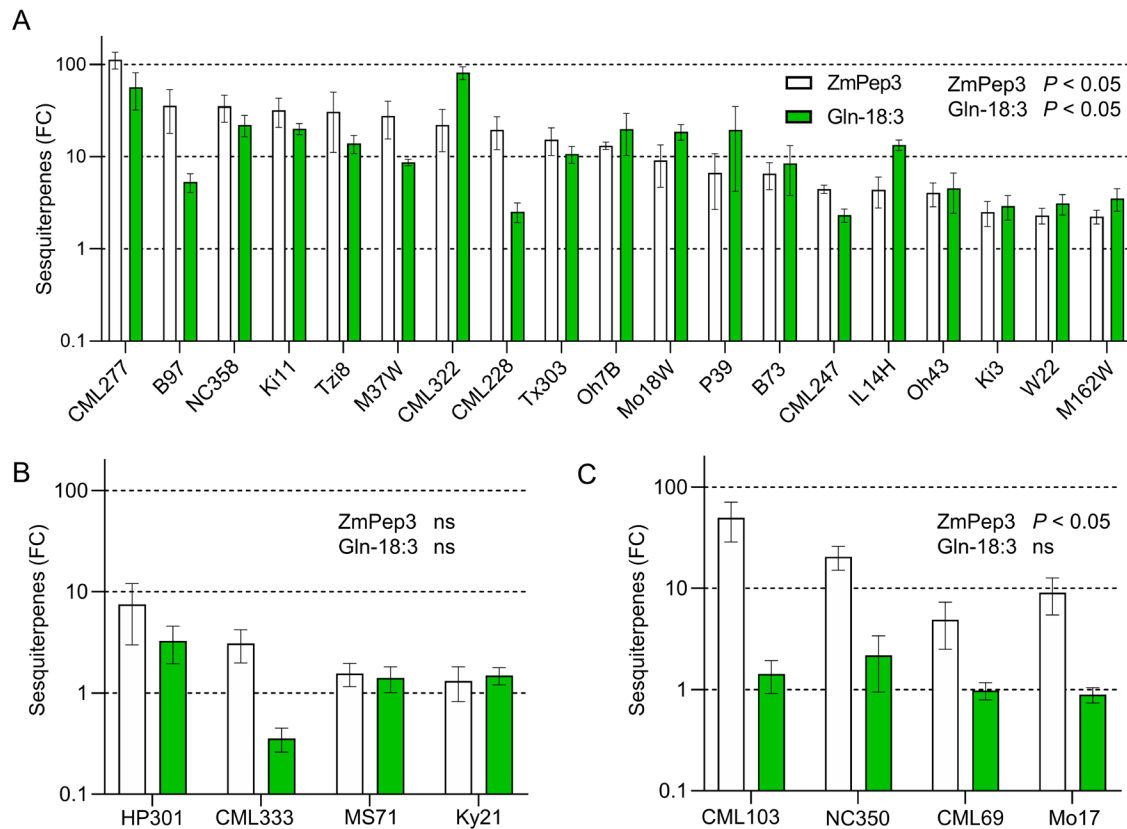


Figure 3. Diverse maize inbred lines display significant variation in HAMP and DAMP elicited defense responses of volatile sesquiterpenes. A total of 27 maize inbred lines including B73, Mo17, W22 and the Nested Association Mapping parent lines were analyzed for elicitor-induced volatile sesquiterpene emission. Maize leaves were treated for 16 hours with water, 1 μ M ZmPep3 or 1 μ M Gln-18:3 and volatiles were analyzed by gas chromatography. Results are presented as the fold-change (FC) in elicitor-induced sesquiterpene emission compared to the water treatment. Sesquiterpenes are defined as the sum of β -caryophyllene, (*E*)- α -bergamotene, (*E*)- β -farnesene and *E*-nerolidol. For all treatments shown, biological replicates ($n=4$) and error bars represent SEM. Significance of elicitor-induced sesquiterpene emission was determined by Student t-tests (two-tailed distribution, unpaired), for each line, and was used to classify the maize inbred lines into three categories: **(A)** Lines with both significant HAMP and DAMP elicitor-induced responses ($P < 0.05$), **(B)** lines where both elicitor-induced responses not statistically significant (ns), and **(C)** lines with significant ZmPep3-induced responses but not significant Gln-18:3 induced responses.

