Primed refers to a mechanism whereby plants are sensitized to respond faster and/or more strongly to future pathogen attack. Here, we demonstrate that preexposure to the green leaf volatile Z-3-hexenyl acetate (Z-3-HAC) primed wheat (Triticum aestivum) for enhanced defense against subsequent infection with the hemibiotrophic fungus Fusarium graminearum. Bioassays showed that, after priming with Z-3-HAC, wheat ears accumulated up to 40% fewer necrotic spikelets. Furthermore, leaves of seedlings showed significantly smaller necrotic lesions compared with nonprimed plants, coinciding with strongly reduced fungal growth in planta. Additionally, we found that F. graminearum produced more deoxynivalenol, a mycotoxin, in the primed treatment. Expression analysis of salicylic acid (SA) and jasmonic acid (JA) biosynthesis genes and exogenous methyl salicylate and methyl jasmonate applications showed that plant defense against F. graminearum is sequentially regulated by SA and JA during the early and later stages of infection, respectively. Interestingly, analysis of the effect of Z-3-HAC pretreatment on SA- and JA-responsive gene expression in hormone-treated and pathogen-inoculated seedlings revealed that Z-3-HAC boosts JA-dependent defenses during the necrotrophic infection stage of F. graminearum but suppresses SA-regulated defense during its biotrophic phase. Together, these findings highlight the importance of temporally separated hormone changes in molding plant health and disease and support a scenario whereby the green leaf volatile Z-3-HAC protects wheat against Fusarium head blight by priming for enhanced JA-dependent defenses during the necrotrophic stages of infection.

Biogenic volatile organic compounds (BVOCs) are known regulators of communication of sedentary plants with their direct environment (Dudareva et al., 2006). Besides attracting pollinators (Pichersky and Gershenzon, 2002), repelling insect herbivores (Birkett et al., 2010), and exerting direct antimicrobial properties (Friedman et al., 2002), BVOCs can act as an alarm signal to warn neighboring plants of an imminent herbivorous or pathogen attack (Heil and Ton, 2008) or serve as an intraplant signal for the induction of resistance (Karban et al., 2006). Engelberth et al. (2004) found that maize (Zea mays) seedlings emitted the green leaf volatiles (GLVs) Z-3-hexenal, Z-3-hexenol (Z-3-HOL), and Z-3-hexenyl acetate (Z-3-HAC) after they had been infested with caterpillars of Spodoptera exigua. Neighboring uninfested seedlings that had been exposed to these GLVs subsequently showed a considerable higher production of the plant defense hormone jasmonic acid (JA) after treatment with caterpillar regurgitant. This form of induced resistance is called priming. Plants in a primed state display faster and/or stronger activation of defense pathways when challenged by microbial pathogens, herbivorous insects, or abiotic stresses (Conrath, 2009). Exposure to these priming signals does not entail a direct activation of costly defense mechanisms but rather a stronger up-regulation of defense pathways when the plant is actually under attack (van Hulten et al., 2006). Besides resulting in a stronger induction of the JA pathway, priming also has been shown to enhance defense associated with the salicylic acid (SA) pathway, which plays a critical role in plant
defense against biotrophic pathogens (Conrath et al., 2006; Jung et al., 2009).

The lion’s share of attention on the use of GLVs in induced resistance has been directed to plant-insect interactions. However, the literature regarding priming by GLVs in plant-pathogen interactions remains scarce (Heil, 2014). Few studies have been performed investigating the effect of priming by GLVs on plant-fungus interactions (Scala et al., 2013a, and refs. therein). For example, hexanoic acid, a molecule with a similar structure to GLVs, has been shown to act as a priming agent in tomato (Solanum lycopersicum) plants against an infection by the necrotrophic fungus Botrytis cinerea, leading to a reduced accumulation of reactive oxygen species in primed plants (Vicedo et al., 2009; Kravchuk et al., 2011; Finiti et al., 2014). Since the GLVs E-2-hexenal (E-2-HAL), Z-3-HOL, E-2-hexeno1, and Z-3-HAC also have been reported to be emitted by perennial ryegrass (Lolium perenne) after infection with Fusarium poae (Parika et al., 2013) and by wheat (Triticum aestivum) seedlings after infection with Fusarium graminearum (Piesik et al., 2011), one may speculate that GLVs not only serve as a priming agent against the impending threat of herbivorous insects but rather constitute a general warning and priming mechanism against insects, bacteria, and fungal alike.

Fusarium head blight (FHB) is an important disease in cereals caused by a complex of Fusarium spp., of which the hemibiotrophic F. graminearum is one of the most prevalent (Parry et al., 1995; Goswami and Kistler, 2004; Audenaert et al., 2009). Besides yield losses of up to 40%, FHB also confers quality losses because of the production of mycotoxins such as deoxynivalenol (DON; Parry et al., 1995; Bottalico and Perrone, 2002; Vanheule et al., 2014).

