

RESEARCH ARTICLE

Open Access



The differential response of cold-experienced *Arabidopsis thaliana* to larval herbivory benefits an insect generalist, but not a specialist

Jana Oberländer^{1,2}, Vivien Lortzing³ , Monika Hilker³ and Reinhard Kunze^{1*}

Abstract

Background: In native environments plants frequently experience simultaneous or sequential unfavourable abiotic and biotic stresses. The plant's response to combined stresses is usually not the sum of the individual responses. Here we investigated the impact of cold on plant defense against subsequent herbivory by a generalist and specialist insect.

Results: We determined transcriptional responses of *Arabidopsis thaliana* to low temperature stress (4 °C) and subsequent larval feeding damage by the lepidopteran herbivores *Mamestra brassicae* (generalist), *Pieris brassicae* (specialist) or artificial wounding. Furthermore, we compared the performance of larvae feeding upon cold-experienced or untreated plants. Prior experience of cold strongly affected the plant's transcriptional anti-herbivore and wounding response. Feeding by *P. brassicae*, *M. brassicae* and artificial wounding induced transcriptional changes of 1975, 1695, and 2239 genes, respectively. Of these, 125, 360, and 681 genes were differentially regulated when cold preceded the tissue damage. Overall, prior experience of cold mostly reduced the transcriptional response of genes to damage. The percentage of damage-responsive genes, which showed attenuated transcriptional regulation when cold preceded the tissue damage, was highest in *M. brassicae* damaged plants (98%), intermediate in artificially damaged plants (89%), and lowest in *P. brassicae* damaged plants (69%). Consistently, the generalist *M. brassicae* performed better on cold-treated than on untreated plants, whereas the performance of the specialist *P. brassicae* did not differ.

Conclusions: The transcriptional defense response of *Arabidopsis* leaves to feeding by herbivorous insects and artificial wounding is attenuated by a prior exposure of the plant to cold. This attenuation correlates with improved performance of the generalist herbivore *M. brassicae*, but not the specialist *P. brassicae*, a herbivore of the same feeding guild.

Keywords: Plant stress response, Herbivore, Defense, Larval feeding, *Pieris brassicae*, *Mamestra brassicae*

Background

Plants have evolved a plethora of mechanisms to cope with abiotic or biotic environmental stress (e.g. [1–4]). Attack by herbivorous insects is a major threat for plants as it can lead to rapid loss of leaf material and thus reduced photosynthetic capacity, often causing severe yield and fitness loss [5–7].

Plant defense responses induced by herbivore attack represent a strategy, which is mobilized only on demand [8, 9]. Inducible defense responses are associated with transcriptional regulation of many genes and shifts in phytohormone levels. Intensively studied key regulators of wounding and herbivore defense responses are the phytohormones jasmonic acid (JA), abscisic acid (ABA), salicylic acid (SA) and ethylene (ET), which are the backbone of the plant immune signaling network [10–15]. Fine-tuning of defense responses to different herbivores is achieved by crosstalk of these signaling pathways and

* Correspondence: reinhard.kunze@fu-berlin.de

¹Freie Universität Berlin, Institute of Biology - Applied Genetics, Dahlem Centre of Plant Sciences, Albrecht-Thaer-Weg 6, 14195 Berlin, Germany
Full list of author information is available at the end of the article



may involve additional plant hormonal regulators like auxins, gibberellins, and brassinosteroids [13, 16].

In natural environments, plants are frequently exposed to simultaneously or consecutively occurring environmental stresses. Combined stresses typically provoke distinct transcriptome reprogramming and plant reactions, which are not simply due to additive effects of the single stresses [17–23]. In case of consecutively occurring environmental stress, plants can “memorize” a past stressful event and benefit from this memory by preparing themselves for a more effective response to upcoming stress. This process has been termed “priming” of a stress response by a past stress experience (reviewed in [24–27]).

Studies on priming of plant responses to insect herbivory especially focused on herbivore-related priming factors, which reliably indicate future herbivory [28]. For example, plant volatiles induced by herbivory and perceived by as yet undamaged plant tissue have been shown to serve as a reliable factor in preparing a plant for improved anti-herbivore defense [29–32]. Furthermore, insect egg depositions on leaves that indicate upcoming larval herbivory have been shown to prepare a plant for more effective defense against the hatching larvae [30]. Previous exposure of plants to herbivory-induced volatiles or to insect egg depositions are known to alter the transcriptional response to herbivory [33–39].

So far, only a few recent studies addressed the influence of an herbivory-unrelated, abiotic stress – especially drought – on the plant’s response to subsequent herbivory by including transcriptional and/or metabolic analysis (e.g. [40–42]). However, stressful conditions such as cold often precede plant attack by herbivorous insects, which usually need warm temperatures for their activities. A study by Firtzloff et al. [43] examined how exposure of *Arabidopsis thaliana* to mild cold affects plant defense against later herbivory by the specialist *Pieris brassicae*. The study showed that a significant subset of cold-regulated genes maintained altered transcript levels even after 1 day of deacclimation. Larval feeding, which started 1 day after deacclimation, induced a different transcriptome in the previously cold-exposed than in previously untreated plants and showed a weakened response of defense genes. However, larval performance of the specialist *P. brassicae* was similar on cold-experienced and untreated plants [43]. These findings are in accordance with some other studies, which also revealed that host plants with attenuated plant defense capacity did not affect the extent of feeding damage inflicted by a specialized herbivorous insect [44] nor the herbivore’s performance [45].

Generalist and specialist herbivorous insects are known to exhibit different tolerances to plant defenses [46]. However, it is unknown as yet whether they are

differentially affected by changes in plant defense that are due to prior exposure of plants to abiotic stress. Here we addressed the questions of whether a generalist insect herbivore shows different sensitivity to cold-mediated changes of feeding-induced host plant defense than a specialist, and if so, which transcriptional differences between cold-treated plants fed on by a generalist or a specialist insect may explain these ecological effects. As in our previous study [43], we used the butterflies *P. brassicae* and *Mamestra brassicae* and the Brassicacea *A. thaliana* as host plant. *Pieris brassicae* is specialized on glucosinolate-containing host plants [47], mostly from the Brassicaceae family. Like other *Pieridae* species it possesses highly specific enzymes for detoxification of the glucosinolates [48–50], which are typical secondary metabolites of the *Brassicales*. As generalist, we studied *Mamestra brassicae*, a moth whose larvae are polyphagous on over 70 plant species in 22 plant families, but exhibit a preference for *Brassica* crops [51]. In contrast to *P. brassicae*, *M. brassicae* detoxifies glucosinolates by general oxidizing enzymes (reviewed by [52]). Both lepidopteran species are active in Europe from early spring to late autumn [53, 54]. They may produce two to three generations per season until they hibernate in the soil as pupae. In the natural habitats of *M. brassicae* and *P. brassicae*, which largely overlap with that of *A. thaliana* (GBIF Secretariat: GBIF Backbone Taxonomy. Accessed via www.gbif.org/species/1920506 and www.gbif.org/species/3052436 on 01 June 2019), in spring and in autumn a succession of cold days followed by a warm period is common.

In a first approach, we compared performance of *M. brassicae* and *P. brassicae* on *A. thaliana* plants previously exposed to mild cold. We found that *M. brassicae* showed improved performance on cold-experienced plants, whereas *P. brassicae* larval performance was the same on cold-experienced and control plants, thus confirming our previous results with this latter species [43]. To elucidate the transcriptional basis of these different ecological effects, we compared the transcriptomes of cold-experienced plants exposed to feeding by the specialist, the generalist or to artificial wounding. Including the artificial wounding treatment allowed disentangling insect species-specific effects from wounding effects on the cold stress-reprogrammed plant transcriptome. We found that transcriptional responses of previously cold-exposed plants to specialist feeding, generalist feeding and artificial wounding differed.

Prior cold experience led to differential regulation of 360 *M. brassicae* feeding damage-responsive genes. In 98% of these the transcriptional response to feeding damage was attenuated. In contrast, the respective fraction of genes was smaller in artificially wounded (681 genes, 84% with attenuated response) and in *P. brassicae*

feeding-damaged plants (125 genes, 69% with attenuated response). These transcriptional changes in conjunction with the larval performance data indicate that the generalist benefits from the cold-mediated attenuation of feeding-induced gene de-regulation, whereas the specialist does not.

Results

Generalist and specialist herbivores show different performances on cold-stressed and control plants

We exposed *A. thaliana* plants to cold (4 °C) for 5 days. After a deacclimation phase (20 °C) of 1 day, larvae of the generalist *M. brassicae* and the specialist *P. brassicae* were allowed feeding upon the previously cold-experienced plants or on control plants. The weight of these larvae on previously cold-treated (Fig. 1: P + T_P and P + T_M) and untreated (Fig. 1: T_P and T_M) plants and the extent of leaf damage inflicted by the larvae were compared.

Weight gain and total weight of *P. brassicae* larvae, their leaf area consumption (Fig. 2) and the relative growth rate (RGR) of the larvae (Additional file 1: Figure S1) did not differ on previously cold-treated compared to untreated plants. In contrast, on previously cold-

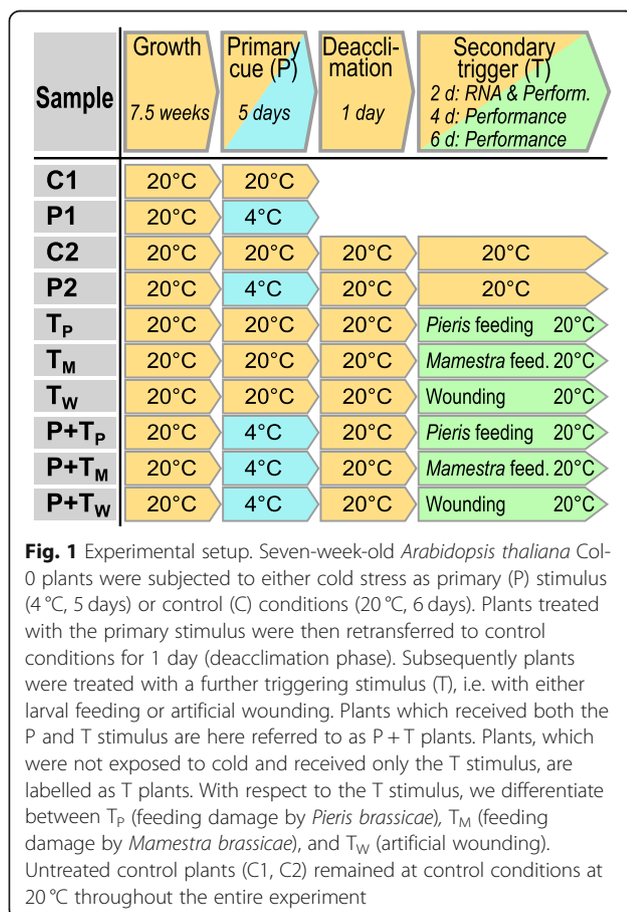
treated plants *M. brassicae* larvae consumed more leaf tissue, gained more weight and were heavier on these plants after a four- and six-day-feeding period than on untreated plants (Fig. 2). Accordingly, the RGR of the larvae was higher on cold-treated plants (Additional file 1: Figure S1). This observation suggests that the cold treatment alters either the metabolic status of the plants or their physiological reaction to leaf tissue damage in a way that is beneficial for the larval development of the generalist herbivore *M. brassicae*, but without consequences for the development of the specialist *P. brassicae*.