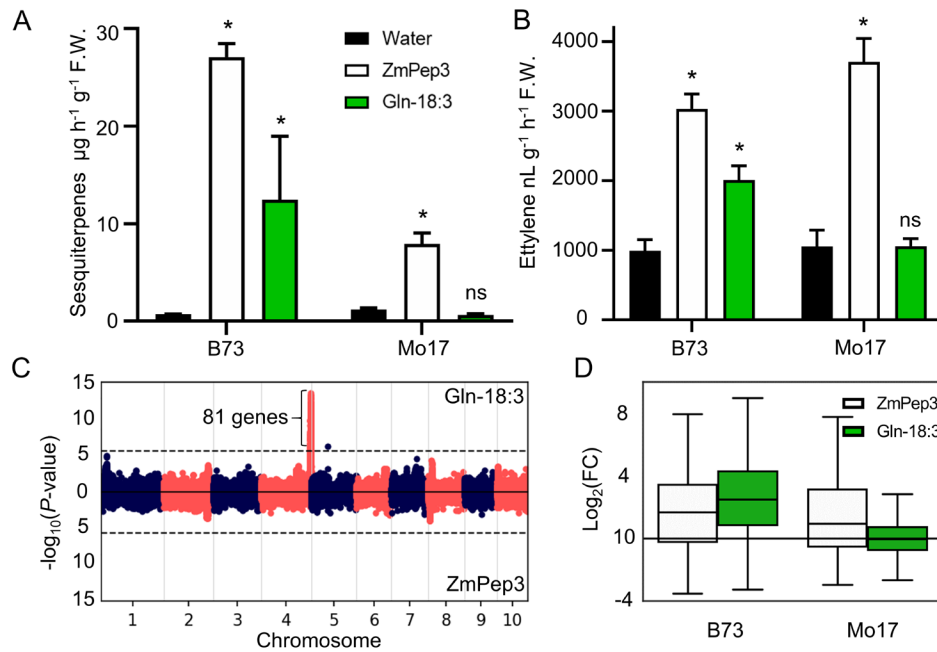


Figure 4. Forward genetics approaches using association analyses identify a maize FAC sensitivity locus based on Gln-18:3 specific response impairments in Mo17. **(A)** Average ($n=4$, +SEM) total sesquiterpene emission after 16 hours and **(B)** average ethylene ($n=6$, +SEM) emission after 2 hours following treatment of B73 and Mo17 leaf 5 with either water, 5 μM ZmPep3 or 5 μM Gln-18:3. Sesquiterpenes are specifically the sum of β -caryophyllene, (*E*)- α -bergamotene, (*E*)- β -farnesene and *E*-nerolidol. **(C)** Metabolite based association analyses using 242 Interrelated B73 x Mo17 (IBM) Recombinant Inbred Lines (RILs) and elicited VOC fold-change values as traits derived from leaf 5 treated with either water, 1 μM ZmPep3- and 1 μM Gln-18:3. VOCs were analyzed 16 hours after treatments. Imputed IBM RIL SNP markers (165,033) were obtained from the Panzea project (www.panzea.org). Fold-change values were calculated using the sum of the total VOC, defined as combination of (*Z*)-3-hexenyl acetate, linalool, (3*E*)-4,8-dimethyl-1,3,7-nonatriene (DMNT), β -caryophyllene, (*E*)- α -bergamotene, (*E*)- β -farnesene and *E*-nerolidol. **(D)** Box plots showing the distribution of elicitor-induced fold-change VOC emission at the top SNP (S4_237322925) associated with FAC sensitivity across IBM RILs. Dashed lines in the Manhattan plots indicate a significance cutoff of $P < 0.05$ based on Bonferroni correction. Number of genes are based on B73 V4 annotations. Error bars represent SEM. Student t-tests (two-tailed distribution, unpaired), was used for detection of significant differences. Asterisks indicate significant difference with $P < 0.05$; ns, not significant.