The hemibiotrophic nature of F. graminearum entails that its lifestyle is characterized by a biotrophic phase followed by a necrotrophic phase. During the biotrophic phase, spores will germinate and hyphae will grow extracellularly and intercellularly. To counteract fungal colonization during the biotrophic phase, the host plant will accumulate hydrogen peroxide (H2O2) to induce programmed cell death. However, H2O2 acts as a signal for F. graminearum to produce DON, which in turn creates a positive feedback loop leading to increased H2O2 and DON production, clearing the way for F. graminearum to further colonize the host plant (Desmond et al., 2008). Plant defense against the biotrophic and necrotrophic phases generally has been linked to SA- and JA-related pathways, respectively (Glazebrook, 2005). This was also found in the study by Ding et al. (2011). They reported higher endogenous SA concentrations during the first hours of infection, followed by a rise in JA concentrations later on. However, plant defense against pathogens is regulated by a whole array of plant hormones, between which an intricate cross talk exists (Pieterse et al., 2012). One of the best-studied antagonistic signaling pathways is between SA and JA (Thaler et al., 2002; Pieterse et al., 2012). Research investigating the hormonal modulation of plant immunity has been done primarily in dicots. The negative relationship between SA and JA also seems to be conserved in rice (Oryza sativa), another monocot (De Vleesschauwer et al., 2013). Because of the presence of this possible antagonistic signaling and the hemibiotrophic lifestyle of F. graminearum, it is important to look more closely to the effect of priming on these two defense pathways in wheat.

Here, we show that preexposure of wheat to the GLV Z-3-HAC primes wheat plants for an enhanced defense against a future infection with F. graminearum. Furthermore, our results indicate that pretreatment with Z-3-HAC leads to a stronger activation of JA-related defense while exerting suppressive effects on SA-responsive gene expression. Lastly, we found evidence that enhanced plant defense led to increased DON production by F. graminearum.

RESULTS

Effect of Z-3-HAC Pretreatment on the Severity of Infection by F. graminearum

To assess whether preexposure to Z-3-HAC results in increased resistance against infection with F. graminearum, we point inoculated 27 spikelets of nine wheat ears, the target tissue of F. graminearum, with 20 µL of a conidia suspension for the preexposed and control treatments. Four days after infection, we observed the first necrotic lesions (Fig. 1A). The preexposed treatment exhibited a significantly lower infection rate than the control treatment, which persisted after 6 d. All inoculated spikelets showed necrosis 8 d after infection for both treatments. Previously, Purahong et al. (2012) reported on the high correlation between FHB resistance levels of wheat ears in field trials and those of detached leaves in a petri dish bioassay. Given the experimental tractability of the latter assays, we next tested the ability of Z-3-HAC to reduce FHB development and severity in a series of detached leaf experiments. At 24 h after infection (hai), we did not find significant differences in lesion length between preexposed and control seedlings. However, at 48 and 72 hai, lesion length was significantly larger (+20%, P < 0.05, +72%, P < 0.01, respectively) in control seedlings compared with preexposed seedlings (Fig. 1B). Additionally, lesions of preexposed seedlings showed an easily distinguishable brown clear center, while control seedlings showed more water-soaked lesions (Fig. 1C).

To investigate the effect of seedling preexposure to Z-3-HAC on fungal growth, we inoculated leaf sheaths with a conidia suspension using the leaf sheath bioassay. At each time point, we consistently found lower fungal biomass in the preexposed seedlings as compared with the control treatment (24 hai, −39%, P = 0.25; 48 hai, −65%, P < 0.05; and 72 hai, −94%, P < 0.05; Fig. 1D).

We inoculated leaf sheaths of wheat seedlings with a conidia suspension in order to establish the time point at which the fungal hyphae form infection structures and invade the plant cell wall. After an incubation period of 24 h, we found a large formation of appressoria-like structures and foot structures (Jansen et al., 2005; Boenisch and Schäfer, 2011; Fig. 1, E and F) and decided to take this time point as a starting point for further experiments.
Effect of Z-3-HAC, SA, and JA on Plant Defense during the Infection Process of *F. graminearum*

**Expression of JA and SA Biosynthesis Genes after *F. graminearum* Infection**

Because of the hemibiotrophic lifestyle of *F. graminearum*, we verified whether a sequential up-regulation of the biosynthesis genes for the SA and JA pathways was present. We selected *PHENYLALANINE AMMONIA LYASE* (*PAL*) and *ISOCHORISMATE SYNTHASE* (*ICS*) as marker genes for the biosynthesis of salicylate (Ding et al., 2011) and *LIPOXYGENASE1* (*LOX1*) and *LOX2* as marker genes for the biosynthesis of JA (Feng et al., 2010). Using the leaf sheath bioassay, we infected...
seedlings with a conidia suspension of *F. graminearum*. Expression analysis revealed at 24 hai a significant up-regulation of PAL ($P < 0.05$), while the expression of ICS, LOX1, and LOX2 was not significantly different from the control treatment (Fig. 2). However, at 48 hai, we saw a significant up-regulation of LOX1 and LOX2 ($P < 0.05$), while the expression of PAL was not different from the control and ICS was even significantly down-regulated ($P < 0.05$; Fig. 2).