Transcriptional response of *Arabidopsis* to feeding damage and artificial wounding

To investigate whether *Arabidopsis* plants respond differently to leaf damage by *P. brassicae* and *M. brassicae* larvae and to artificial wounding, we analyzed the transcriptomes in leaves from plants grown at 20 °C (Fig. 1, samples T_P, T_M, T_W and C2). A Principal Component Analysis (PCA) based on gene expression values of the differently treated plants revealed that the patterns of plants exposed to *P. brassicae* feeding, *M. brassicae* feeding and artificial wounding were clearly separated from untreated control samples. However, the patterns of the treated samples partially overlapped with each other, indicating that expression of a fraction of genes is similarly regulated in the treated samples (Fig. 3a).

Overall, 1975, 1695, and 2239 differentially expressed genes (DEGs) were identified that showed ≥ 2 fold expression change after 2 days feeding damage by *P. brassicae* (T_P vs C2), *M. brassicae* (T_M vs C2) or wounding (T_W vs C2), respectively (Fig. 3b, Additional file 2: Table S1). As the majority of these genes responded qualitatively and quantitatively similarly to the three damage types (Additional file 1: Figure S2), the magnitude of the plant's transcriptional response to herbivory or artificial wounding was similar. However, feeding damage by the generalist *M. brassicae* resulted in a larger fraction of upregulated genes (62% of 1695 genes in T_M vs C2; Fig. 3b) than by the specialist *P. brassicae* (43% of 1975 genes in T_P vs C2, Fig. 3b). In total, 507 DEGs were regulated in all three sample types (central intersection in Fig. 3b). 176 DEGs specifically responded to larval feeding by either species but not to artificial wounding (Fig. 3b; intersection of T_P vs C2 and T_M vs C2 but not T_W vs C2), and 639, 700 and 767 genes were uniquely regulated upon *P. brassicae* feeding, *M. brassicae* feeding and artificial wounding, respectively. In the intersections, almost all DEGs (94–99%) were regulated in the same direction (Fig. 3b).

To disentangle common and unique regulated processes, an enrichment analysis of biological process-Gene Ontology (GO) terms was conducted (Fig. 3d).



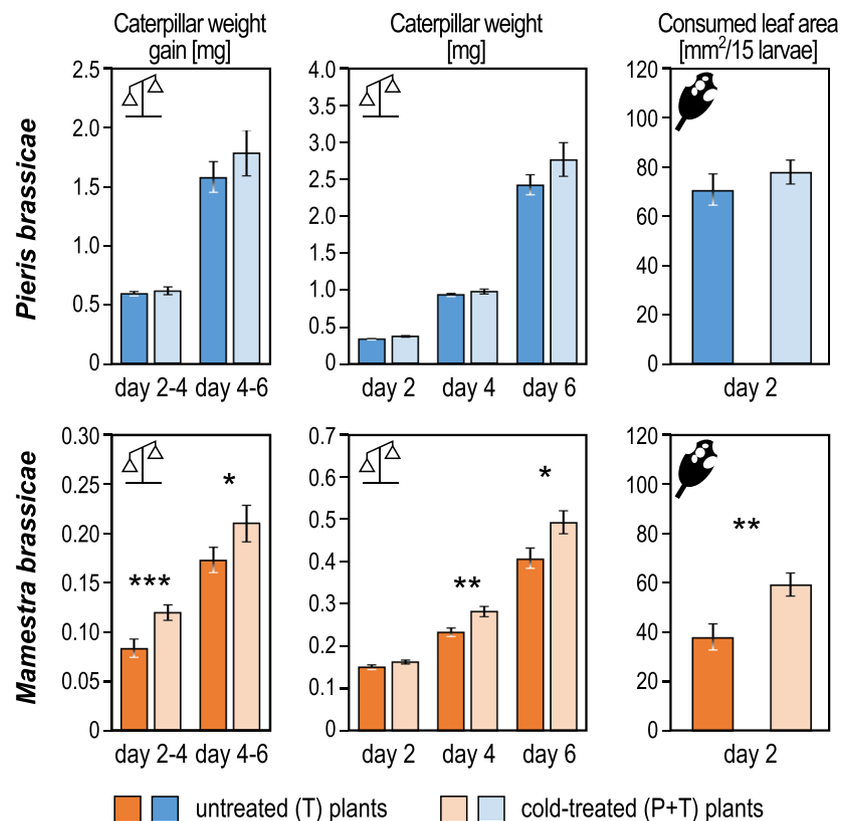


Fig. 2 Performance of *Pieris brassicae* and *Mamestra brassicae* neonate larvae after 2, 4 and 6 days feeding upon previously cold-treated or untreated plants. Larvae were placed onto plants as neonates. Measured parameters (mean values \pm SE) are caterpillar weight after 2, 4 and 6 days feeding, weight gain between day 2–4 and day 4–6, and consumed leaf area per plant (each with a group of 15 larvae) after 2 days feeding (from day 0 to day 2). Asterisks indicate significantly different values; * $P < 0.05$, ** $P < 0.01$, *** $P < 0.001$ as calculated by Student's t -tests. N (*P. brassicae*): T = 14 plants, P + T = 14 plants; N (*M. brassicae*): T = 11 plants, P + T = 11 plants

Among the 255 genes downregulated by all three damage types (Fig. 3b; central intersection), 13 GO terms were significantly enriched, which associate predominantly with responses to light, transcription, and growth. Among the 245 commonly upregulated genes, 28 GO terms were enriched, including several defense-related processes, such as response to and regulation of jasmonic acid, glucosinolate metabolism and response to insects, herbivores, bacteria, and fungi. These defense-related processes include many well described wounding- and feeding-responsive genes, i.e. *JAZ10*, *VSP1*, *VSP2*, *LOX2*, *CYP79B2*, *CYP79B3*, *IGMT1*.

Among the DEGs that were specifically responding to *P. brassicae*-feeding, two GO terms associated with abiotic stress were enriched in the 265 upregulated genes (“response to ABA”, “response to water deprivation”) and four GO terms were enriched in the downregulated genes, including the biological process “response to salicylic acid”.

Many *M. brassicae* feeding-specific upregulated genes fall into GO terms related to transcription and defense, including processes like “DNA replication initiation”,

“response to jasmonic acid” and “response to salicylic acid”, whereas the six GO terms overrepresented among *M. brassicae*-specific downregulated genes are associated with development and growth.

Artificial wounding-specific responses were overall more generic. Out of 401 upregulated genes only one GO term (“secondary metabolic process”) consisting of 19 genes was weakly enriched. The eight GO terms associated with downregulated genes ranged from protein folding to “defense response to bacterium” to “response to abiotic stress”.

Feeding by *P. brassicae* evoked only a weak upregulation of two indole-glucosinolate biosynthesis genes, *CYP79B2* and *CYP79B3*, the indole-glucosinolate O-methyltransferases *IGMT1* and *IGMT5* [55, 56] and the nitrile specifier gene *NSP3* (Additional file 2: Table S1). Feeding by *M. brassicae* induced a stronger and more complex transcriptional response in the glucosinolate pathway. In addition to the *P. brassicae*-induced genes, *MYB51*, *NSP1*, *CYP81F2*, *CYP81F4* and *IGMT2* were upregulated. Yet, the strongest effects on the glucosinolate system, upregulation of indole-glucosinolate synthesis

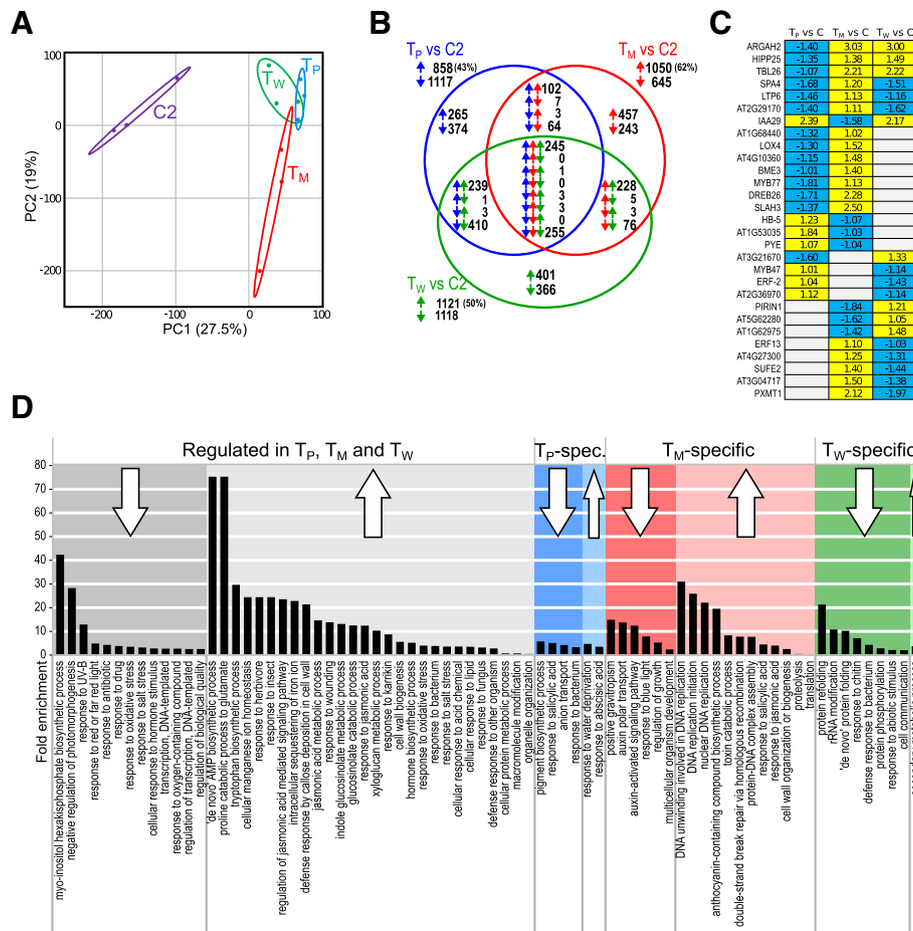


Fig. 3 Regulation of gene expression in response to herbivory or artificial wounding in *Arabidopsis thaliana* leaves compared to untreated control leaves. Plants were exposed to feeding by *P. brassicae* larvae (T_P), *M. brassicae* larvae (T_M), artificial wounding (T_W) or were left untreated (C2). Here, the T_P, T_M and T_W treatments were adjusted in such a way that we obtained comparable extent of leaf damage (about 60 mm² per plant; see Additional file 1: Figure S4). Plant material for microarray analysis was collected 2 days later. N = 3 biological replicates of each sample type. **a** Principle component analysis (PCA) of transcriptomic patterns of individual samples collected for microarray analysis. Samples originated from untreated control plants (C2, purple), feeding-damaged plants by either *P. brassicae* (T_P, red) or *M. brassicae* (T_M, blue) or artificially wounded plants (T_W, green). The first two principal components, which explain most of the changes, are depicted (explained variances are shown at the axes). Ellipses indicate the 95% confidence interval. **b** The Venn diagram shows the number of genes, which were upregulated (upwards pointing arrows) or downregulated (downwards pointing arrows) in T_P, T_M and T_W samples compared to C2 samples. **c** Heatmap depicting genes, which show opposed regulation in at least two treatments. Yellow = upregulated, blue = downregulated, grey = not regulated (log₂ fold changes). **d** Gene Ontology terms associated with commonly or uniquely up- and downregulated genes

genes and nitrile specifier protein genes and downregulation of aliphatic glucosinolate synthesis genes, was observed upon artificial wounding of the leaf.

