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Identification and characterisation of hypomethylated DNA loci controlling quantitative resistance in Arabidopsis

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27 Abstract

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Variation in DNA methylation enables plants to inherit traits independently of changes to DNA 29 sequence. Here, we have screened an Arabidopsis population of epigenetic recombinant inbred 30 lines (epiRILs) for resistance against Hyaloperonospora arabidopsidis (Hpa). These lines 31 share the same genetic background, but show variation in heritable patterns of DNA 32 methylation. We identified 4 epigenetic quantitative trait loci (epiQTLs) that provide 33 34 quantitative resistance without reducing plant growth or resistance to other (a)biotic stresses. Phenotypic characterisation and RNA-sequencing analysis revealed that *Hpa*-resistant epiRILs 35 are primed to activate defence responses at the relatively early stages of infection. Collectively, 36 our results show that hypomethylation at selected pericentromeric regions is sufficient to 37 provide quantitative disease resistance, which is associated with genome-wide priming of 38 defence-related genes. Based on comparisons of global gene expression and DNA methylation 39 between the wild-type and resistant epiRILs, we discuss mechanisms by which the 40 pericentromeric epiQTLs could regulate the defence-related transcriptome. 41

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57 Introduction

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59 Eukaryotic cytosine methylation plays an important role in the regulation of gene expression 60 and genome stability. In plants, this form of DNA methylation occurs at three sequence contexts: CG, CHG and CHH, where H indicates any base except guanine (G)^{1,2}. Patterns of 61 plant DNA methylation in the plant genome can remain stable over multiple generations and 62 influence heritable phenotypes³. Recent evidence has suggested that reduced DNA methylation 63 increases the responsiveness of the plant immune system ⁴ This 'priming' of plant defence 64 enables an augmented induction of defence-related genes after pathogen attack, causing 65 increased levels of quantitative resistance⁵⁻⁸. In some cases, priming of defence-related genes 66 is associated with post-translational histone modifications that mark a more open chromatin 67 68 structure^{9,10}. Additional evidence for epigenetic regulation of plant immunity has come from independent studies reporting that disease-exposed Arabidopsis produces progeny that 69 expresses transgenerational acquired resistance (TAR), which is associated with priming of 70 defence-related genes^{10,11}. Furthermore, Arabidopsis mutants that are impaired in the 71 establishment or maintenance of DNA methylation mimic TAR-related priming without prior 72 priming stimulus¹²⁻¹⁴. By contrast, the hyper-methylated *ros1-4* mutant, which is impaired in 73 74 active DNA de-methytation, is more susceptible to biotrophic pathogens, affected in defence gene responsiveness, and impaired in TAR^{14,15}. Thus, DNA (de)methylation determines 75 quantitative disease resistance by influencing the responsiveness of defence-related genes. 76 However, causal evidence that selected hypomethylated DNA loci are responsible for the 77 78 meiotic transmission of this form of quantitative disease resistance is lacking.

79 Epigenetic Recombinant Inbred Lines (epiRILs) have been developed with the aim to study the epigenetic basis of heritable plant traits^{16,17}. EpiRILs show little differences in DNA 80 sequence, but vary substantially in DNA methylation. A commonly used population of epiRILs 81 is derived from a cross between the Arabidopsis wild-type (Wt) accession Col-0 and the 82 decreased DNA methylation1-2 (ddm1-2) mutant¹⁷. The DDM1 protein is a chromatin re-83 modelling enzyme that provides DNA methyltransferase enzymes access to heterochromatic 84 transposable elements $(TEs)^{18-20}$. Accordingly, the ddm1-2 mutation causes loss of 85 pericentromeric heterochromatin and reduced DNA methylation in all sequence contexts^{21,22}. 86 Although the epiRILs from the *ddm1-2* x Col-0 cross do not carry the *ddm1-2* mutation, they 87 contain stably inherited hypomethylated DNA regions from the ddm1-2 parent, which are 88 maintained up to 16 generations of self-pollination^{17,23,24}. A core set of 123 epiRILs from this 89

population at the 8th generation of self-pollination in the wild-type (Wt) background has been
characterized for differentially methylated region (DMR) markers, enabling linkage mapping
of heritable hypomethylated loci controlling root growth, flowering and abiotic stress
tolerance^{8,25,26}.