**Effect of Exogenous Methyl Salicylate and Methyl Jasmonate on Disease Development**

As defense against *F. graminearum* has been attributed to both SA- and JA-related defense pathways and following our previous results, we assessed whether preexposure to methyl salicylate (MeSA) or methyl jasmonate (MeJA) contributed to smaller lesions in leaves of infected wheat seedlings. Remarkably, while seedlings that had been preexposed to MeSA exhibited significantly smaller lesions at 48 hai ($-29\%, P < 0.05$) and 72 hai ($-21\%, P < 0.05$) compared with the control seedlings (Fig. 1B), preexposure to MeJA led to significantly longer lesions at 48 hai ($+29\%, P < 0.05$) and 72 hai ($+53\%, P < 0.05$), showing enhanced susceptibility (Fig. 1B). However, because our previous observations showed an induction of JA biosynthesis genes at 24 hai, we verified whether treating the seedlings with MeJA at 24 hai would lead to enhanced defense. At 24 and 48 hai, there were no significant differences between the treatments (Fig. 3). Nonetheless, at 72 hai, the treatment with MeJA led to lower lesion length. Additionally, lesion length for the seedlings that had been preexposed to Z-3-HAC was lower ($-57\%, P < 0.05$) than that of nonpreexposed seedlings treated with MeJA ($-20\%, P < 0.05$; Fig. 3).

In conjunction with our gene expression results, these observations support an important role of both SA and JA in plant defense against *F. graminearum*, with SA mainly contributing to resistance during the pathogen’s early biotrophic growth and JA conditioning plant immunity during later stages of infection.

![Figure 2](image_url). Expression profiles of PAL, ICS, LOX1, and LOX2 at 24 and 48 h after challenge with a conidia suspension of *F. graminearum*. Data represent means of four biological replicates, each consisting of four pooled leaf sheaths. Error bars represent SE. Significant differences between the treatments per time point ($P < 0.05$) are depicted with asterisks.
Gene Expression of Preexposed Seedlings after Treatment with MeSA or MeJA

To elucidate whether the preexposure of Z-3-HAC leads to a direct activation of plant innate immunity or rather primes for an enhanced defense response following pathogen attack, we first investigated the impact of Z-3-HAC preexposure on MeSA- and MeJA-inducible gene expression.

We selected PAL and ICS as marker genes for the salicylate pathway (Ding et al., 2011) and LOX1 and LOX2 for the jasmonate pathway (Feng et al., 2010). Additionally, we selected different plant defense genes encoding pathogenesis-related proteins, which are known to play a role in the defense against infection with F. graminearum, namely, the pathogenesis-related proteins PR1 (Makandar et al., 2012), PR2, β-1,3-glucanase (Gao et al., 2013), PR4, shown to possess antifungal properties against Fusarium spp. (Bertini et al., 2009), and PR5 (thaumatin-like protein; Gao et al., 2013). As F. graminearum is known to interfere with the redox state of plant cells through the action of the mycotoxin DON, we also analyzed the expression of PEROXIDASE (PEROX) and NADPH OXIDASE (NADPHox; Desmond et al., 2008). Because cell wall reinforcement is a component of plant defense against fungal pathogens, we selected CINNAMOYL COENZYME A REDUCTASE3 (CCR3) and CINNAMYL ALCOHOL DEHYDROGENASE1 (CAD1) as marker genes for lignin biosynthesis (Bi et al., 2011).

Quantitative PCR analysis of the above-mentioned genes revealed that there were no significant differences between the control treatment and the Z-3-HAC-pretreated seedlings in both bioassays (Figs. 3 and 4), suggesting that Z-3-HAC does not function as a direct activator of plant defense.

After treatment with MeSA, compared with the control treatment, we observed a significant up-regulation of PR1, PR4, and PR5 in both preexposed and nonpreexposed seedlings at both time points (Fig. 4). Additionally, PEROX showed a significant up-regulation in the nonpreexposed seedlings at 24 hai (20-fold, \( P < 0.05 \)). Interestingly, at 24 h after challenge with MeSA, preexposed seedlings showed a significantly lower up-regulation of PR4 (877-fold versus 1,799-fold, \( P < 0.05 \)) and PR5 (35-fold versus 68-fold, \( P < 0.05 \)) compared with the nonpreexposed seedlings (Fig. 4). As SA- and JA-regulated defense have mainly been reported to act antagonistically (Glazebrook, 2005), we investigated whether a similar or opposite trend was present in preexposed and control seedlings after treatment with MeJA. MeJA treatment did not result in significant differences in the expression of PR1 between the different treatments, but at 24 hai, MeJA did induce a significantly up-regulation of PR4 (877-fold versus 1,799-fold, \( P < 0.05 \)) and PR5 (35-fold versus 68-fold, \( P < 0.05 \)) compared with the nonpreexposed seedlings (Fig. 4). As SA- and JA-regulated defense have mainly been reported to act antagonistically (Glazebrook, 2005), we investigated whether a similar or opposite trend was present in preexposed and control seedlings after treatment with MeJA. MeJA treatment did not result in significant differences in the expression of PR1 between the different treatments, but at 24 hai, MeJA did induce a significantly up-regulation of PR4 (877-fold versus 1,799-fold, \( P < 0.05 \)) and PEROX in the preexposed treatment compared with the control (Fig. 4). Contrary to treatment with MeSA, treatment with MeJA resulted in a significantly stronger up-regulation in the preexposed seedlings compared with the nonpreexposed seedlings for PR4 (46-fold versus 12-fold, \( P < 0.05 \)), PR5 (294-fold versus 17-fold,
For both the MeSA and MeJA treatments, PR2, NADPHox, CCR3, and CAD1 were not significantly induced (data not shown). Additionally, expression of the biosynthesis genes PAL and ICS was not affected by MeSA or MeJA treatment, suggesting that SA biosynthesis was not affected by these compounds.