In total, 1648 genes were responsive to at least two damage types. The vast majority of these genes were regulated in the same direction, only 29 genes were regulated in opposite directions (intersections in Fig. 3b). The genes with highest regulation differences (15- to 21-fold difference) between at least two treatments were ARGAH2, IAA29, SLAH3, DREB26, and PXMT1 (Fig. 3c). ARGAH2, one of two arginase proteins known in *Arabidopsis*, is involved in defense responses, as its expression is inducible by methyl jasmonate treatment

[57]; this gene was clearly downregulated only by *P. brassicae* feeding, but not by *M. brassicae* damage nor by artificial wounding.

JA is a major signaling molecule involved in response to wounding and defense against chewing herbivores [58–60]. Concordantly, artificially wounded leaves showed upregulation of most of the genes involved in JA biosynthesis (i.e. LOX2, AOS, AOC1 to AOC4, OPR3), JA homeostasis and turnover (i.e. JAZ2, JAZ9, JA10, IAR3, ILL6, CYP94B3) [61–64] and JA signaling (i.e. VSP1, VSP2) [58]. The JA-responsive defensin PDF1.2a [65] was upregulated as well in artificially wounded leaves. Furthermore, several JA-responsive genes

involved in biosynthesis (i.e. *CYP81F4*, *IGMT5*) [55, 66] and metabolism of glucosinolates (i.e. *PYK10*, *NSP1*) [67, 68] were upregulated.

Prior cold treatment affects the transcriptional response to tissue damage

To analyze the influence of a preceding cold stress on the transcriptional response to artificial wounding or feeding by a generalist or specialist herbivore, transcriptome analyses of leaf material from plants subjected to the treatments described in the Methods section and in Fig. 1 were performed.

A principle component analysis of gene expression values revealed a clear separation of the C2 control plant transcriptome from that of the other plant treatments, except for the transcriptome of *M. brassicae* feeding-damaged plants (T_M), whose 95% confidence interval overlapped slightly with that of the C2 control (Fig. 4a). The cold-treated plants (P2) showed a transcriptome shift relative to the C2 control, displayed in the first principle component (PC1), which accounts for ~25% of sample variances in all three sample groups. This indicates that deacclimation was not yet completed at the time of sampling. Subsequent feeding damage by *P. brassicae* ($P + T_P$) or *M. brassicae* ($P + T_M$) led to a separation of the transcriptome from that of P2 plants, whereas artificial wounding ($P + T_W$) did not. This suggests that a prior cold treatment results in a different plant transcriptional response to continuous two-day-larval feeding damage than to discontinuous artificial wounding. Moreover, the T- and P + T-induced transcriptomes differed also in a species-specific manner, indicating that *Arabidopsis* can distinguish between damage by *P. brassicae* or *M. brassicae* (Additional file 1: Figure S3).

In cold-treated plants, the total number of regulated genes ranged from 1367 in *M. brassicae*-damaged leaves to 2341 in *P. brassicae*-damaged leaves to 3293 in artificially wounded leaves relative to untreated and undamaged control plants (Fig. 4b, d; $P + T$ vs C2; the regulated genes are listed in Additional file 2: Table S1). Following prior cold stress, 446, 793, and 2439 genes were differentially regulated compared to untreated plants upon *P. brassicae* or *M. brassicae* feeding or artificial wounding, respectively (Fig. 4b, d; $P + T$ vs T). Thus, the total number of genes which were differentially expressed due to prior cold stress was higher in artificially wounded than in larval feeding-damaged plants. In general, roughly equal fractions of DEGs were up- or downregulated in $P + T_P$, $P + T_M$, and $P + T_W$ plants compared to the respective T plants.

Of particular interest are those genes that were differentially regulated in cold-treated and damaged plants relative to damaged plants ($P + T$ vs T) and also in

untreated and damaged (T vs C2) and/or cold-treated and damaged plants ($P + T$ vs C2) relative to control plants (Fig. 4b, colored intersections in Venn diagrams). These gene sets comprise 284, 490, and 1768 genes in *P. brassicae* feeding-, *M. brassicae* feeding- and wounding-damaged leaves, respectively (Additional file 2: Table S1). The 80, 270, and 465 genes in the intersection of T vs C2 and $P + T$ vs T, but not $P + T$ vs C2 (Fig. 4b, underlined numbers) were regulated by tissue damage, however, the magnitude of the transcriptional response to damage was diminished when the plants had previously experienced cold. In contrast, genes exclusively occurring in the overlapping intersection of $P + T$ vs T and $P + T$ vs C2 were regulated only upon sequential experience of cold and tissue damage by feeding or wounding, but not by damage of untreated control plants. The intersections of T vs C2, $P + T$ vs T and $P + T$ vs C2 consist of genes that respond to feeding or wounding, and this response was significantly different when plants had been exposed to a prior cold phase.

We further investigated whether genes were specifically or commonly regulated by the three cold / damage combinations. Upon prior cold treatment, 46 DEGs were commonly regulated (40 up, 6 down), i.e. their transcriptional response was independent of the insect species and type of wounding (larval feeding, artificial damage) (Fig. 4c, Additional file 2: Table S1). Additionally, 15 genes were differentially regulated (7 up, 8 down) after cold exposure and subsequent feeding damage by both herbivore species, but not after cold exposure and subsequent artificial wounding (Fig. 4c).

The prior cold treatment also affected the magnitude of the transcriptional response to subsequent tissue damage. In *Pieris*-damaged leaves, the cold pre-treatment caused a significant intensification of damage-induced up- or downregulation in 39 of the 125 damage-induced genes (31%), whereas in the remaining genes the magnitude of regulation was diminished or even turned into opposite regulation (Fig. 4e; Additional file 2: Table S1). In artificially wounded local leaves, regulation of 84% of the damage-induced genes was attenuated. In leaves damaged by *M. brassicae*, almost all (98%) feeding-induced genes exhibited attenuated regulation upon prior cold treatment. Only 2% of the feeding-induced genes exhibited intensified expression changes in cold pre-treated plants (Fig. 4e). These results show that a cold phase attenuated the transcriptional response to subsequent leaf damage in the majority of damage-induced genes. However, the degree of attenuation was dependent on the type of damage.

Leaf tissue damage affects the cold deacclimation process

To investigate whether leaf tissue damage by larval feeding and artificial wounding has an impact on gene

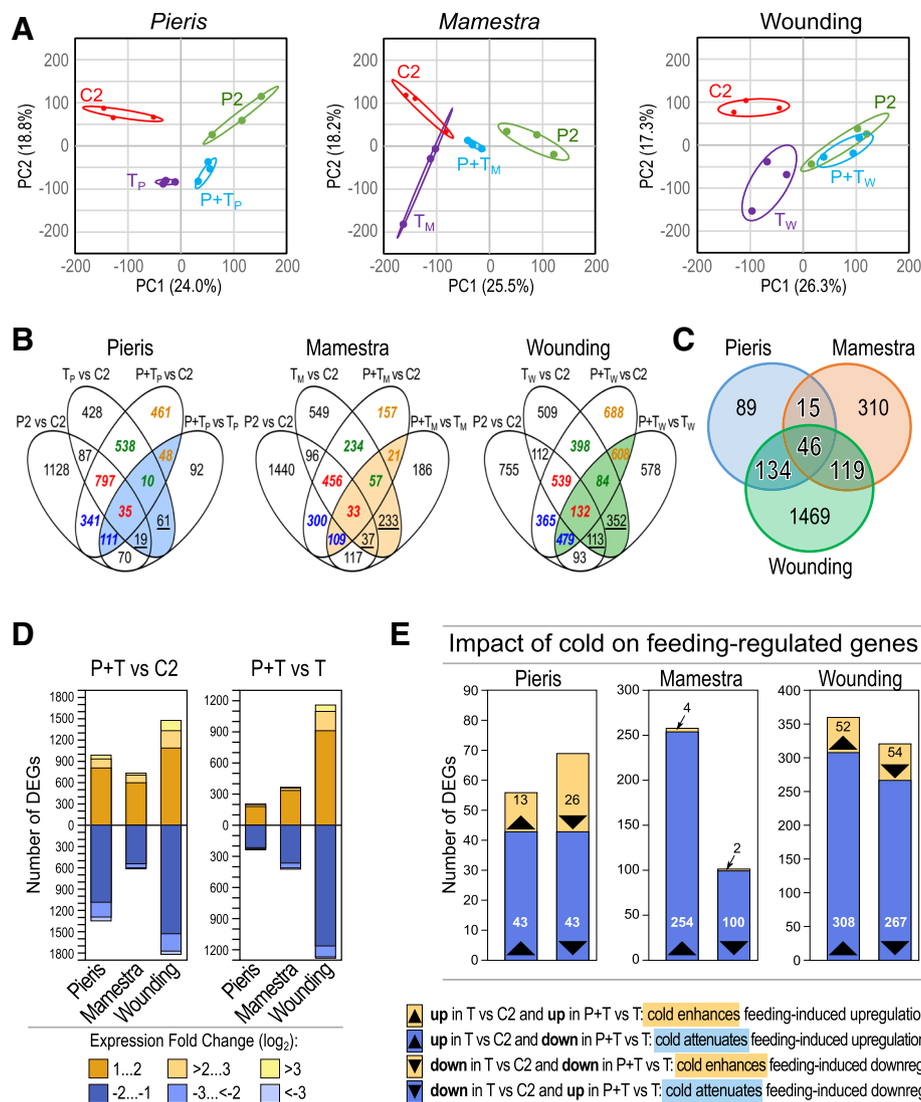


Fig. 4 Previous cold treatment alters the transcriptional response to herbivory and artificial wounding. Leaves of *A. thaliana* were untreated (C2), cold-treated (P2), damaged by *Pieris brassicae* (T_P) or *Mamestra brassicae* feeding (T_M) or artificial wounding (T_W), or cold-treated followed by feeding/wounding damage (P + T_P, P + T_M and P + T_W). N = 3 biological replicates of each sample type. **a** Principle component analysis (PCA) of the normalized gene expression of individual experimental leaf samples. The first two components, which explain most of the changes, are depicted (explained variances are shown at the axes). Ellipses indicate the 95% confidence interval. **b** Venn diagrams of genes regulated in response to larval feeding or artificial wounding with and without prior cold treatment. Blue characters, genes specifically regulated upon cold treatment; green characters, genes specifically regulated upon damage; red characters, genes regulated upon both cold per se and damage per se; orange characters, genes regulated only when the plant had been exposed to the combination of prior cold and subsequent damage; colored intersections, genes that were differentially regulated in P + T plants relative to T plants (P + T vs T) and also in untreated (T vs C2) or cold-treated, damaged plants (P + T vs C2) relative to control plants. **c** Venn diagram with the genes in the colored sectors in panel (b). **d** Numbers of differentially expressed genes (DEGs) in cold-treated and herbivore- or wounding-damaged plants compared to untreated plants (P + T vs C2; left panel) and in cold-treated herbivore- or wounding-damaged plants compared to untreated, damaged plants (P + T vs T; right panel). **e** Genes responsive to herbivory or artificial wounding with enhanced or attenuated expression changes in cold-treated relative to untreated plants

expression during deacclimation, we compared the transcriptomes of cold-treated plants during deacclimation with or without experience of tissue damage. First, we compared the transcriptome of plants at the end of the cold-period (Fig. 1; P1 plants) with that of plants after 3

days of deacclimation (Fig. 1; P2 plants). In the P2 plants we found more than 1500 newly regulated genes with 25 significantly enriched biological process GO terms, indicating that deacclimation also involves activation of cellular processes (Fig. 5a). Eleven GO terms are enriched

only for downregulated genes, nine terms only for up-regulated genes, and six terms are enriched for up- and downregulated genes (Fig. 5b). Interestingly, the down-regulated terms include the ‘glucosinolate biosynthesis process’. A closer look reveals that in this category especially genes with function in aliphatic glucosinolate biosynthesis were downregulated, like *MAM3*, *CYP79F1*,

CYP79F2, *SOT18*, *IPM11*, *IPM12* and *CYP83A1*. Noticeably, with the exception of *MAM3*, none of these genes were differentially regulated after 5 days cold in P1 plants.