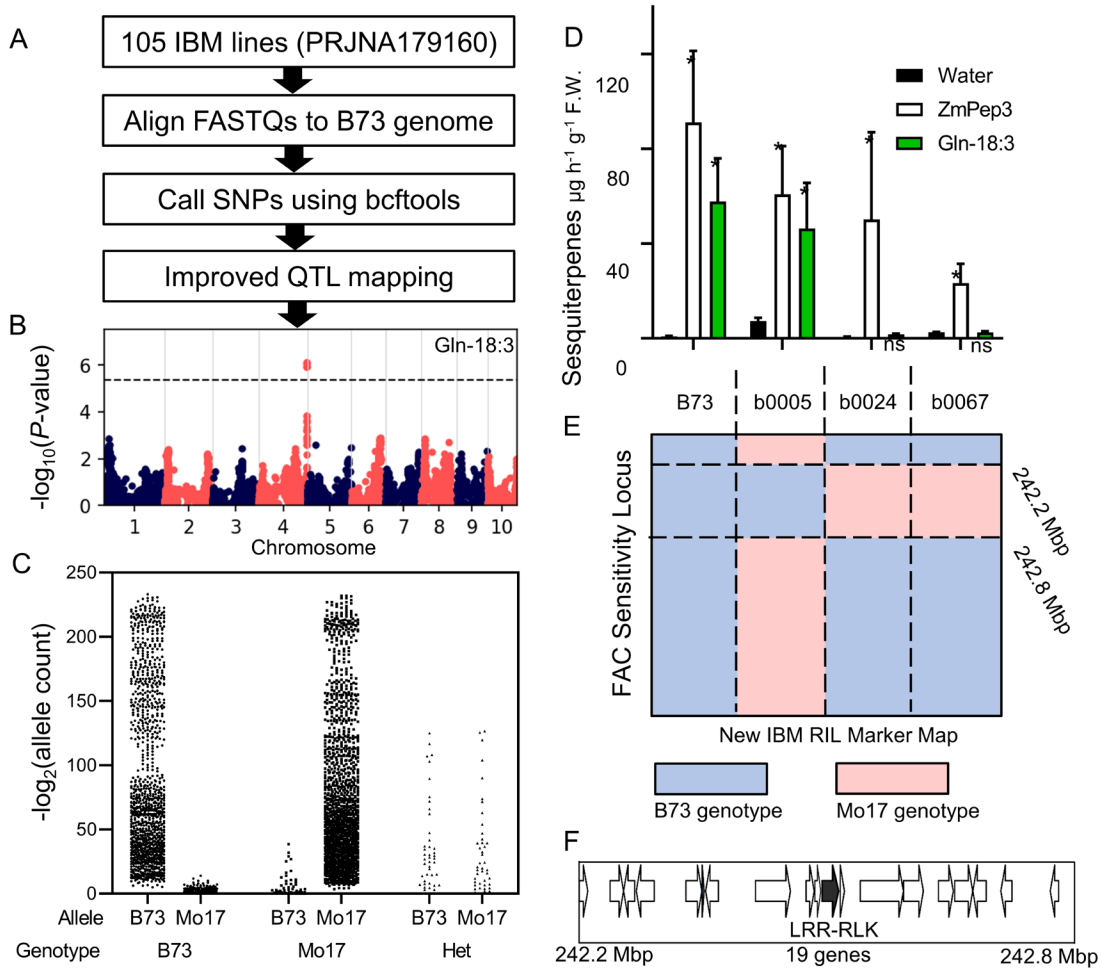


Figure 6. Fine-mapping the FAC sensitivity locus using a newly generated marker map for 105 IBM RILs. **(A)** Summarized procedural workflow describing creation of the new IBM RIL marker map using RNA-seq data from 105 IBM RILs obtained from NCBI aligned to the B73 RefGen_4 genome. **(B)** Association analyses using the the new IBM RIL map (10,043 marker genes) with total VOC fold-change values from leaf 5 of 86 RILs treated with either water, 1 μ M ZmPep3- or 1 μ M Gln-18:3 analyzed 16 hours after treatment. Association analyses (General Linear Model) with newly assigned markers and VOC fold-change traits follows from Fig. 4C. Dashed lines in Manhattan plots indicate significance cutoff of $P < 0.05$ based on Bonferroni correction. **(C)** Aggregated allele counts of all SNPs within the FAC sensitivity locus that were used for generating the new IBM RIL marker map compared to the final assigned marker genotype. Het stands for heterozygous. **(D)** Average ($n=3$, +SEM) elicitor-induced sesquiterpene emission, as measured by the sum of β -caryophyllene, α -bergamotene, and E - β -farnesene 16 hours after treatment of B73 and IBM RILs b0005, b0024 and b0067 leaves with water, 5 μ M ZmPep3 or 5 μ M Gln-18:3. **(E)** Corresponding newly created IBM-RIL marker map of the FAC sensitivity locus. **(F)** The relative location of 19 genes present in the refined FAC sensitivity locus with the LRR-RLK marked as a black arrow. Coordinates and gene numbers are based on the B73 V4 annotations. Student t-tests (two-tailed distribution, unpaired), were used for detection of significant differences. Asterisks indicate significant difference with $P < 0.05$; ns, not significant.