Figure 4. Expression profiles of PR1, PR4, PR5, and PEROX at 24 and 48 h after challenge with MeSA or MeJA. Data represent means of three biological replicates, each consisting of four pooled leaf sheaths. Error bars represent ± se. Different letters per time point indicate significant differences between the treatments (P < 0.05).
Expression of Defense Genes upon Infection with *F. graminearum*

In order to elucidate whether the delay in disease progression of *F. graminearum* in the primed seedlings can be attributed to a stronger activation of JA-associated defense mechanisms, we analyzed the expression of the defense genes (Fig. 5) upon infection with *F. graminearum*.

Both primed and nonprimed seedlings showed a stronger up-regulation of PR1, PR4, PR5, and PEROX. Consistent with our results from the MeSA experiment, at 24 hai, we saw a higher up-regulation in the nonprimed treatment compared with the primed treatment of PR1 (30-fold versus 10-fold, \( P < 0.05 \)), PR4 (109-fold versus 37-fold, \( P = 0.059 \)), PR5 (90-fold versus 33-fold, \( P < 0.05 \)), and PEROX (9-fold versus 5-fold, \( P = 0.056 \); Fig. 5). At 48 hai, the expression pattern followed a similar trend as in MeJA-challenged seedlings at 24 hai. Namely, the expression of PR4 (10-fold versus 1-fold, \( P < 0.05 \)) and PR5 (5-fold versus 2-fold, \( P < 0.05 \)) was significantly higher in the primed treatment compared with the nonprimed treatment (Fig. 5). PR2, NADPHox, CCR3, and CAD1 were not significantly induced after infection with *F. graminearum* (data not shown).

DON Analysis

Owing to its hemibiotrophic character, *F. graminearum* is able to switch from a biotrophic to a necrotrophic lifestyle through the production of DON. As we intended to clarify if the enhanced defense through priming resulted in a higher production of DON, we quantified DON content using ultra-high-performance liquid chromatography-mass spectrometry (U-HPLC-MS).

At 24 hai, DON was present in two out of four biological repeats of the nonprimed treatment (23.4 pg mg\(^{-1}\) plant dry weight), whereas in the nonprimed treatment, DON was present in two out of four biological repeats (51.5 pg mg\(^{-1}\) plant dry weight; Fig. 6). Remarkably, at 48 hai, DON content was significantly higher in the primed seedlings than in the nonprimed seedlings (7,838.4 versus 324.5 pg mg\(^{-1}\) plant dry weight, \( P < 0.05 \)).

DISCUSSION

**Exposure to Z-3-HAC Resulted in Enhanced Resistance against Infection by *F. graminearum***

Priming by GLVs has already been shown for maize (Engelberth et al., 2004), lima bean (*Phaseolus lunatus*; Heil and Kost, 2006), poplar (*Populus* spp.; Frost et al., 2008), and tomato (Finiti et al., 2014). However, despite the fact that wheat is one of the most produced cereals in the world, no studies exist that investigate priming by GLVs in wheat. Additionally, the above-mentioned studies mainly investigated the effect of priming by GLVs on defense against herbivores, while little research has been done on the potential of GLVs to prime crops against fungal pathogens. Therefore, expanding our knowledge on priming in a plant-fungus interaction is of paramount importance to obtain a better understanding of the mechanisms and potential of plant priming by GLVs. To our knowledge, this study is the first to investigate whether GLVs can act as a priming agent for wheat against a fungal infection. We found that exposure of wheat to the GLV Z-3-HAC caused wheat seedlings and ears to become primed against a subsequent infection of *F. graminearum*. We observed for the primed wheat plants a delay in infection in both ears (Fig. 1A) and seedlings (Fig. 1B) as well as lower fungal biomass accumulation (Fig. 1D). As *F. graminearum* is a fungus that mainly thrives on the floral organs of wheat instead of the leaves (Walter et al., 2010), it can be expected that it is better adapted to resist plant defense in spikelets compared with the leaf sheaths, possibly explaining the larger effect of priming we saw in the seedlings compared with the spikelets.