It was therefore interesting to investigate how larval feeding or wounding affects this cold deacclimation response, especially with respect to the genes involved in glucosinolate biosynthesis. Overall, the regulation of 14–19% of all 2588 DEGs in deacclimating P2 plants was attenuated when feeding or wounding occurred (Fig. 5c, Additional file 2: Table S1), resulting in a faster decay of the cold deacclimation response. However, feeding damage by *P. brassicae* larvae resulted in higher expression of five of the seven above mentioned aliphatic glucosinolate biosynthesis genes (*MAM3*, *CYP79F1*, *CYP79F2*, *SOT18* and *IPM11*) in P + T_p compared to deacclimating P2 plants (expression of *IPM12* and *CYP83A1* is not altered). In contrast, feeding by *M. brassicae* larvae increased the expression of only two of the seven genes (*IPM11* and *CYP79F1*). Wounding alone did not increase the transcription level of any of the seven genes.

Stress- and stress combination-dependent transcriptional regulation of biological processes

The transcriptome analyses revealed that (i) a preceding cold phase leads to a modified transcriptional response of feeding- or wounding-regulated genes (Fig. 4e) and (ii) leaf damage by feeding or wounding modifies the transcription profile of cold-regulated genes during deacclimation (Fig. 5c). This raised the question which biological process GO terms contributed to the overall transcriptional status of P + T plants. We thus determined the enriched GO terms (Fig. 6) among the genes differentially regulated solely by cold treatment (blue characters in Figs. 4b and 6), by damage (green characters in Figs. 4b and 6), by cold or damage (red characters in Figs. 4b and 6) and by the combination of prior cold and subsequent damage (orange characters in Figs. 4b and 6), respectively. Enhanced gene regulation in many biological process GO terms was triggered almost exclusively by the single stresses cold (P2), damage (T), or the combined stressors cold+damage (P + T). Other GO terms, though, were enriched in cold exposed plants but also after damage (P2 or T). For example, leaf damage exclusively contributed to upregulation of the ‘response to JA’ process. In contrast, in the process ‘response to wounding’ some genes were induced by cold or damage, while other genes were upregulated only by damage.

Of all regulated genes in P + T plants, 13% (*Mamestra*), 22% (*Pieris*) and 39% (Wounding) only changed in expression if a cold treatment preceded the tissue damage (Fig. 4b, orange characters). These genes can be considered as primable for improved damage-triggered induction by prior cold exposure.

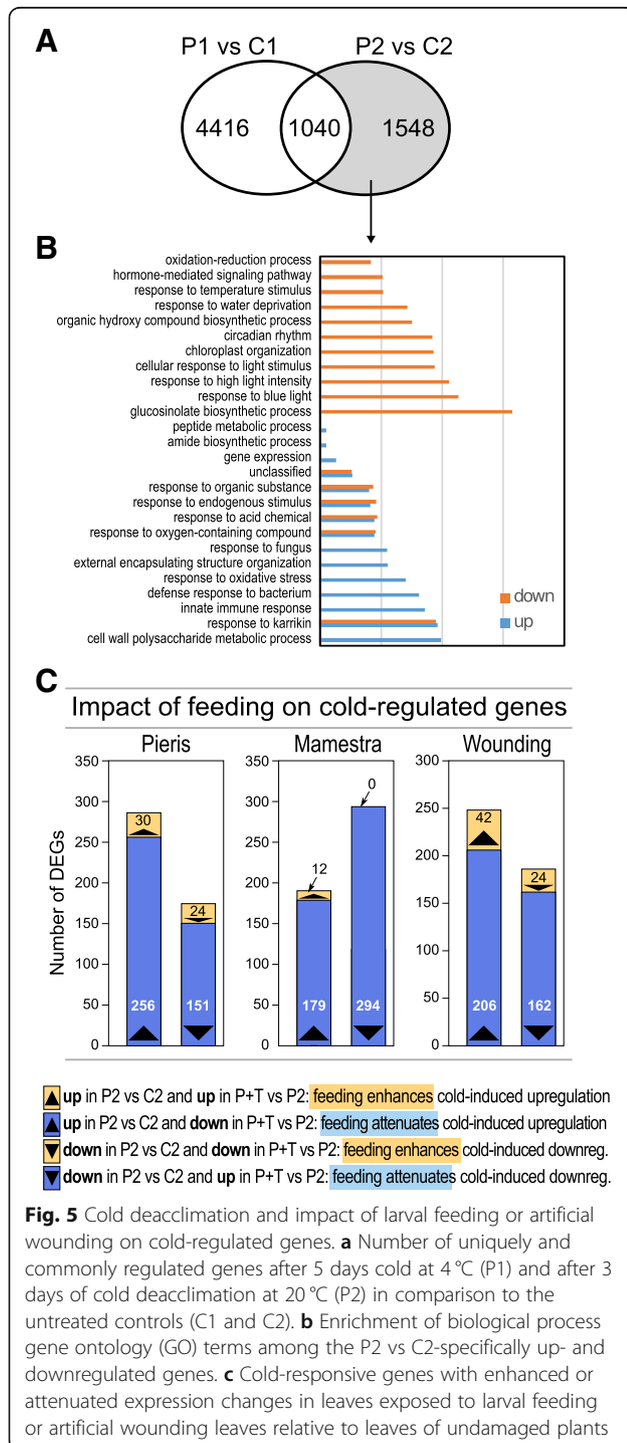




Fig. 6 Treatment-specific enrichment of biological process GO terms of regulated genes in P + T plants. Fold enrichment of biological process GO terms in P + T samples relative to C2 (P + T vs C2) samples of DEG subgroups that are solely deregulated by cold (P2; corresponding to the sectors marked with blue characters in Fig. 4b), by either cold or feeding / artificial wounding (P2 or T; corresponding to the sectors marked with red characters in Fig. 4b), by larval feeding or artificial wounding (T; corresponding to the sectors marked with green characters in Fig. 4b), or only by the combination of cold and larval feeding or artificial damage (P + T; corresponding to the sectors marked with orange characters in Fig. 4b)

Damage of *A. thaliana* leaves by *P. brassicae*, *M. brassicae* and artificial wounding resulted in significant transcriptional changes in 187, 173, and 235 defense- or glucosinolate synthesis-related genes (Additional file 2:

Table S1). Of these, 30% (*Pieris*), 43% (*Mamestra*) and 30% (Wounding) were upregulated. Remarkably, 70% of the defense- or glucosinolate synthesis-related genes upregulated by *P. brassicae* feeding were also upregulated

by *M. brassicae* feeding, whereas 75% of the genes downregulated upon *P. brassicae* feeding were not regulated by *M. brassicae* feeding. When preceded by cold, the transcriptional response of only 4% of these *P. brassicae* feeding-induced genes was attenuated by ≥ 2 -fold. In contrast, responses of much greater fractions of genes deregulated by *M. brassicae* feeding (29%) or wounding (32%) were attenuated by a factor of 2 or more, when the plants had previously been exposed to cold.

Thus, drought or cold stress preceding tissue damage apparently affects similar biological processes of *A. thaliana*, but not necessarily the same genes.

Discussion

Similarities and differences in *A. thaliana* transcriptional response to leaf damage by *P. brassicae* feeding, *M. brassicae* feeding and artificial wounding

After a prior cold treatment of *Arabidopsis* plants, larvae of the generalist herbivore *M. brassicae* performed better than on untreated plants whereas larvae of the specialist *P. brassicae* did not benefit, indicating that the cold treatment induced changes in the plant that promoted larval development of *M. brassicae*, but not of the specialist *P. brassicae*. It is conceivable that after a cold phase the plant's metabolic status or response to differences in leaf tissue damage patterns is altered.

For each of the three leaf damage scenarios approximately two thirds of all regulated genes were also regulated in one or both of the other two damage types. Remarkably, > 98% of these genes were regulated in the same direction, only 29 genes showed opposite regulation upon different damage types (Fig. 3b). Several of the genes with the largest regulation differences are known to be involved in plant responses to phytopathogens. For example, *argah2* mutants show increased susceptibility to pathogens inducing clubroot disease [69, 70]. SLAH3 is an anion channel expressed in guard cells and involved in stomatal immunity by closure of guard cells in response to pathogen attack [71]. DREB26 is responsive to infection by the necrotrophic fungus *Botrytis cinerea* and to various abiotic stresses as well [72]. PXMT1 is a target of miR163, a microRNA which promotes in a light-dependent manner seed germination and primary root length [73] and modulates defense responses against bacterial pathogens [74]. The Aux/IAA protein IAA29 is a transcription factor acting as repressor of the auxin signaling pathway [75].

Several studies addressed the hypothesis that highly specialized herbivores are more tolerant towards defenses of their host plants than generalists (reviewed by [46]). However, plant defense responses are multifaceted and not by default more effective against generalists than specialists. Thus, to identify plant responses specifically induced or suppressed by generalist and specialist

herbivores, a treatment like artificial wounding can provide a baseline for changes at the molecular level [46, 76]. Here, 23–30% of genes transcriptionally responding to leaf damage by the specialist *P. brassicae*, the generalist *M. brassicae* or artificial wounding were shared (Fig. 3b, central intersection), including upregulation of JA-responsive defense-related genes. This shows that part of the responses to feeding damage overlapped with the reaction to artificial wounding. The majority of wound-responding genes could not be assigned to a distinct, significantly regulated process, indicating a generic “panic” response of *A. thaliana* to artificial wounding [77].

Artificial wounding resulted in upregulation of many JA biosynthesis genes. Most of these genes were also upregulated to very similar levels in response to feeding by *M. brassicae* larvae, whereas transcriptional induction was attenuated or lacking upon feeding by *P. brassicae* larvae (Additional file 2: Table S1). Interestingly though, many SA-responsive genes are downregulated in response to *P. brassicae* feeding. Salicylic acid can act antagonistically to JA-mediated plant defense responses [12, 14, 60, 78], but it can also positively modulate the plant defense against herbivores [35, 79]. We found that after two days feeding by 10 *P. brassicae* larvae eight SA-associated WRKY transcription factors are downregulated. The SA-responsive factors *WRKY38*, *WRKY60* and *WRKY70* are only downregulated upon *P. brassicae* feeding, but not upon *M. brassicae* feeding or artificial wounding. It will thus be interesting to investigate whether *P. brassicae* oral secretions negatively affect the plant's SA-response pathway towards a diminished herbivore defense [80–82].

Strikingly, opposite to *P. brassicae* feeding, *M. brassicae* feeding was accompanied by more up- than downregulated SA-response genes, and seven of the eight *WRKY* genes downregulated upon *P. brassicae* feeding were not responding to *M. brassicae* feeding. It is tempting to speculate that, in contrast to *P. brassicae*, *M. brassicae* oral secretions do not dampen the plant's SA-response pathway. Moreover, *M. brassicae* feeding induced in *Arabidopsis* leaves a stronger and more complex transcriptional response of glucosinolate biosynthesis-associated genes. Striking is the upregulation of *MYB51*, a regulator of indole glucosinolate biosynthesis [83], the nitrile specifier protein *NSPI* [84], the P450 monooxygenases *CYP81F2* [85] and *CYP81F4* [66] and the indole glucosinolate methyltransferase *IGMT2* [55]. Elevated expression of plant specifier proteins has been found to promote *A. thaliana*'s defense against *P. rapae* larvae, a close relative of *P. brassicae*, as it deters *P. rapae* from egg deposition on the plants. In addition, the endoparasitoid *Cotesia rubecula*, which prefers *P. rapae* larvae as hosts, is more attracted to *P. rapae*-infested plants overexpressing specifier proteins than to

P. rapae-infested Col-0 wild-type plants. In contrast to Col-0, the specifier overexpressors accumulate mainly simple nitriles from glucosinolate hydrolysis [86]. *CYP81F2* encodes a P450 monooxygenase involved in 4MI3G (4-methoxyindol-3-ylmethylglucosinolate) synthesis and antifungal defense [85].