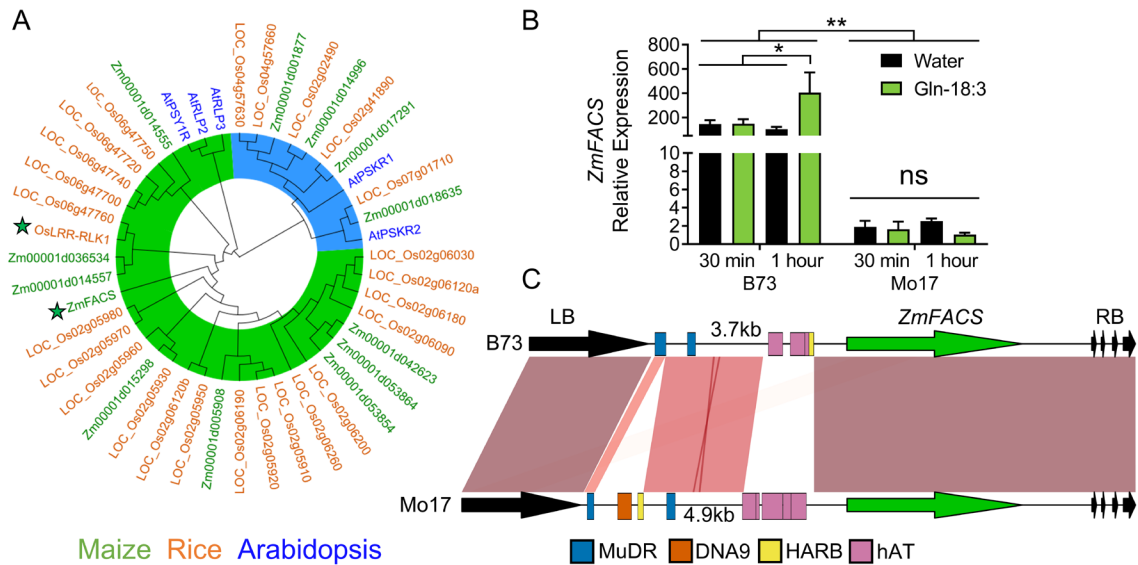


Figure 7. Candidate gene within the FAC sensitivity locus, denoted *ZmFACS*, is an ortholog of *Oryza sativa* *OsLRR-RLK1* necessary for antiherbivore responses. **(A)** Phylogenetic tree generated from the protein sequences of genes related to *ZmFACS* and *OsLRR-RLK1* from *Arabidopsis thaliana* (blue font), *Oryza sativa* (orange font) and *Zea mays* (green font). Based on similarity to the *A. thaliana* genes, the phylogeny tree reveals two clades of *AtPSKR1/2*-related genes (highlighted in blue) and *AtPSY1R*-related genes (highlighted in green). *ZmFACS* and *OsLRR-RLK1* proteins are designated by a green star. **(B)** Average ($n=3$, +SEM) relative expression of *ZmFACS* in leaf 5 of the maize B73 and Mo17 inbred lines treated with either water or Gln-18:3 for 0.5 and 1 hours determined by quantitative RT-PCR in comparison with the expression of the control gene *Rpl17*. **(C)** Comparative promoter sequence similarity visualization of the *ZmFACS* (blue arrows) in B73 and Mo17 including the neighboring genes upstream (LB - Left border) and downstream (RB - Right border) from *ZmFACS* for reference. The 3.7kb and 4.9kb sequences represent the genomic distance between the *ZmFACS* start codon and the LB stop codon and were used for transposable element prediction using the Poaceae database at the Repbase Censor to identify a number of transposable element fragments from the MuDR family (blue), DNA9 (orange), Harbinger (HARB; yellow) and hAT (purple) families. Student t-tests (two-tailed distribution, unpaired), was used for detection of significant differences. Asterisks indicate significant difference with $P < 0.05$; ns, not significant.