**SA and JA Contribute to Defense against *F. graminearum***

The resistance of wheat against infection by *F. graminearum* has primarily been attributed to SA-mediated (Makandar et al., 2012) and JA-mediated (Li and Yen, 2008; Qi et al., 2012) defense pathways. Because of the hemibiotrophic lifestyle of *F. graminearum*, it can be expected that SA and JA will play a sequential role in the plant defense against *F. graminearum*. This biphasic defense response has already been shown by Ding et al. (2011), who reported a peak in SA content followed by a peak in JA content. At the gene level, this was confirmed in our study. At 24 hai, biosynthesis genes of the SA pathway were more strongly up-regulated, indicating an activation of SA signaling (Fig. 2). However, at 48 hai, the up-regulation of SA biosynthesis genes was diminished, and *ICS* expression was even down-regulated. This coincided with a stronger up-regulation at 48 hai of *LOX1* and *LOX2*, which are indicators of JA biosynthesis. Additionally, preexposure to MeSA resulted in smaller lesions, while preexposure to MeJA exacerbated the infection by *F. graminearum* (Fig. 1B), suggesting that during the early infection stages, defense is regulated mainly by SA. The negative effect of MeJA preexposure on disease development may be attributed to a negative cross talk between the SA and JA pathways (Robert-Seilaniantz et al., 2011; Pieterse et al., 2012), in which activation of the JA defense pathway would suppress the SA defense pathway, which is critical in the early defense response against *F. graminearum*. The importance of SA-related defense contradicts the study of Li and Yen (2008), which found at 24 hai a significant up-regulation of *LOX* and *ALLENE OXIDE SYNTHASE (AOS)*, an enzyme more upstream of JA biosynthesis. In addition, they reported an up-regulation of ethylene-responsive genes that led them to conclude that FHB tolerance in wheat is primarily mediated by JA and ethylene signaling, while SA mediated resistance is insignificant. However, it should be remarked here that, as the above-mentioned study investigated gene expression at...
24 hai, it is possible that, at this time point, *F. graminearum* had already entered into the necrotrophic phase of its life cycle (Walter et al., 2010), which consequently would result in augmented JA-related defense of the plant. This was confirmed in our experiment. The addition of exogenous MeJA 24 hai rendered the seedlings more resistant (Fig. 3). These data stress the importance of timing in studying the mechanism of plant-pathogen interaction, especially if the pathogen has a hemibiotrophic character. The importance of the jasmonate-mediated defense against infection by *Fusarium verticilloides* also has been shown for another monocot, maize. Christensen et al. (2014) characterized the

**Figure 5.** Expression profiles of PAL, ICS, LOX1, LOX2, PR1, PR4, PR5, and PEROX at 24 and 48 h after challenge with a conidia suspension of *F. graminearum*. Data represent means of four biological replicates, each consisting of four pooled leaf sheaths. Error bars represent SE. Different letters per time point indicate significant differences between the treatments (*P* < 0.05).
9-LOX gene, ZmLOX12, and showed that mutants exhibited increased susceptibility toward infection accompanied by diminished levels of JA. Our results indicate that SA plays an important role during the early stages of infection, while JA contributes to resistance during the necrotrophic stage. By precisely switching from SA- to JA-related defense at the onset of the necrotrophic phase of the pathogen, the plant can defend itself more effectively against infection by F. graminearum.

Z-3-HAC Primes for a Stronger Activation of JA-Related Defense

Priming does not directly activate costly defense mechanisms but entails a stronger plant defense upon infection (van Hulten et al., 2006; Conrath, 2009). We did not observe significant effects on gene expression by a preexposure to Z-3-HAC (Figs. 4 and 5), thus showing that Z-3-HAC did not act as a direct activator of plant defense. Furthermore, as preexposure to Z-3-HAC led to enhanced defense after treatment with MeJA or F. graminearum, we can conclude that exposure to Z-3-HAC rendered wheat seedlings in a primed state.

Defense against F. graminearum is a sequential and meticulously regulated mechanism in which the plant will consecutively employ SA- and JA-mediated defense against the biotrophic and necrotrophic phases, respectively, between which a negative cross talk exists (Glazebrook, 2005; Robert-Seilaniantz et al., 2011).

To elucidate whether the GLV Z-3-HAC targets SA- or JA-related defense, we analyzed the expression of defense genes of primed and nonprimed seedlings after challenge with MeSA and MeJA, respectively. Our results demonstrated that priming of wheat seedlings by Z-3-HAC resulted in a stronger up-regulation of PR4, PR5, and PEROX after challenge with MeJA, while expression of these genes was suppressed in the primed seedlings after challenge with MeSA, both compared with the nonprimed seedlings (Fig. 4). These results strongly suggest that Z-3-HAC promotes JA-related defense pathways but antagonizes SA-related immunity. It remains unclear whether Z-3-HAC acts as a direct repressor of SA-regulated defense or as an indirect repressor through a stronger activation of JA-related defense. Although we failed to see a priming effect of Z-3-HAC on MeJA-induced LOX1 or LOX2 expression, it cannot be ruled out that Z-3-HAC indirectly suppresses SA action by stimulating the JA pathway downstream of JA biosynthesis. Elucidating the exact mechanism(s) by which Z-3-HAC taps into the SA and JA pathways is a key challenge for future research.

A stronger defense against insects also has been shown in several studies that investigate the effect of BVOCs (Karban et al., 2006; Kessler et al., 2006; Ton et al., 2007) and GLVs in particular (Engelberth et al., 2004; Kost and Heil, 2006; Frost et al., 2008). Engelberth et al. (2007) also showed that maize plants that were preexposed to Z-3-HAC had higher production of JA and 12-oxophytodienoic acid after application with caterpillar regurgitant. This is in accordance with a similar study of Ton et al. (2007), where maize plants were exposed to the BVOCs of Spodoptera littoralis-infested plants seedlings. They reported a higher defense gene expression after these exposed seedlings were subsequently infested with S. littoralis. Similar to our study, these BVOCs contained several GLVs and

Figure 6. DON concentrations (pg mg$^{-1}$ plant dry weight) at 24 and 48 h after challenge with a conidia suspension of F. graminearum. Data represent means of four biological replicates, each consisting of six to eight pooled leaf sheaths. Error bars represent $\pm$ s. A significant difference between treatments per time point ($P < 0.05$) is depicted with an asterisk.
enhanced a specific subset of JA-inducible genes. Therefore, it is conceivable that pretreatment with GLVs confers increased resistance against insects and pathogens that are susceptible to JA-related plant defense. Hence, we want to remark here that, since it is known that most plant tissue biting-chewing insects such as caterpillars activate the JA-related defense while piercing-sucking herbivores such as aphids induce the SA-related defense (Heil, 2008; Smith et al., 2010; Liu et al., 2011), it would be interesting to investigate the priming potential of GLVs against insect herbivores with different modes of feeding.