Studies of plant interactions with the lepidopteran generalist *Spodoptera littoralis* and with specialists (including *P. rapae* and *P. brassicae*) revealed that application of larval oral secretion of these insects results in suppression of plant defense gene expression [87, 88]. Among the genes with attenuated expression in our study the protease inhibitor DR4 and extracellular lipase 3 EXL3 showed a more pronounced attenuation upon *P. brassicae* than upon *M. brassicae* feeding (Additional file 2: Table S1). These genes are also suppressed upon feeding by *S. littoralis* [87]. It is thus conceivable that the expression attenuation we observed was caused by oral secretions of the herbivores. It is known that plants can distinguish between damage by different herbivores and by artificial wounding [89] because their oral secretions contain species-specific herbivore-associated molecular patterns (HAMPs) that enable plants to modulate their defense responses (reviewed in [90–92]). It will be interesting to investigate in the future whether the observed differences between the expression patterns upon *P. brassicae* or *M. brassicae* feeding depend on such HAMPs.

Prior low temperature exposure causes attenuated regulation of genes responsive to leaf damage

The comparison of expression changes in herbivory- or wounding-responsive genes in plants, which had previously experienced 5 days at 4 °C, revealed similarities, but also striking damage type-dependent differences in transcriptional reprogramming. Among the 46 genes that were regulated after each of the three damage types, several were reported to function in stress responses. The flavonol monooxygenase 1 (*FMO1*) is known to be essential for the establishment of systemic acquired resistance (SAR) and therefore systemic defenses against pathogens like *Pseudomonas syringae* [93]. *UGT72E2* and *UGT72E3* are involved in glucosylation of monolignols, which results in increased content of coniferin, syringin, and other phenylpropanoids [94–97]. ALLENE OXIDE CYCLASE1 (*AOC1*), a key enzyme in JA biosynthesis, is known to be rapidly responding to cold stress ([98, 99], reviewed by [100]).

Fifteen genes were differentially regulated only upon leaf damage by either of the two herbivores but not upon artificial wounding. Among the eight commonly downregulated genes is a terpene synthase (*TPS03*), which is known to be inducible by wounding and herbivory [101]. The transcription factors *RAP2.9* and *ZAT10*

function as regulators in biotic and abiotic stress responses as well as in stress combinations [102–104]. *ORA59* is involved in JA/ET synergistic regulation and important for pathogen defense via *PDFI.2* activation [15, 105]. Commonly upregulated genes (7) include *LOX5*, a member of the 9-lipoxygenases involved in pathogen defense [106] and *PIL1*, a transcription factor known to be cold- and high light-stress responsive with functions in shade avoidance. It is also JA responsive in a COI-dependent manner [107–109]. The gene *ST2A* displayed increased expression in previously cold-experienced plants responding to *P. brassicae* feeding, while the response to *M. brassicae* was opposite. *ST2A*, one of 18 sulfotransferases in *Arabidopsis*, is involved in JA metabolism by sulfating 11-OH-JA and 12-OH-JA [110].

Common for all three types of tissue damage was that a smaller fraction of damage-responsive genes was more strongly up- or downregulated, whereas in the majority of them the transcriptional response was attenuated after a prior cold treatment. The difference in the fractions of genes with altered regulation between *P. brassicae* and *M. brassicae* is striking, though. In leaves damaged by *P. brassicae* larvae 31% of the genes are more intensely and 69% more weakly regulated. In contrast, upon herbivory by *M. brassicae* larvae, only 2% of the damage-responsive genes are more intensely regulated whereas in 98% of these genes the expression change is lower than in plants that were not exposed to cold.

Feeding and wounding promote a decline of the cold acclimation status

Cold acclimation and subsequent deacclimation are known to be accompanied by extensive transcriptomic and metabolomic reorganization. Not only acclimation but also deacclimation is an active and tightly regulated process, which involves metabolic changes in lipid and cell wall components, downregulation of protein synthesis, and transcriptional reprogramming of jasmonate, brassinosteroid and other hormonal pathways [111, 112]. Pagter et al. [111] found that the deacclimation-associated responses of *A. thaliana* Col-0 proceed most rapidly during the first 12 h after shifting 4 °C-acclimated plants to 20 °C. However, deacclimation is only in part a reversion of cold acclimation, and even after 24 h the plant metabolism and transcriptome have not yet fully reverted to the non-acclimated status [111]. It is thus conceivable that after 24 h of deacclimation the plant response to herbivore attack differs from that of untreated plants, but it is not predictable whether the prior cold treatment results in an unspecific or herbivore-specific, improved or compromised defense.

Although the cold deacclimation response is considered to be rapid and mainly passive [112, 113], more than 1500 genes were newly regulated 3 days after

terminating the plant's exposure to cold. Similar results were obtained in an earlier study by Firtzloff et al. [43]. Conspicuously, among the newly regulated genes the GO term 'glucosinolate biosynthesis process' is downregulated. A weaker expression of these genes could imply a reduced aliphatic glucosinolate content in P2 plants and therefore provide advantageous conditions for the larvae of the generalist herbivore species, *M. brassicae*. Performance of this generalist species is negatively affected by aliphatic glucosinolates [114, 115]. In contrast, the specialist *P. brassicae* is well known to effectively detoxify glucosinolates (e.g. [48]).

In addition, the differences in the transcriptional response of cold-treated *Arabidopsis* plants to feeding by the two herbivores support the notion that the generalist *M. brassicae*, but not the specialist *P. brassicae*, benefits from a cold phase prior to hatching of the larvae. For instance, AOS (allene oxide synthase), a key gene in JA biosynthesis [116], the antifungal/antimicrobial defense thionin gene *THI2.1* [117], and the indolic glucosinolate synthesis genes *CYP81F4*, *CYP81F2* and *IGMT1* [55, 66, 118] were induced in plants not exposed to cold by *M. brassicae* feeding, but not by *P. brassicae* feeding. In cold-treated plants, expression of these genes was attenuated upon *M. brassicae* feeding, but not altered upon *P. brassicae* feeding. This is consistent with the observation that the performance of *P. brassicae* larvae is identical on cold-treated and control plants, whereas *M. brassicae* larvae perform better on cold-treated plants. Yet, the plant's defense response invoked by the feeding damage of *P. brassicae* larvae is comparable in untreated and cold-treated *A. thaliana* plants. Since the specialist *P. brassicae* is well adapted to the defense measures [119, 120] it was expected that its performance is not impaired.

Since *M. brassicae* is more sensitive to the defense compounds of *A. thaliana* [114], its performance in untreated plants is negatively affected. In cold-treated plants, though, the *M. brassicae* feeding damage pattern elicited an attenuated defense reaction. These results are in accordance with two other studies that addressed the question of how the experience of prior abiotic stress influences later defense responses against herbivores [42, 43]. Common results of the three studies are: (i) prior exposure of plants to abiotic stress caused a reduced transcriptional induction of tissue damage-inducible defense genes, including attenuated gene expression of e.g. JA- and glucosinolate metabolism-related genes; (ii) the performance of the specialist herbivores *P. rapae* [42] and *P. brassicae* (this study and [43]) was not affected by prior drought or cold treatment of *A. thaliana*; (iii) herbivory led to a shift from the drought- or cold-adapted transcriptome towards herbivore defense, thus accelerating the abiotic stress deacclimation. Yet, the

differentially regulated genes in feeding-damaged plants with prior drought or cold experience differed to a great extent. Only two genes, a glutathione S-transferase (*GSTU8*) and *UPF0496* were transcriptionally responding to all tissue damage types when preceded by drought or cold.

Conclusions

We show that a prior cold treatment of *A. thaliana* differentially reprogrammed the transcriptional response to leaf tissue damage by artificial wounding and feeding by the specialist herbivore *P. brassicae* or the generalist herbivore *M. brassicae*. The cold-treatment resulted at the transcriptional level in an attenuation of the plant's damage-induced defense response. We suggest that this attenuation is responsible for the improved larval performance of the generalist *M. brassicae*. In contrast, the specialist *P. brassicae* is unaffected by the damage-induced *A. thaliana* defense measures and accordingly does not benefit from the defense attenuation by a preceding cold treatment of the plants.

Methods

Plant growth

Arabidopsis thaliana Columbia Col-0 seeds (Stock No. N1093) were obtained from the Nottingham Arabidopsis Stock Centre (NASC). Seeds were sown on soil type A (2:2:1, Einheitserde CL P: Einheitserde CL T: Sand) and stratified for 2 days at 4 °C. Thereafter, plants were grown in a growth chamber at short day conditions (8 h/16 h light/dark cycle, 120 μE), 20 °C and 50% relative humidity for 7 weeks. Three-week-old seedlings were transplanted in pots containing soil type B (7:7:3, Einheitserde CL P: Einheitserde CL T: Perlite).

Insect rearing

Pieris brassicae larvae from in-house captive breeding were reared on savoy cabbage (*Brassica oleracea* convar. *Capitata* var. *sabauda*) as described by [43]. *Mamestra brassicae* were obtained from N. Fatouros (Biosystematics Group, Wageningen University and Research, Wageningen, Netherlands). Larvae were reared on cabbage plants (*Brassica oleracea* var. *sabellica* L.) until pupation. Soil was provided to last instar *M. brassicae* larvae for pupation, while *P. brassicae* pupae were kept on cardboard. Adults of *M. brassicae* were offered water and a sugar-water solution (1:5 w/v). Adult *P. brassicae* butterflies were fed with an aqueous honey solution.

Plant treatments

The experimental design is depicted in Fig. 1. Seven-week-old plants were subjected to (i) 5 days cold at 4 °C (P samples), (ii) leaf damage by *P. brassicae* larvae (T_P samples), *M. brassicae* larvae (T_M samples) or artificial

wounding (T_W samples), (iii) cold followed by leaf damage ($P + T_D$, $P + T_M$ or $P + T_W$ samples), or (iv) no stimulus (C samples). The stimulus ‘cold’ was applied for 5 days, followed by 1 day under normal growth conditions (20°C) as memory/deacclimation phase. P1 samples were taken directly after 5 days of cold and P2 samples 3 days after transferring plants back to 20°C (Fig. 1). The second stress (larval herbivory or artificial wounding) was applied for 2 days following the 1 day memory/deacclimation phase. For treatment with larvae, neonate *P. brassicae* or *M. brassicae* larvae were added in a clipcage to leaf number 17. For control, an empty clipcage was placed on leaf number 17 of untreated control (C) and cold-pretreated (P) plants. Artificial wounding was applied by damaging leaf number 17 with forceps for 30 s two times a day for 2 days. The damaged area almost matched the area of damage that larvae feeding inside a clipcage inflicted to a leaf.