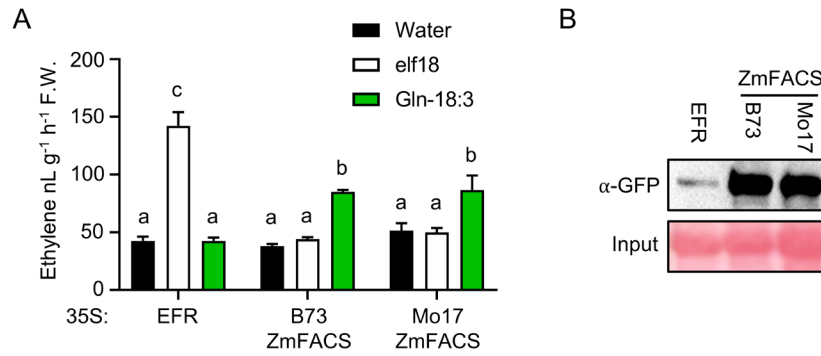


Figure 8. Transient heterologous expression in tobacco (*N. benthamiana*) of both B73 and Mo17 ZmFACS increases plant responsiveness to Gln-18:3 elicitation. **(A)** Average (n=4, +SEM) elicited ethylene emission 2 hours following treatment with water, 1 μ M elf18 or 1 μ M Gln-18:3 in *N. benthamiana* leaves expressing either YFP-fusion proteins with *Arabidopsis thaliana* EFR (control), B73 ZmFACS and Mo17 ZmFACS. **(B)** Detection of EFR and ZmFACS cloned from B73 and Mo17 protein expression with α -GFP using a western blot. ANOVA $P < 0.05$, Tukey honestly significant difference (HSD) tests ($\alpha = 0.05$) with different letters (a-c) representing significant differences.

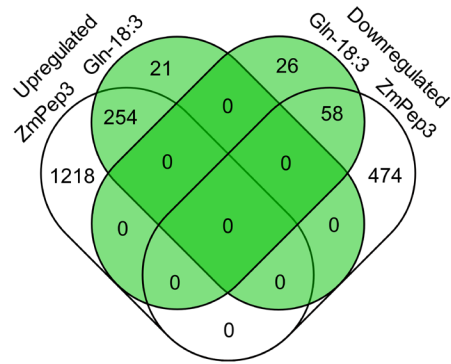


Figure S1. Comparison of ZmPep3- and Gln-18:3-induced DEGs in B73 maize leaves reveals large scale overlap in HAMP and DAMP elicited regulation. Venn diagram comparison of the upregulated and downregulated DEGs following ZmPep3 and Gln-18:3 treatments found using RNA-seq data after 2 hour treatment of maize B73 leaves. The upregulated and downregulated DEGs for each treatment were used to identify enriched GO terms and Mapman bins using the hypergeometric test.