A study by Scala et al. (2013b) investigated the mode of action of another GLV, E-2-HAL, in the defense of Arabidopsis (Arabidopsis thaliana) against the biotrophic bacterium Pseudomonas syringae. They found that plant mutants impaired in the production of GLVs were more resistant against P. syringae and exhibited higher SA and lower JA levels. They also showed that expression of MYC2 was not influenced by E-2-HAL but expression of the transcription factor AP2/ERF domain OCTADECANOID-RESPONSIVE ARABIDOPSIS (ORA59) was. Both genes are important players in the JA signaling pathway, the former contributing to defense against herbivorous insects and the latter promoting resistance to necrotrophic pathogens by integrating the ethylene and jasmonate pathways (Pieterse et al., 2012). Ethylene also plays an important role in the defense of the monocot rice against both (hemi)biotrophic and necrotrophic fungi, contributing to both increased resistance and susceptibility (De Vleesschauwer et al., 2013). More research is needed to elucidate whether Z-3-HAC also interacts with the ethylene signaling pathway or other plant defense hormones in wheat.

Nonprimed Seedlings Exhibit a Stronger Up-Regulation of Defense Genes upon F. graminearum Infection

Our finding that priming by Z-3-HAC activates a stronger JA-related defense response concurs with the gene expression of seedlings after F. graminearum infection. At 24 hai, we found a significantly stronger up-regulation of defense genes of the nonprimed seedlings compared with the primed seedlings, following the pattern after MeSA challenge. Nevertheless, primed seedlings were able to slow the infection process of F. graminearum (Fig. 1, A and B), suggesting that another mechanism was influenced by Z-3-HAC. At 48 hai, the expression pattern of PR1, PR4, PR5, and PEROX (Fig. 5) followed the expression pattern of the seedlings at 24 h after treatment with MeJA (Fig. 4). Even though at 24 hai, the expression of these defense genes was higher in the nonprimed seedlings, expression might still have been high enough to contribute to defense in both primed and nonprimed seedlings. However, as F. graminearum switched to a necrotrophic phase, the higher gene expression in primed seedlings at 48 hai might have contributed to the enhanced defense of primed seedlings we saw in our infection experiments (Fig. 1).

The molecular mechanisms for priming remain largely elusive. Recent reviews have attributed the enhanced defense to the accumulation of dormant mitogen-activated protein kinases, chromatin modifications, modifications of primary metabolism, accumulation of inactive defense metabolite conjugates, and activation of a second reactive oxygen species burst (Conrath, 2011; Pastor et al., 2013). The direct effect of BVOCs as defense signals in plant-insect interactions has already been shown in different studies (Gatehouse, 2002; Arimura et al., 2005; Kessler et al., 2006). Other studies found minor inductions of gene expression after treatment with GLVs. For example, Bate and Rothstein (1998) exposed Arabidopsis seedlings to the GLV E-2-HAL. They found induction of PAL, LOX, and AOS but no induction of PAL and PR2. Additionally, the effect of E-2-HAL was only moderate compared with treatment with the volatile MeJA. Engelberth et al. (2013) performed a microarray analysis of maize seedlings at 20 and 60 min after exposure to the GLV Z-3-HOL. They found a significant expression of genes involved in transcriptional regulation and signaling (AOS, WRKY12, and MYC7, an

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ortholog of MYC2 in Arabidopsis). Furthermore, they suggest that these early regulators serve as a main switch for the subsequent remodeling through the activation of a second-tier level of genes. Thus, these early responses might play a role in the underlying mechanism of defense priming. Contrary to Bate and Rothstein (1998), they found Z-3-HOL to be a more potent inducer of defense genes than MeJA, MeSA, and ethylene. In our study, we did not observe a significantly differential gene response between the control treatment and the wheat seedlings exposed to Z-3-HAC but not infected at the two time points (Fig. 4). This shows that overnight exposure to Z-3-HAC had no direct effect on gene expression.