In this study, we have characterised the core set of 123 lines from the *ddm1-2* x Col-0 94 epiRIL population for resistance against the biotrophic downy mildew pathogen 95 Hyaloperonospora arabidopsidis (Hpa) to search for heritable hypomethylated loci controlling 96 disease resistance. We identified 4 of these epigenetic quantitative trait loci (epiQTLs), 97 accounting for 60% of the variation in disease resistance. None of these epiQTLs were 98 associated with growth impairment, indicating that the resistance does not incur major 99 physiological costs on plant development. Further phenotypic characterisation and 100 101 transcriptome analysis of selected Hpa-resistant epiRILs revealed that their resistance is associated with genome-wide priming of defence-related genes. Interestingly, bisulfite 102 103 sequencing did not reveal defence regulatory genes inside the epiQTL regions that were simultaneously primed and hypomethylated, suggesting that DDM1-dependent DNA 104 methylation at the epiQTLs trans-regulates the responsiveness of distant defence genes. 105

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107 **Results**

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109 Identification of epiQTLs controlling quantitative resistance against *Hpa*.

To examine the role of DDM1-dependent DNA methylation in heritable disease resistance, 123 110 epiRILs from the *ddm1-2* x Col-0 cross were analysed for *Hpa* resistance and compared to 111 siblings of the *ddm1-2* parent (Figure 1a, red), the Wt parent (Col-0), and five progenies thereof 112 113 (Figure 1a, green). Leaves of three-week old plants were inoculated with Hpa conidiospores and then collected for trypan-blue staining at six days post inoculation (dpi). Microscopic 114 classification of leaves into 4 classes of *Hpa* colonisation (Figure 1-figure supplement 1) 115 revealed 51 epiRILs with statistically enhanced levels of resistance compared to each 116 susceptible Wt line (Pearson's Chi-squared tests, p < 0.05). Of these, 8 epiRILs showed similar 117 levels of Hpa resistance as the ddm1-2 line (Figure 1a, dark blue triangles; Pearson's Chi-118 squared test, p>0.05), whereas 43 epiRILs showed intermediate levels of resistance. To identify 119 the epiQTL(s) responsible for the observed variation in Hpa resistance, the categorical 120 classification of *Hpa* infection was converted into a single value numerical resistance index 121

(RI; Figure 1a, bottom graph). Using a linkage map of stably inherited DMR markers²³ 122 (Supplementary dataset S1), interval mapping revealed 4 statistically significant epiQTLs on 123 chromosomes I, II, IV and V (Figure 1b). The epiQTL on chromosome II had the highest 124 logarithm of odds (LOD) value. For all epiQTLs, the DMR markers with the highest LOD 125 scores ('peak markers') showed a positive correlation between *ddm1-2* haplotype and RI 126 (Figure 1c), indicating that the hypomethylated haplotype from ddm1-2 increases resistance 127 against *Hpa*. A linear regression model to calculate the percentage of RI variance explained by 128 each peak marker $(R^2(g))^{25}$ confirmed that the DMR peak marker of the epiQTL on 129 chromosome II had the strongest contribution to RI variation. Using an additive model, the 130 combined contribution of all epiQTL peak markers to RI variation $(R^2(G))^{25}$ was estimated at 131 60.0% (Figure 1d). 132

133 DNA methylation maintains genome stability by preventing transposition of TEs. In the Col-0 x ddml-2 epiRIL population, reduced methylation at the ddml-2 haplotype occurs 134 predominantly at long transposons in heterochromatic pericentromeric regions^{20,23}. Frequent 135 transposition events in the epiRILs are nevertheless rare as most DNA hypomethylation occurs 136 at relic transposons that have lost the ability to transpose, and the occurrence of independent 137 transposition events at similar loci is extremely unlikely^{23,27}. However, it is possible that 138 transposition events originating from the heavily hypomethylated *ddm1-2* parent were crossed 139 into the population, resulting into shared transposition events (STEs) between multiple 140 epiRILs, which could have contributed to variation in resistance. To account for this possibility, 141 we compared the genomic DNA sequences of the 4 epiQTL intervals from 122 epiRILs (LOD 142 drop-off = 2) for the presence of STEs in more than two epiRILs, using TE-tracker software²⁸. 143 This analysis revealed three STEs in the epiQTL interval on chromosome I (Supplementary 144 dataset S2), while no STEs could be detected in the other epiQTL intervals. None of the STEs 145 in the epiQTL on chromosome I showed statistically significant linkage with RI 146 (Supplementary dataset S2). Accordingly, we conclude that the segregating *Hpa* resistance in 147 the epiRIL population is caused by epigenetic variation in DNA methylation, rather than 148 genetic variation by STEs. 149

Effects of the resistance epiQTLs on plant growth and resistance against other (a)biotic stresses.