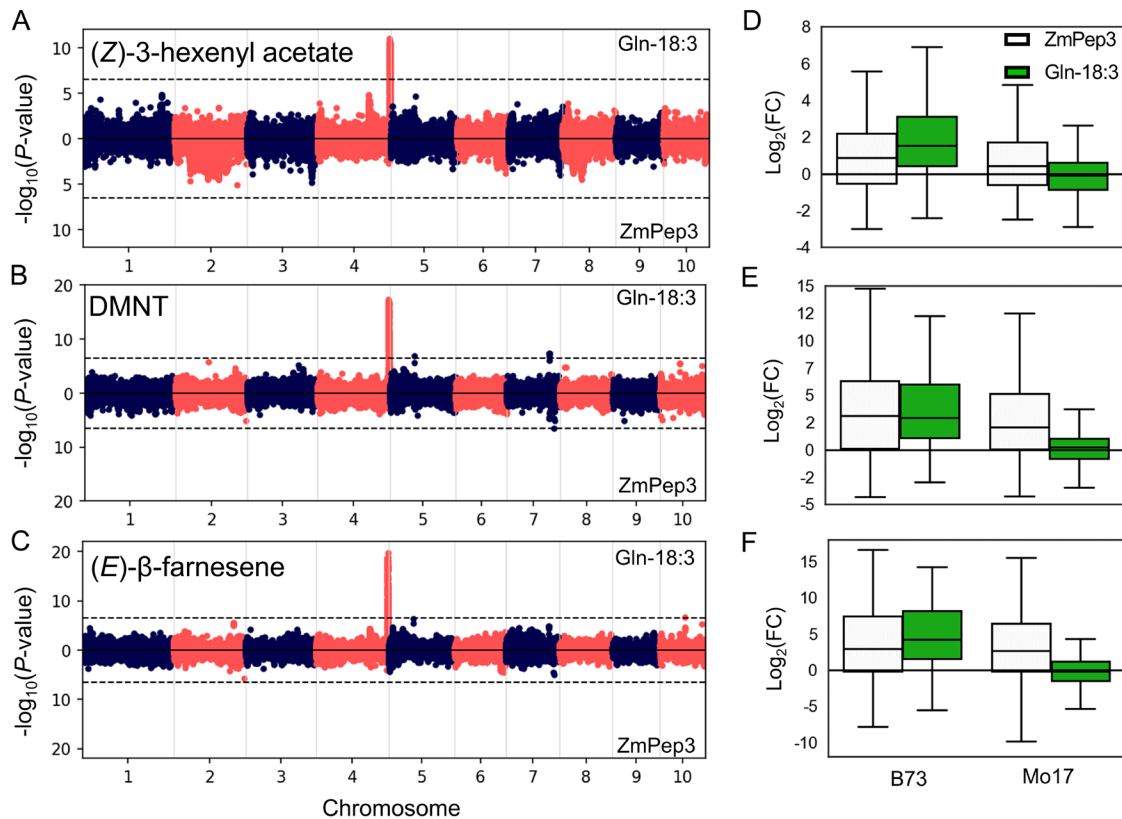


Figure S2. A forward genetics study using association analyses supports a shared maize Gln-18:3 sensitivity locus using elicited VOCs from three different biosynthetic pathways. Association analyses using imputed SNP markers (165,033) obtained from the Panzea project for the IBM RIL population and VOC fold-change values as traits derived from leaf 5 of 242 RILs treated with water, 1 μ M ZmPep3- and 1 μ M for 16 hours. Fold change (FC) values were calculated for individual elicited VOCs from different biosynthetic pathways, including (A) (Z)-3-hexenyl acetate, (B) (3E)-4,8-dimethyl-1,3,7-nonatriene (DMNT) and (C) (E)-β-farnesene. Box plots showing the distribution of elicitor-induced fold-change VOC emission for (D) (Z)-3-hexenyl acetate, (E) DMNT and (F) (E)-β-farnesene at the highest statistically significant SNP (S4_237322925) associated with FAC sensitivity across the 242 IBM RILs. Dashed lines in Manhattan plots indicate significance cutoff of $P < 0.05$ based on Bonferroni correction.

FACS (B73) MLVPIPTRSHAATPRFMQEKPEPEPEPEHTARTSFHRPRALALALLLISISWESCASACGEPERASLLQFLAELSYDAGL 80
FACS (Mo17) MLVPIPTRSHAATPRFMQEKPEPEPEPEHTARTSFHRPRALALALLLISISWESCASACGEPERASLLQFLAELSYDAGL 80

FACS (B73) TGLWRGTDCCKWEGITCDDQYGTAVTVSAISLPGRGLEGRISQSLASLAGLRRRLNLSYNSLSGDLPLGLVSASGSVAVLD 160
FACS (Mo17) TGLWRGTDCCKWEGITCDDQYGTAVTVSAISLPGRGLEGRISQSLASLAGLRRRLNLSYNSLSGDLPLGLVSASGSVAVLD 160