**F. graminearum Produces More DON in Primed Seedlings**

Pathogens have evolved different mechanisms to evade or hijack plant defenses in order to successfully infect plant tissue. It has generally been accepted that SA-mediated defense provides protection against biotrophic pathogens and JA-mediated defense provides protection against necrotrophic pathogens (Thaler et al., 2012). However, besides SA and JA, plant defense against pathogens is regulated by an intricate network of different plant hormones between which a complicated cross talk exists (López et al., 2008; Robert-Seilaniantz et al., 2011; De Vleesschauwer et al., 2013). By manipulating this cross talk, pathogens can use the host’s own defense to their own benefit and successfully infect the plant. This phenomenon also has been described for pathogenic fungi. The necrotrophic fungus *Alternaria alternata* is known to produce different host-specific toxins and cause disease on different host plants (Ito et al., 2004). Prasad and Upadhyay (2010) showed that the toxin produced by *A. alternata* f. sp. *lycopersici* triggers the production of H2O2 and ethylene in tomato leaves. The induced production of ethylene is known to further increase the production of the mycotoxin DON. Namely, after hyphal growth in the apoplast, plants accumulate H2O2 to induce programmed cell death in order to counteract the necrotrophic phase of *Fusarium* spp. However, H2O2 acts as a signal for *F. graminearum* to produce DON, which in turn creates a positive feedback loop leading to increased H2O2 and DON production, thus successfully hijacking the plant defense system and clearing the path for the necrotrophic phase of *F. graminearum* (Walter et al., 2010; Audenaert et al., 2013). Because of the health risks associated with mycotoxins (Bennett and Klich, 2003), future research efforts should be focused on exploring the impact of enhanced plant defense on mycotoxin production by different fungi.

**CONCLUSION**

In summary, we have found that the GLV Z-3-HAC primes wheat for enhanced defense against the hemibiotrophic fungus *F. graminearum*, resulting in slower disease progress, reduced symptom development, lower fungal growth, and higher DON production in planta. Furthermore, we show that defense against *F. graminearum* is sequenially regulated by SA and JA and propose a model whereby Z-3-HAC treatment boosts JA-dependent defenses to block the pathogen during its necrotrophic growth stage.

**MATERIALS AND METHODS**

**Fusarium graminearum** and *Conidia Spore Suspension*

A GFP transformant of *F. graminearum* strain 8/1 (Jansen et al., 2005; kindly provided by Dr. Karl Heinz-Kogel) was grown on potato dextrose agar for 7 to 10 d at 20°C under a regime of 12 h of dark and 12 h of combined UVC and UVA light (2x TUV 8W T5 and 1x TL 8W BLB; Philips). Macronidia were harvested by adding a solution of 0.01% (v/v) Tween 80 to the potato dextrose agar plates and rubbing the mycelium with a Drigalski spatula. Subsequently, the suspension was diluted to a final concentration of 5 × 10⁵ conidia mL⁻¹.

**Plant Material**

Six seeds of wheat (*Triticum aestivum* var Sahara) were germinated and grown in pots (8.5-cm diameter × 6.5-cm height) in a growth chamber (18°C, 16-h-light/8-h-dark regime) for 2 weeks. When the seedlings reached wheat GS 12 (Lancashire et al., 1991), plants were selected for the leaf sheath bioassay.

**Experimental Design**

**Treatments and Cuvette System**

Throughout this study, the same methodology was used. We designate primed plants as wheat plants that have been preexposed to Z-3-HAC. Unless stated otherwise, four different treatments were used: (1) a control treatment;
(2) a priming treatment in which wheat plants were primed with Z-3-HAC; (3) a treatment in which primed wheat plants were subsequently challenged with a conidia suspension of *F. graminearum*, MeSA, or MeJA; and (4) a treatment in which nonprimed plants were challenged with a conidia suspension of *F. graminearum*, MeSA, or MeJA.

To expose wheat to Z-3-HAC, a dynamic push-pull cuvette system was used (Tholl et al., 2006). Wheat plants were placed in one of four naphalan (Foodpack) cuvettes, each of which was assigned to one of the four above-mentioned treatments. We applied 70 μL of Z-3-HAC (Sigma-Aldrich) on a piece of filter paper inside the two cuvettes that were assigned to the priming treatment (treatments 2 and 3; 5 μg). To eliminate a direct effect of Z-3-HAC on *F. graminearum*, the filter paper was removed the following day (8 AM) prior to infection. Additionally, the cuvettes were allowed to flush in order to eliminate trace amounts of Z-3-HAC. Subsequently, the plants were challenged with a suspension of *F. graminearum*, MeSA, or MeJA according to the respective treatments (treatments 3 and 4) and experiments described below.

To supply the cuvettes with sufficient air, we used a slightly modified design by Joo et al. (2010). A membrane pump (N035AN; KNF Neuberger), which was installed after a dust filter (2-μm pore size Zellfluor PTFE Membrane Filter), provided a continuous air stream of 600 mL min⁻¹ (GP40; Brooks Instrument). In order to purify the incoming air of pollutants and ozone, air passed through two active carbon filters (Airpel 10; Desotec) and an ozone filter (ETO342FC0202; Ansreco), respectively. Tubing and connections were made out of perfluoroalkoxy (Swagelok).

**Ear Infection Experiment**

To evaluate the effect of preexposure to the GLV Z-3-HAC on infection of wheat ears by *F. graminearum*, we performed an infection assay. We placed a total of nine wheat ears (var Sahara) for each treatment in cuvettes made out of naphalan (Foodpack). We applied 70 μL of Z-3-HAC (Sigma-Aldrich) on a piece of filter paper inside the cuvette that was assigned to the priming treatment and removed it the following day. Subsequently, we point inoculated three spikelets of each ear with 20 μL of a conidia suspension of 5 × 10⁵ conidia mL⁻¹. Each 2 d, we evaluated the inoculated spikelets for signs of necrosis.