Larval performance measurement

Individual seven-week-old plants treated with or without prior cold were subjected to feeding by 15 freshly hatched *M. brassicae* or *P. brassicae* larvae on leaf 17. The experiments were repeated 11 times ($N = 11$ plants) with *M. brassicae* and 15 times ($N = 15$ plants) with *P. brassicae*. Larvae were confined in clipcages with a diameter of 3 cm. Two days later, larval weight and weight gain were determined. Furthermore, the consumed leaf area was assessed by comparing pictures of the leaves taken before and after 2 days feeding using ImageJ [121]. The leaf expansion during the 2 days feeding period was marginal and not taken into account. Subsequently larvae were returned to the plants and allowed to feed upon the whole plant for another 4 days. Two and 4 days later larval weight and weight gain were measured again. Larval performance data were evaluated with “R” [122] and subjected to statistical analysis [123, 124]. Data were tested for normal distribution (Shapiro-Wilk test) and homogenous variances (Levene’s test). If larval weight and weight gain values were not normally distributed and/or did not show variance homogeneity, data were \log_2 transformed to fulfil the prerequisites for applying unpaired Student’s *t*-test.

Transcriptome analyses

We analyzed the transcriptome of untreated plants (C), cold-exposed plants (P), damaged plants (T) and cold-exposed and feeding-damaged plants (P + T). We standardized the extent of damage by insects and artificial wounding to be able to ascribe damage-induced transcriptomic changes to the type of damage rather than to the extent of damage. Therefore, plant leaves were exposed to 10 *P. brassicae* larvae or 20 *M. brassicae* larvae in a clip cage. After 2 days feeding, the leaf area

consumed by the two species was almost identical (Additional file 1: Figure S4). The artificially wounded area was similar as well. For RNA extraction, a 1 cm wide strip from leaf number 17 of C1, C2, P1, P2, T_D , T_M , T_W , $P + T_D$, $P + T_M$ and $P + T_W$ plants was harvested. The stripe was located proximal to the clipcage or wounding site. To minimize effects of circadian clock-dependent transcriptional regulation, all samples were collected 4 to 6 h after the onset of the daylight phase, i.e. at a time when larvae are actively feeding in nature. After harvesting, the strips were kept frozen in liquid nitrogen. Leaf material of three individual plants was pooled to obtain one biological replicate, and three biological replicates of each sample type were analyzed.

Frozen leaf material was ground in liquid nitrogen, and total RNA was extracted according to Onate-Sanchez [125]. Total RNA was DNase I-digested according to manufacturer’s instructions (Thermo Fisher Scientific). Yield and quality of extracted RNA was determined spectrophotometrically and by denaturing agarose gel electrophoresis.

Genome-wide expression analyses were conducted on ArrayXS Arabidopsis v2 microarrays (series XS-5010; GEO accession GPL19779; Oaklabs GmbH, Hennigsdorf, Germany). Microarray data were processed and analyzed with the *Bioconductor Linear Models* for microarray data (limma) software package [126, 127] as described in Firtzlaff et al. [43]. In short, microarray signals were background-corrected and interarray-normalized. Genes with ≥ 2 -fold expression change and adjusted *P*-values ≤ 0.05 (Benjamini and Hochberg false discovery rate procedure) were defined to be differentially expressed genes (DEGs) (Additional file 1: Figure S2). Gene expression data are deposited in the NCBI GEO repository under the accession number GSE114211.

Principle component analysis (PCA) of the transcriptomic data sets was performed using the “ggplot” and “ggbiplot” packages of “R” [122, 128]. Enriched gene ontology (GO) terms were identified using the *TAIR GO Term Enrichment for Plants* tool (www.arabidopsis.org) provided by PANTHER DB (<http://pantherdb.org>). If not mentioned otherwise, a Bonferroni correction for multiple testing was applied to reduce false positives.

Additional files

Additional file 1: Figure S1. Relative growth rates of *Pieris brassicae* and *Mamestra brassicae* neonate larvae on previously cold-treated or untreated plants. **Figure S2.** Gene expression changes in plants exposed to larval feeding or artificial wounding compared to untreated control plants. **Figure S3.** Principle component analysis of transcriptomes of plants exposed to individual treatments. **Figure S4.** Leaf area consumption by *Pieris brassicae* and *Mamestra brassicae* neonate larvae after 2 days feeding upon previously cold-treated or untreated plants. (PDF 348 kb)

Additional file 2: Table S1. List of genes differentially expressed in response to cold treatment and/or feeding by *Pieris brassicae* larvae, feeding by *Mamestra brassicae* larvae or artificial wounding. (XLSX 1843 kb)

Abbreviations

ABA: Abscisic acid; DEG: Differentially expressed gene; ET: Ethylene; GO: Gene ontology; JA: Jasmonic acid; PCA: Principal component analysis; SA: Salicylic acid

Acknowledgments

We thank Nina Fatouros for providing *Mamestra brassicae*, Charlotte Thomas for general lab support and the members of the Hilker and Kunze labs for valuable discussions.

Authors' contributions

JO, VL, MH and RK conceived and designed the study. JO performed the experiments and analyzed the data. JO, VL, MH and RK wrote the manuscript. All authors read and approved the final manuscript.

Funding

The research was funded by the Deutsche Forschungsgemeinschaft, Collaborative Research Centre 973, project B4 (www.sfb973.de). The funding body was not involved in the design of the study and collection, analysis, and interpretation of data and in writing the manuscript.

Availability of data and materials

Microarray transcription raw data are deposited in the NCBI Gene Expression Omnibus (GEO) repository under the accession number GSE114211.

Ethics approval and consent to participate

Not applicable.

Consent for publication

Not applicable.

Competing interests

The authors declare that they have no competing interests

Author details

¹Freie Universität Berlin, Institute of Biology - Applied Genetics, Dahlem Centre of Plant Sciences, Albrecht-Thaer-Weg 6, 14195 Berlin, Germany.

²Present address: University of Bern, Molecular Plant Physiology, Altenbergrain 21, CH-3013 Bern, Switzerland. ³Freie Universität Berlin, Institute of Biology - Applied Zoology / Animal Ecology, Dahlem Centre of Plant Sciences, Haderslebener Str. 9, 12163 Berlin, Germany.

Received: 16 April 2019 Accepted: 23 July 2019

Published online: 02 August 2019

References

- Walling LL. The myriad plant responses to herbivores. *J Plant Growth Regul.* 2000;19(2):195–216.
- Hirayama T, Shinozaki K. Research on plant abiotic stress responses in the post-genome era: past, present and future. *Plant J.* 2010;61(6):1041–52.
- Baxter A, Mittler R, Suzuki N. ROS as key players in plant stress signalling. *J Exp Bot.* 2014;65(5):1229–40.
- Zebelo SA, Maffei ME. Role of early signalling events in plant-insect interactions. *J Exp Bot.* 2015;66(2):435–48.
- Nabity PD, Zavala JA, DeLucia EH. Indirect suppression of photosynthesis on individual leaves by arthropod herbivory. *Ann Bot.* 2009;103(4):655–63.
- Núñez-Farfán J, Fornoni J, Valverde PL. The evolution of resistance and tolerance to herbivores. *Annu Rev Ecol Evol S.* 2007;38:541–66.
- Strauss SY, Rudgers JA, Lau JA, Irwin RE. Direct and ecological costs of resistance to herbivory. *Trends Ecol Evol.* 2002;17(6):278–85.
- Schaller A. Induced plant resistance to herbivory. Stuttgart: Springer; 2008.
- Karban R, Baldwin IT. Induced responses to herbivory. Chicago: The University of Chicago Press; 1997. 330 p.
- Howe GA, Jander G. Plant immunity to insect herbivores. *Annu Rev Plant Biol.* 2008;59:41–66.
- Koo AJ, Howe GA. The wound hormone jasmonate. *Phytochemistry.* 2009;70(13–14):1571–80.
- Pieterse CM, Leon-Reyes A, Van der Ent S, Van Wees SC. Networking by small-molecule hormones in plant immunity. *Nat Chem Biol.* 2009;5(5):308–16.
- Verhage A, van Wees SC, Pieterse CM. Plant immunity: it's the hormones talking, but what do they say? *Plant Physiol.* 2010;154(2):536–40.
- Pieterse CM, Van der Does D, Zamioudis C, Leon-Reyes A, Van Wees SC. Hormonal modulation of plant immunity. *Annu Rev Cell Dev Biol.* 2012;28:489–521.
- Wasternack C, Hause B. Jasmonates: biosynthesis, perception, signal transduction and action in plant stress response, growth and development. An update to the 2007 review in *annals of botany.* *Ann Bot.* 2013;111(6):1021–58.
- Diezel C, von Dahl CC, Gaquerel E, Baldwin IT. Different lepidopteran elicitors account for cross-talk in herbivory-induced phytohormone signaling. *Plant Physiol.* 2009;150(3):1576–86.
- Mittler R. Abiotic stress, the field environment and stress combination. *Trends Plant Sci.* 2006;11(1):15–9.
- Prasch CM, Sonnewald U. Signaling events in plants: stress factors in combination change the picture. *Environ Exp Bot.* 2015;114:4–14.
- Stam JM, Kroes A, Li Y, Gols R, van Loon JJ, Poelman EH, et al. Plant interactions with multiple insect herbivores: from community to genes. *Annu Rev Plant Biol.* 2014;65:689–713.
- Atkinson NJ, Urwin PE. The interaction of plant biotic and abiotic stresses: from genes to the field. *J Exp Bot.* 2012;63(10):3523–43.
- Ramegowda V, Senthil-Kumar M. The interactive effects of simultaneous biotic and abiotic stresses on plants: mechanistic understanding from drought and pathogen combination. *J Plant Physiol.* 2015;176:47–54.
- Suzuki N, Rivero RM, Shulaev V, Blumwald E, Mittler R. Abiotic and biotic stress combinations. *New Phytol.* 2014;203(1):32–43.
- Voelckel C, Baldwin IT. Herbivore-induced plant vaccination. Part II. Array-studies reveal the transience of herbivore-specific transcriptional imprints and a distinct imprint from stress combinations. *Plant J.* 2004;38(4):650–63.
- Hilker M, Schwachtje J, Baier M, Balazadeh S, Baurle I, Geiselhardt S, et al. Priming and memory of stress responses in organisms lacking a nervous system. *Biol Rev Camb Philos Soc.* 2016;91(4):1118–33.
- Conrath U, Beckers GJ, Langenbach CJ, Jaskiewicz MR. Priming for enhanced defense. *Annu Rev Phytopathol.* 2015;53:97–119.
- Martinez-Medina A, Flors V, Heil M, Mauch-Mani B, Pieterse CM, Pozo MJ, et al. Recognizing plant defense priming. *Trends Plant Sci.* 2016;21(10):818–22.
- Pastor V, Luna E, Mauch-Mani B, Ton J, Flors V. Primed plants do not forget. *Environ Exp Bot.* 2013;94:46–56.
- Mescher MC, De Moraes CM. Plant biology: pass the ammunition. *Nature.* 2014;510(7504):221–2.
- Heil M, Ton J. Long-distance signalling in plant defence. *Trends Plant Sci.* 2008;13(6):264–72.
- Hilker M, Fatouros NE. Plant responses to insect egg deposition. *Annu Rev Entomol.* 2015;60:493–515.
- Hilker M, Fatouros NE. Resisting the onset of herbivore attack: plants perceive and respond to insect eggs. *Curr Opin Plant Biol.* 2016;32:9–16.
- Karban R, Yang LH, Edwards KF. Volatile communication between plants that affects herbivory: a meta-analysis. *Ecol Lett.* 2014;17(1):44–52.
- Altmann S, Muino JM, Lortzing V, Brandt R, Himmelbach A, Altschmied L, et al. Transcriptomic basis for reinforcement of elm antiherbivore defence mediated by insect egg deposition. *Mol Ecol.* 2018;27(23):4901–15.
- Geuss D, Stelzer S, Lortzing T, Steppuhn A. *Solanum dulcamara's* response to eggs of an insect herbivore comprises ovidical hydrogen peroxide production. *Plant Cell Environ.* 2017;40(11):2663–77.
- Lortzing V, Oberländer J, Lortzing T, Tohge T, Steppuhn A, Kunze R, et al. Insect egg deposition renders plant defence against hatching larvae more effective in a salicylic acid-dependent manner. *Plant Cell Environ.* 2019;42:1019–32.
- Büchel K, McDowell E, Nelson W, Descour A, Gershenson J, Hilker M, et al. An elm EST database for identifying leaf beetle egg-induced defense genes. *BMC Genomics.* 2012;13:242.
- Engelberth J, Contreras CF, Dalvi C, Li T, Engelberth M. Early transcriptome analyses of Z-3-hexenol-treated *Zea mays* revealed distinct transcriptional networks and anti-herbivore defense potential of green leaf volatiles. *PLoS One.* 2013;8(10):e77465.
- Ton J, D'Alessandro M, Jourdie V, Jakob G, Karlen D, Held M, et al. Priming by airborne signals boosts direct and indirect resistance in maize. *Plant J.* 2007;49(1):16–26.