FACS (B73) VSFNQSLGDLSPAPGQRPLQLQVLNISSNSFTGQLTSTAWERMRSVLNALNASNNSLTGQIPDQFCATAPSFVLELSYN 240
FACS (Mo17) VSFNQSLGDLSPAPGQRPLQLQVLNISSNSFTGQLTSTAWERMRSVLNALNASNNSLTGQIPDQFCATAPSFVLELSYN 240

FACS (B73) KFSGGVPPGLGNCMSLRVLRAGHNLSGTLPRELFNATSLERLSFSNLFHGTVDGAHVAKLSNLVLDLGDNSFGGKIP 320
FACS (Mo17) KFSGGVPPGLGNCMSLRVLRAGHNLSGTLPRELFNATSLERLSFSNLFHGTVDGAHVAKLSNLVLDLGDNSFGGKIP 320

FACS (B73) DTIGQLKRLQELHLDYNSMYGELPPALSNCDDLITLDRSNGFSGELSRVDFSNMPSLRTIDLMLNFFSGTIPESIYSCR 400
FACS (Mo17) DTIGQLKRLQELHLDYNSMYGELPPALSNCDDLITLDRSNGFSGELSRVDFSNMPSLRTIDLMLNFFSGTIPESIYSCR 400

FACS (B73) NLTAIRLASNKFHGLSEGLGNLKSLSFLSLTNNSLSNITNALQILRSSKNLTLLGINFFEETIPDDAVIYGFENLQV 480
FACS (Mo17) NLTAIRLASNKFHGLSEGLGNLKSLSFLSLTNNSLSNITNALQILRSSKNLTLLGINFFEETIPDDAVIYGFENLQV 480

FACS (B73) LDIGNCLLSGEIPLWISKLVNLEMLFLDGNRLSGSIPTWIHTLEYLFYLDISNNSLTGEIPKEVVSIPMLTSERTAHL 560
FACS (Mo17) LDIGNCLLSGEIPLWISKLVNLEMLFLDGNRLSGSIPTWIHTLEYLFYLDISNNSLTGEIPKEVVSIPMLTSERTAHL 560

FACS (B73) ASVFDLPVYDGPSRQYRIPIAFPKVLNLSNRFTGQIPPEIGQLKGLSLDISNNSLTGPIPTSICNLNLLVLDLSSND 640
FACS (Mo17) ASVFDLPVYDGPSRQYRIPIAFPKVLNLSNRFTGQIPPEIGQLKGLSLDISNNSLTGPIPTSICNLNLLVLDLSSND 640

FACS (B73) LTGKIPVALENLHFLSTFNVSNNDLEGIPTGGQFGTFQNSSFLGNPKLCGFMIGRRCSADVPLVSTGGRNKKAILAIA 720
FACS (Mo17) LTGKIPVALENLHFLSTFNVSNNDLEGIPTGGQFGTFQNSSFLGNPKLCGFMIGRRCSADVPLVSTGGRNKKAILAIA 720

FACS (B73) FGVFFAMIAILLWRLVLSIRINRLTAQGRREDNGYLETSTFNSSLEHGVIMVPQKGKGNENKLTFSDIVKATNNFNKEN 800
FACS (Mo17) FGVFFAMIAILLWRLVLSIRINRLTAQGRREDNGYLETSTFNSSLEHGVIMVPQKGKGNENKLTFSDIVKATNNFNKEN 800

FACS (B73) IIGCGGYGLVYKAELPDGCKLAIKKLNDEMCLMEREFTAEEALSMAQHDHLVPLWGYCIQGNRSFLIYSYMENGLDDW 880
FACS (Mo17) IIGCGGYGLVYKAELPDGCKLAIKKLNDEMCLMEREFTAEEALSMAQHDHLVPLWGYCIQGNRSFLIYSYMENGLDDW 880

FACS (B73) LHNRRDDASTFLDWPTLRRIAQGASRGLSYIHNDCKPIVHRDIKCSNILLDKELKAYVADFGLSRLILPNKTHVTTEL 960
FACS (Mo17) LHNRRDDASTFLDWPTLRRIAQGASRGLSYIHNDCKPIVHRDIKCSNILLDKELKAYVADFGLSRLILPNKTHVTTEL 960