**Detached Leaf Assay**

To investigate whether preexposure of wheat seedlings to Z-3-HAC will impact DON production by *F. graminearum*, DON concentration levels were measured using U-HPLC-MS based on a method described by Van Meulebroek et al. (2012). In short, 200 mg of six to eight pooled leaf sheaths was crushed using liquid nitrogen. Afterward, 1 mL of cold-modified Bieleski extraction buffer (pH 2.5) consisting of methanol, ultrapure water, and formic acid (75:20:5, v/v/v), was added. Additionally, the suspension was amended with a deuterium-labeled internal standard of 100 pg L⁻¹ d₉-absicic acid (D9OChemlin). Subsequently, the samples were vortexed and placed at −20°C for 12 h of cold extraction. The samples were centrifuged, and 500 μL of the supernatant was transferred to a 30-kD Amicon Ultra centrifugal filter unit (Merck, Millipore). The purified extract was subsequently reduced under vacuum at 35°C to one-fourth of the original volume (Gyrovap). Finally, the extract was transferred to an HPLC vial, and 10 μL was injected directly on column.

**U-HPLC-MS**

The U-HPLC-MS system consisted of an Acceela U-HPLC pumping system (Thermo Fisher Scientific) coupled to an Exactive Orbitrap mass spectrometer (Thermo Fisher Scientific) and equipped with a heated electrospray ionization source, operating in both the positive and negative modes (switching polarity mode). Chromatographic separation of the compounds was achieved with a gradient elution program using a reverse-phase Nucleodur Gravity C18 column (1.8 μm, 50-mm × 2.1-mm i.d.; Macherey-Nagel). The column oven temperature was set at 30°C. The mobile phase consisted of a binary solvent system: 0.1% (v/v) formic acid in ultrapure water (solvent A) and methanol (solvent B) at a constant flow rate of 300 μL min⁻¹. A linear gradient profile with the following proportions (v/v) of solvent A was applied: 0 to 1 min at 98%, 1 to 2.5 min from 98% to 60%, 2.5 to 4 min from 60% to 50%, 4 to 5 min from 50% to 20%, 5 to 7 min at 20%, 7 to 7.1 min from 20% to 0%, 7.1 to 8 min at 0%, and 8 to 8.01 min from 0% to 98%, followed by 2 min of reequilibration. The instrumental parameters for heated electrospray ionization can be found in Van Meulebroek et al. (2012). DON was identified based on both the retention time relative to the internal standard and the accurate mass (mass-to-charge ratio of 297.1337). After identification, concentrations were calculated by fitting the area ratios into a seven-point calibration curve set up in a leaf sheath matrix. DON was kindly provided by Dr. Marc Lemmens.

**RNA Extraction and Quantitative Reverse Transcription-PCR**

RNA from the leaf sheaths was extracted using TRI reagent (Sigma-Aldrich) according to the manufacturer’s specifications and quantified with a spectrophotometer (ND1000; Nanodrop). For each sample, four leaf sheaths were pooled to a total of three to four biological repeats. First strand complementary DNA was synthesized from 500 ng of total RNA using the SuperScript III First-Strand Synthesis Supermix kit (Life Technologies). The presence of genomic DNA was checked using gel electrophoresis. The primers used for quantitative reverse transcription (qRT)-PCR analysis are listed in Table I. qRT-PCR analysis was performed using a CFX96 system (Bio-Rad). The thermal profile consisted of an initial denaturation step for 3 min at 95°C followed by 40 cycles of 95°C for 30 s and 60°C for 60 s. Finally, melting curve analysis was performed using a temperature profile of 95°C for 10 s, cooling to 65°C for 5 s, and subsequently heating to 95°C at a rate of 0.5°C per 10 s. Fungal biomass was quantified using a pre-mRNA slicing factor of *F. graminearum* (FGSG_01244; Becher et al., 2011). Normalization of wheat defense genes was carried out using cell division control protein (Ta44277) and protein transport protein Sec23A (Ta55284) as reference genes (Paolacci et al., 2009). All calculations and analysis of the quality of the reference genes were performed using qBase+ software (Biogazelle).

**Sample Preparation**

To investigate whether preexposure of wheat seedlings to Z-3-HAC will impact DON production by *F. graminearum*, DON concentration levels were measured using U-HPLC-MS based on a method described by Van Meulebroek et al. (2012). In short, 200 mg of six to eight pooled leaf sheaths was crushed using liquid nitrogen. Afterward, 1 mL of cold-modified Bieleski extraction buffer (pH 2.5) consisting of methanol, ultrapure water, and formic acid (75:20:5, v/v/v), was added. Additionally, the suspension was amended with a deuterium-labeled internal standard of 100 pg L⁻¹ d₉-absicic acid (D9OChemlin). Subsequently, the samples were vortexed and placed at −20°C for 12 h of cold extraction. The samples were centrifuged, and 500 μL of the supernatant was transferred to a 30-kD Amicon Ultra centrifugal filter unit (Merck, Millipore). The purified extract was subsequently reduced under vacuum at 35°C to one-fourth of the original volume (Gyrovap). Finally, the extract was transferred to an HPLC vial, and 10 μL was injected directly on column.

**Data Analysis**

Gene expression data were checked for normality using the Kolmogorov-Smirnov test, and statistical comparisons between different treatments were calculated using proc mixed (SAS 9.0). Statistical differences between the primed and nonprimed treatments in the spikelet infection experiment were calculated using the χ² test (SPSS 20; IBM).

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