39. Bonnet C, Lassueur S, Ponzio C, Gols R, Dicke M, Reymond P. Combined biotic stresses trigger similar transcriptomic responses but contrasting resistance against a chewing herbivore in *Brassica nigra*. *BMC Plant Biol*. 2017;17(1):127.
40. Coolen S, Proietti S, Hickman R, Davila Olivas NH, Huang PP, Van Verk MC, et al. Transcriptome dynamics of *Arabidopsis* during sequential biotic and abiotic stresses. *Plant J*. 2016;86(3):249–67.
41. Weldegergis BT, Zhu F, Poelman EH, Dicke M. Drought stress affects plant metabolites and herbivore preference but not host location by its parasitoids. *Oecologia*. 2015;177(3):701–13.
42. Davila Olivas NH, Coolen S, Huang P, Severing E, van Verk MC, Hickman R, et al. Effect of prior drought and pathogen stress on *Arabidopsis* transcriptome changes to caterpillar herbivory. *New Phytol*. 2016;210(4):1344–56.
43. Firtzlaff V, Oberländer J, Geiselhardt S, Hilker M, Kunze R. Pre-exposure of *Arabidopsis* to the abiotic or biotic environmental stimuli "chilling" or "insect eggs" exhibits different transcriptomic responses to herbivory. *Sci Rep*. 2016;6:28544.
44. Stotz HU, Pittendrigh BR, Kroymann J, Weniger K, Fritsche J, Bauke A, et al. Induced plant defense responses against chewing insects. Ethylene signaling reduces resistance of *Arabidopsis* against Egyptian cotton worm but not diamondback moth. *Plant Physiol*. 2000;124(3):1007–18.
45. Müller R, de Vos M, Sun JY, Sønderby IE, Halkier BA, Wittstock U, et al. Differential effects of indole and aliphatic glucosinolates on lepidopteran herbivores. *J Chem Ecol*. 2010;36(8):905–13.
46. Ali JG, Agrawal AA. Specialist versus generalist insect herbivores and plant defense. *Trends Plant Sci*. 2012;17(5):293–302.
47. Feltwell J. The large white butterfly: the biology, biochemistry and physiology of *Pieris brassicae* (Linnaeus). The Hague: Dr. W. Junk Publishers; 1982.
48. Wittstock U, Agerbirk N, Stauber EJ, Olsen CE, Hippler M, Mitchell-Olds T, et al. Successful herbivore attack due to metabolic diversion of a plant chemical defense. *Proc Natl Acad Sci U S A*. 2004;101(14):4859–64.
49. Kumar R, Bhardwaj U, Kumar P, Mazumdar-Leighton S. Midgut serine proteases and alternative host plant utilization in *Pieris brassicae* L. *Front Physiol*. 2015;6:95.
50. Stauber EJ, Kuczka P, van Ohlen M, Vogt B, Janowitz T, Piotrowski M, et al. Turning the 'mustard oil bomb' into a 'cyanide bomb': aromatic glucosinolate metabolism in a specialist insect herbivore. *PLoS One*. 2012;7(4):e35545.
51. Rojas JC, Wyatt TD, Birch MC. Flight and oviposition behavior toward different host plant species by the cabbage moth, *Mamestra brassicae* (L.) (Lepidoptera : Noctuidae). *J Insect Behav*. 2000;13(2):247–54.
52. Hopkins RJ, van Dam NM, van Loon JJ. Role of glucosinolates in insect-plant relationships and multitrophic interactions. *Annu Rev Entomol*. 2009;54:57–83.
53. Grüner C, Sauer KP. Aestival dormancy in the cabbage moth *Mamestra brassicae* L. (Lepidoptera: Noctuidae) : 1. Adaptive significance of variability of two traits: Day length thresholds triggering aestival dormancy and duration of aestival dormancy. *Oecologia*. 1988;74(4):515–23.
54. Held C, Spieth HR. First evidence of pupal summer diapause in *Pieris brassicae* L.: the evolution of local adaptedness. *J Insect Physiol*. 1999;45(6):587–98.
55. Pfalz M, Mikkelsen MD, Bednarek P, Olsen CE, Halkier BA, Kroymann J. Metabolic engineering in *Nicotiana benthamiana* reveals key enzyme functions in *Arabidopsis* indole glucosinolate modification. *Plant Cell*. 2011;23(2):716–29.
56. Pfalz M, Mukhaimar M, Perreau F, Kirk J, Hansen CI, Olsen CE, et al. Methyl transfer in glucosinolate biosynthesis mediated by indole glucosinolate O-methyltransferase 5. *Plant Physiol*. 2016;172(4):2190–203.
57. Brownfield DL, Todd CD, Deyholos MK. Analysis of *Arabidopsis* arginase gene transcription patterns indicates specific biological functions for recently diverged paralogs. *Plant Mol Biol*. 2008;67(4):429–40.
58. Zhang L, Zhang F, Melotto M, Yao J, He SY. Jasmonate signaling and manipulation by pathogens and insects. *J Exp Bot*. 2017;68(6):1371–85.
59. Campos ML, Kang JH, Howe GA. Jasmonate-triggered plant immunity. *J Chem Ecol*. 2014;40(7):657–75.
60. Erb M, Meldau S, Howe GA. Role of phytohormones in insect-specific plant reactions. *Trends Plant Sci*. 2012;17(5):250–9.
61. Widemann E, Miesch L, Lugan R, Holder E, Heinrich C, Aubert Y, et al. The amidohydrolases IAR3 and ILL6 contribute to jasmonoyl-isoleucine hormone turnover and generate 12-hydroxyjasmonic acid upon wounding in *Arabidopsis* leaves. *J Biol Chem*. 2013;288(44):31701–14.
62. Heitz T, Widemann E, Lugan R, Miesch L, Ullmann P, Desaubry L, et al. Cytochromes P450 CYP94C1 and CYP94B3 catalyze two successive oxidation steps of plant hormone jasmonoyl-isoleucine for catabolic turnover. *J Biol Chem*. 2012;287(9):6296–306.
63. Kitaoka N, Matsubara T, Sato M, Takahashi K, Wakuta S, Kawaide H, et al. *Arabidopsis* CYP94B3 encodes jasmonoyl-L-isoleucine 12-hydroxylase, a key enzyme in the oxidative catabolism of jasmonate. *Plant Cell Physiol*. 2011;52(10):1757–65.
64. Koo AJ, Cooke TF, Howe GA. Cytochrome P450 CYP94B3 mediates catabolism and inactivation of the plant hormone jasmonoyl-L-isoleucine. *Proc Natl Acad Sci U S A*. 2011;108(22):9298–303.
65. Penninckx IA, Thomma BP, Buchala A, Mettraux JP, Broekaert WF. Concomitant activation of jasmonate and ethylene response pathways is required for induction of a plant defensin gene in *Arabidopsis*. *Plant Cell*. 1998;10(12):2103–13.
66. Kai K, Takahashi H, Saga H, Ogawa T, Kanaya S, Ohta D. Metabolomic characterization of the possible involvement of a cytochrome P450, CYP81F4, in the biosynthesis of indolic glucosinolate in *Arabidopsis*. *Plant Biotechnology*. 2011;28(4):379–85.
67. Nakano RT, Pislewska-Bednarek M, Yamada K, Edger PP, Miyahara M, Kondo M, et al. PYK10 myrosinase reveals a functional coordination between endoplasmic reticulum bodies and glucosinolates in *Arabidopsis thaliana*. *Plant J*. 2017;89(2):204–20.
68. Wittstock U, Meier K, Dorr F, Ravindran BM. NSP-dependent simple nitrile formation dominates upon breakdown of major aliphatic glucosinolates in roots, seeds, and seedlings of *Arabidopsis thaliana* Columbia-0. *Front Plant Sci*. 2016;7:1821.
69. Gravot A, Deleu C, Wagner G, Lariagon C, Lugan R, Todd C, et al. Arginase induction represses gall development during clubroot infection in *Arabidopsis*. *Plant Cell Physiol*. 2012;53(5):901–11.
70. Brauc S, De Vooght E, Claeys M, Geuns JM, Hofte M, Angenon G. Overexpression of arginase in *Arabidopsis thaliana* influences defence responses against *Botrytis cinerea*. *Plant Biol (Stuttg)*. 2012;14(Suppl 1):39–45.
71. Zheng X, Kang S, Jing Y, Ren Z, Li L, Zhou JM, et al. Danger-associated peptides close stomata by OST1-independent activation of anion channels in guard cells. *Plant Cell*. 2018;30(5):1132–46.
72. Sham A, Moustafa K, Al-Ameri S, Al-Azzawi A, Iratni R, AbuQamar S. Identification of *Arabidopsis* candidate genes in response to biotic and abiotic stresses using comparative microarrays. *PLoS One*. 2015;10(5):e0125666.
73. Chung PJ, Park BS, Wang H, Liu J, Jang IC, Chua NH. Light-inducible miR163 targets PXM1 transcripts to promote seed germination and primary root elongation in *Arabidopsis*. *Plant Physiol*. 2016;170(3):1772–82.
74. Chow HT, Ng DW. Regulation of miR163 and its targets in defense against *Pseudomonas syringae* in *Arabidopsis thaliana*. *Sci Rep*. 2017;7:46433.
75. Liscum E, Reed JW. Genetics of aux/IAA and ARF action in plant growth and development. *Plant Mol Biol*. 2002;49(3–4):387–400.
76. Lortzing T, Firtzlaff V, Nguyen D, Rieu I, Stelzer S, Schad M, et al. Transcriptomic responses of *Solanum dulcamara* to natural and simulated herbivory. *Mol Ecol Resour*. 2017;17(6):e196–211.
77. Avramova Z. The jasmonic acid-signalling and abscisic acid-signalling pathways cross talk during one, but not repeated, dehydration stress: a non-specific 'panicky' or a meaningful response? *Plant Cell Environ*. 2017;40(9):1704–10.
78. Thaler JS, Humphrey PT, Whiteman NK. Evolution of jasmonate and salicylate signal crosstalk. *Trends Plant Sci*. 2012;17(5):260–70.
79. Rostas M, Winter TR, Borkowski L, Zeier J. Copper and herbivory lead to priming and synergism in phytohormones and plant volatiles in the absence of salicylate-jasmonate antagonism. *Plant Signal Behav*. 2013;8(6):e24264.
80. Chung SH, Rosa C, Scully ED, Peiffer M, Tooker JF, Hoover K, et al. Herbivore exploits orally secreted bacteria to suppress plant defenses. *Proc Natl Acad Sci U S A*. 2013;110(39):15728–33.
81. Acevedo FE, Smith P, Peiffer M, Helms A, Tooker J, Felton GW. Phytohormones in fall armyworm saliva modulate defense responses in plants. *J Chem Ecol*. 2019; <https://doi.org/10.1007/s10886-019-01079-z>.
82. Mason CJ, Jones AG, Felton GW. Co-option of microbial associates by insects and their impact on plant-herbivore interactions. *Plant Cell Environ*. 2019;42(3):1078–86.
83. Gogolashvili T, Berger B, Mock HP, Müller C, Weisshaar B, Flügge UI. The transcription factor HIG1/MYB51 regulates indolic glucosinolate biosynthesis in *Arabidopsis thaliana*. *Plant J*. 2007;50(5):886–901.
84. Burow M, Losansky A, Muller R, Plock A, Kliebenstein DJ, Wittstock U. The genetic basis of constitutive and herbivore-induced ESP-independent nitrile formation in *Arabidopsis*. *Plant Physiol*. 2009;149(1):561–74.