FACS (B73) GTLGYPPEYAHGWVATLRGDIYSFGVVLLELLTGLRPVPLVLTSSKELVPWVLEMSSQGLVDVLDPTLCGTGHEEQMLK 1040
FACS (Mo17) GTLGYPPEYAHGWVATLRGDIYSFGVVLLELLTGLRPVPLVLTSSKELVPWVLEMSSQGLVDVLDPTLCGTGHEEQMLK 1040

FACS (B73) VLGLACKCVNNNPIMRPHIMEVVTCLSEINVGLQAQKSVKTIQLASYT 1088
FACS (Mo17) VLGLACKCVNNNPIMRPHIMEVVTCLSEINVGLQAQKSVKTIQLASYT 1088

Figure S3. Alignment of the encoded amino acid ZmFACS sequences of B73 and Mo17. The amino acid sequences for B73 ZmFACS (Zm00001d053867) and Mo17 ZmFACS (Zm00014a001621) were aligned using Clustal Omega and visualized by highlighting in grey all matching amino-acids. The signal peptide, colored red, was predicted for both B73 ZmFACS and Mo17 ZmFACS using Signal-3L. The predicted LRR sequences, with a single predicted island domain between them, were manually annotated based on the consensus LRR motif sequence and labeled using a blue arrow below the matching LRR sequences. Transmembrane domain was predicted using Phobius and labeled using a black arrow below the matching sequence. Two predicted protein domains by InerPro using the amino-acid of B73 ZmFACS were included, the ATP binding site (IPR017441) and the Serine/threonine-protein kinase active site (IPR008271) and labeled using an orange arrows below the matching sequence.

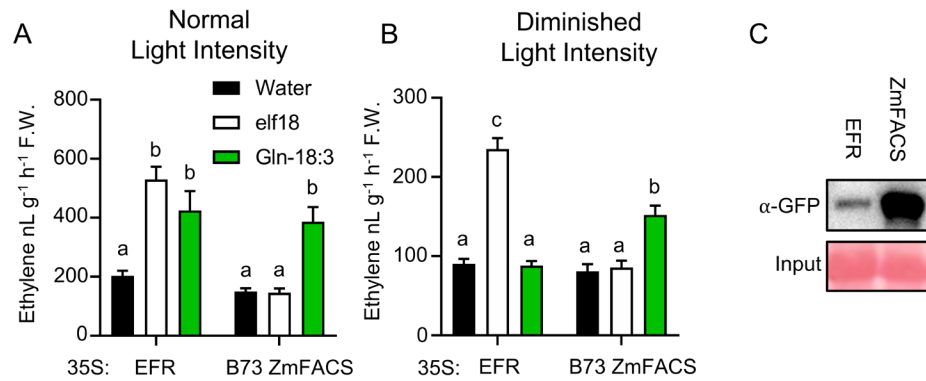


Figure S4. Native Gln-18:3 response sensitivity in *N. benthamiana* is reduced in plants grown under diminished light intensity; however, transient heterologous expression of ZmFACS increases and restores responsiveness to Gln-18:3. **(A)** Average (n=4, +SEM) ethylene emission after 2-hour treatments with either water, 1 μ M elf18 or 1 μ M Gln-18:3 in *N. benthamiana* leaves expressing YFP-fusion proteins with either *Arabidopsis thaliana* EFR or B73 ZmFACS. Plants were grown under normal light intensity, as described in the method section. **(B)** Average (n=4, +SEM) ethylene emission after 2-hour treatments with either water, 1 μ M elf18 or 1 μ M Gln-18:3 in *N. benthamiana* leaves expressing YFP-fusion proteins with either *Arabidopsis thaliana* EFR or B73 ZmFACS. Plants were grown under diminished light intensity, as described in the method section. **(C)** Detection of YFP-fusion *Arabidopsis thaliana* EFR and ZmFACS cloned from B73 using α -GFP western blot. Different letters (a, b, c) represent significant differences (All ANOVAs $P < 0.05$ followed by Tukey honestly significant difference (HSD), $\alpha = 0.05$).