85. Bednarek P, Pislewska-Bednarek M, Svatos A, Schneider B, Doudsky J, Mansurova M, et al. A glucosinolate metabolism pathway in living plant cells mediates broad-spectrum antifungal defense. *Science*. 2009;323(5910):101–6.
86. Mumm R, Burow M, Bukovinskine'kiss G, Kazantzidou E, Wittstock U, Dicke M, et al. Formation of simple nitriles upon glucosinolate hydrolysis affects direct and indirect defense against the specialist herbivore, *Pieris rapae*. *J Chem Ecol*. 2008;34(10):1311–21.
87. Consales F, Schweizer F, Erb M, Gouhier-Darimont C, Bodenhausen N, Bruessow F, et al. Insect oral secretions suppress wound-induced responses in *Arabidopsis*. *J Exp Bot*. 2012;63(2):727–37.
88. Reymond P, Bodenhausen N, Van Poecke RM, Krishnamurthy V, Dicke M, Farmer EE. A conserved transcript pattern in response to a specialist and a generalist herbivore. *Plant Cell*. 2004;16(11):3132–47.
89. Shinya T, Hojo Y, Desaki Y, Christeller JT, Okada K, Shibuya N, et al. Modulation of plant defense responses to herbivores by simultaneous recognition of different herbivore-associated elicitors in rice. *Sci Rep*. 2016;6:32537.
90. Mithöfer A, Boland W. Recognition of herbivory-associated molecular patterns. *Plant Physiol*. 2008;146(3):825–31.
91. Maffei ME, Arimura G, Mithofer A. Natural elicitors, effectors and modulators of plant responses. *Nat Prod Rep*. 2012;29(11):1288–303.
92. Basu S, Varsani S, Louis J. Altering plant defenses: herbivore-associated molecular patterns and effector arsenal of chewing herbivores. *Mol Plant-Microbe Interact*. 2018;31(1):13–21.
93. Mishina TE, Zeier J. The *Arabidopsis* flavin-dependent monooxygenase FMO1 is an essential component of biologically induced systemic acquired resistance. *Plant Physiol*. 2006;141(4):1666–75.
94. König S, Feussner K, Kaefer A, Landesfeind M, Thurow C, Karlovsky P, et al. Soluble phenylpropanoids are involved in the defense response of *Arabidopsis* against *Verticillium longisporum*. *New Phytol*. 2014;202(3):823–37.
95. Lanot A, Hodge D, Jackson RG, George GL, Elias L, Lim EK, et al. The glucosyltransferase UGT72E2 is responsible for monolignol 4-O-glucoside production in *Arabidopsis thaliana*. *Plant J*. 2006;48(2):286–95.
96. Miao YC, Liu CJ. ATP-binding cassette-like transporters are involved in the transport of lignin precursors across plasma and vacuolar membranes. *Proc Natl Acad Sci U S A*. 2010;107(52):22728–33.
97. Zhao Q, Nakashima J, Chen F, Yin Y, Fu C, Yun J, et al. Laccase is necessary and nonredundant with peroxidase for lignin polymerization during vascular development in *Arabidopsis*. *Plant Cell*. 2013;25(10):3976–87.
98. Hannah MA, Heyer AG, Hinch DK. A global survey of gene regulation during cold acclimation in *Arabidopsis thaliana*. *PLoS Genet*. 2005;1(2):e26.
99. Hu Y, Jiang L, Wang F, Yu D. Jasmonate regulates the INDUCER OF CBF EXPRESSION-C-REPEAT BINDING FACTOR/DRE BINDING FACTOR1 cascade and freezing tolerance in *Arabidopsis*. *Plant Cell*. 2013;25(8):2907–24.
100. Kazan K. Diverse roles of jasmonates and ethylene in abiotic stress tolerance. *Trends Plant Sci*. 2015;20(4):219–29.
101. Huang M, Abel C, Sohrabi R, Petri J, Haupt I, Cosimano J, et al. Variation of herbivore-induced volatile terpenes among *Arabidopsis* ecotypes depends on allelic differences and subcellular targeting of two terpene synthases, TPS02 and TPS03. *Plant Physiol*. 2010;153(3):1293–310.
102. Mittler R, Kim Y, Song L, Couto J, Couto A, Ciftci-Yilmaz S, et al. Gain- and loss-of-function mutations in Zat10 enhance the tolerance of plants to abiotic stress. *FEBS Lett*. 2006;580(28–29):6537–42.
103. Prasch CM, Sonnewald U. Simultaneous application of heat, drought, and virus to *Arabidopsis* plants reveals significant shifts in signaling networks. *Plant Physiol*. 2013;162(4):1849–66.
104. Tsutsui T, Kato W, Asada Y, Sako K, Sato T, Sonoda Y, et al. DEAR1, a transcriptional repressor of DREB protein that mediates plant defense and freezing stress responses in *Arabidopsis*. *J Plant Res*. 2009;122(6):633–43.
105. Pre M, Atallah M, Champion A, De Vos M, Pieterse CM, Memelink J. The AP2/ERF domain transcription factor ORA59 integrates jasmonic acid and ethylene signals in plant defense. *Plant Physiol*. 2008;147(3):1347–57.
106. Vicente J, Cascon T, Vicedo B, Garcia-Agustin P, Hamberg M, Castresana C. Role of 9-lipoxygenase and alpha-dioxygenase oxylipin pathways as modulators of local and systemic defense. *Mol Plant*. 2012;5(4):914–28.
107. Ciolfi A, Sessa G, Sassi M, Possenti M, Salvucci S, Carabelli M, et al. Dynamics of the shade-avoidance response in *Arabidopsis*. *Plant Physiol*. 2013;163(1):331–53.
108. Rasmussen S, Barah P, Suarez-Rodriguez MC, Bressendorff S, Friis P, Costantino P, et al. Transcriptome responses to combinations of stresses in *Arabidopsis*. *Plant Physiol*. 2013;161(4):1783–94.
109. Robson F, Okamoto H, Patrick E, Harris SR, Wasternack C, Brearley C, et al. Jasmonate and phytochrome a signaling in *Arabidopsis* wound and shade responses are integrated through JAZ1 stability. *Plant Cell*. 2010;22(4):1143–60.
110. Gidda SK, Miersch O, Levitin A, Schmidt J, Wasternack C, Varin L. Biochemical and molecular characterization of a hydroxyjasmonate sulfotransferase from *Arabidopsis thaliana*. *J Biol Chem*. 2003;278(20):17895–900.
111. Pagter M, Alpers J, Erban A, Kopka J, Zuther E, Hinch DK. Rapid transcriptional and metabolic regulation of the deacclimation process in cold acclimated *Arabidopsis thaliana*. *BMC Genomics*. 2017;18(1):731.
112. Zuther E, Juszczyk I, Lee YP, Baier M, Hinch DK. Time-dependent deacclimation after cold acclimation in *Arabidopsis thaliana* accessions. *Sci Rep*. 2015;5:12199.
113. Byun YJ, Koo MY, Joo HJ, Ha-Lee YM, Lee DH. Comparative analysis of gene expression under cold acclimation, deacclimation and reacclimation in *Arabidopsis*. *Physiol Plantarum*. 2014;152(2):256–74.
114. Beekwilder J, van Leeuwen W, van Dam NM, Bertossi M, Grandi V, Mizzi L, et al. The impact of the absence of aliphatic glucosinolates on insect herbivory in *Arabidopsis*. *PLoS One*. 2008;3(4):e2068.
115. Jeschke V, Kearney EE, Schramm K, Kunert G, Shekhov A, Gershenzon J, et al. How glucosinolates affect generalist lepidopteran larvae: growth, development and glucosinolate metabolism. *Front Plant Sci*. 2017;8:1995.
116. Schaller F. Enzymes of the biosynthesis of octadecanoid-derived signalling molecules. *J Exp Bot*. 2001;52(354):11–23.
117. Eppe P, Apel K, Bohlmann H. Overexpression of an endogenous thionin enhances resistance of *Arabidopsis* against *Fusarium oxysporum*. *Plant Cell*. 1997;9(4):509–20.
118. Pfalz M, Vogel H, Kroymann J. The gene controlling the indole glucosinolate modifier1 quantitative trait locus alters indole glucosinolate structures and aphid resistance in *Arabidopsis*. *Plant Cell*. 2009;21(3):985–99.
119. Smallegange RC, van Loon JJ, Blatt SE, Harvey JA, Agerbirk N, Dicke M. Flower vs. leaf feeding by *Pieris brassicae*: glucosinolate-rich flower tissues are preferred and sustain higher growth rate. *J Chem Ecol*. 2007;33(10):1831–44.
120. Schweizer F, Heidel-Fischer H, Vogel H, Reymond P. *Arabidopsis* glucosinolates trigger a contrasting transcriptomic response in a generalist and a specialist herbivore. *Insect Biochem Mol Biol*. 2017;85:21–31.
121. Schneider CA, Rasband WS, Eliceiri KW. NIH image to ImageJ: 25 years of image analysis. *Nat Methods*. 2012;9(7):671–5.
122. R Development Core Team. R : A language and environment for statistical computing. ed. Vienna University of Economics and Business. Vienna: The R Foundation for Statistical Computing; 2015. Retrieved from <http://www.R-project.org>.
123. Revelle W. Psych: procedures for personality and psychological research. Evanston: Northwestern University; 2018. <https://CRAN.R-project.org/package=psych>. Version = 1.8.12.
124. Fox J, Weisberg S. An R companion to applied Regression, Third Edition. Thousand Oaks: Sage; 2019. <https://socialsciences.mcmaster.ca/jfox/Books/Companion/>.
125. Onate-Sanchez L, Vicente-Carbajosa J. DNA-free RNA isolation protocols for *Arabidopsis thaliana*, including seeds and siliques. *BMC Res Notes*. 2008;1:93.
126. Ritchie ME, Phipson B, Wu D, Hu Y, Law CW, Shi W, et al. *limma* powers differential expression analyses for RNA-sequencing and microarray studies. *Nucleic Acids Res*. 2015;43(7):e47.
127. Smyth GK. Linear models and empirical bayes methods for assessing differential expression in microarray experiments. *Stat Appl Genet Mol Biol*. 2004;3(1):Article3.
128. Wickham H. ggplot2: elegant graphics for data analysis. Gentleman R, Hornik K, Parmigiani G, editors. New York: Springer; 2009. 213 p.

Publisher's Note

Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.