Expression of inducible defence mechanisms is often associated with physiological costs, 153 resulting in reduced plant growth²⁹. To determine whether the resistance that is controlled by 154 the 4 epiQTLs is associated with costs to plant growth, we quantified the green leaf area (GLA) 155 of 12-15 individual plants per line at the stage of *Hpa* inoculation (Figure 1-figure supplement 156 2). Subsequent interval mapping revealed one statistically significant epiQTL on chromosome 157 I (Figure 1b). The corresponding peak marker (MM150) showed a negative correlation between 158 159 GLA and ddm1-2 haplotype (Figure 1c), indicating that the hypomethylated ddm1-2 allele at this locus represses plant growth. The growth epiQTL mapped to a different region than the 160 resistance epiQTL on chromosome I (Figure 1b, inset). Furthermore, none of the 8 most 161 resistant epiRILs showed significant growth reduction compared to all Wt lines in the screen 162 (Figure 1-figure supplement 2). Hence, the resistance provided by the 4 hypomethylated 163 164 epiQTLs is not associated with major physiological costs to plant growth.

Enhanced defence to one stress can lead to enhanced susceptibility to another stress, 165 which is caused by antagonistic cross-talk between defence signalling pathways³⁰. To examine 166 whether Hpa resistance in the epiRIL population is associated with increased susceptibility to 167 other stresses, we compared the 8 most Hpa-resistant epiRILs (Figure 1a; Figure 1-figure 168 supplement 3a) for resistance against the necrotrophic fungus *Plectosphaerella cucumerina* 169 (Pc) and tolerance to salt (NaCl). At nine dpi with Pc spores, epiRIL#193 showed a statistically 170 significant reduction in necrotic lesion size compared to the Wt (line #602), indicating 171 enhanced resistance (Figure 1-figure supplement 3b). The seven other epiRILs showed 172 unaffected levels of Pc resistance that were similar to the Wt. Salt tolerance was quantified by 173 the percentage of seedlings with fully developed cotyledons at six days after germination on 174 agar medium with increasing NaCl concentrations. Remarkably, all Hpa-resistant epiRILs 175 showed varying degrees of tolerance to the highest NaCl concentration compared to Wt plants 176 177 (Figure 1-figure supplement 3c). Thus, the quantitative resistance to Hpa in the epiRIL population does not compromise resistance against necrotrophic pathogens or abiotic stress. 178

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180 *Hpa*-resistant epiRILs are primed to activate different defence mechanisms.

181 Basal resistance against *Hpa* involves a combination of salicylic acid (SA)-dependent and SA-182 independent defence mechanisms^{31,32}. To examine the role of SA-dependent defences, we

profiled the expression of the SA-inducible marker gene PR1 at 48 and 72 hours post 183 inoculation (hpi), which represents a critical time-window for host defence against $Hpa^{33,34}$. 184 None of the epiRILs showed a statistically significant increase in basal PR1 expression after 185 mock inoculation (Figure 2a; Figure 1-figure supplement 4a), indicating that the resistance is 186 not based on constitutive up-regulation of SA-dependent defence signalling. However, in 187 comparison to the Wt line, epiRILs #71, #148, #193, #229 and #508 showed augmented 188 induction of *PR1* at 48 and/or 72 hpi with *Hpa* (Figure 2a; Figure 1-figure supplement 4a), 189 indicating priming of SA-inducible defences⁷. To assess the role of cell wall defence, all lines 190 191 were analysed for effectiveness of callose deposition, which is a pathogen-inducible defence mechanism that is largely controlled by SA-independent signalling³⁵. Compared to the Wt line, 192 all but one epiRIL (#193) showed a statistically significant increase in the proportion of callose-193 arrested germ tubes (Figure 2a; Figure 1-figure supplement 4b). Hence, the 8 most Hpa-194 resistant epiRILs are primed to activate differentially regulated defence responses, which 195 explains the lack of major costs on growth and compatibility with other types of (a)biotic stress 196 resistance in the epiRILs (Figures 1b and 2a; Figure 1-figure supplements 2-4). 197

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199 Transgenerational stability of the resistance.

The 123 epiRILs analysed for Hpa resistance had been self-pollinated for 8 generations in a 200 Wt (Col-0) genetic background since the F1 x Col-0 backcross (F9)¹⁷. To examine the 201 transgenerational stability of the resistance phenotype over one more generation, 5 individuals 202 from the 8 most resistant epiRILs and the Wt line (Figure 1a, Figure 1-figure supplement 3a) 203 204 were selected to generate F10 families, which were then tested for Hpa resistance. Comparing distributions of pooled leaves from all five families per line confirmed that each epiRIL 205 maintained a statistically enhanced level of resistance (Figure 1-figure supplement 5; Pearson's 206 Chi-squared test, p < 0.05; top asterisks). However, when comparing individual F10 families to 207 the Wt, 2 of the 40 F10 families (line #71-2 and line #148-2) exhibited Wt levels of 208 susceptibility, indicating that they had lost Hpa resistance from the F9 to the F10 generation. 209 Furthermore, 4 of the 8 epiRILs tested (#71, #148, #545, and #508) displayed statistically 210 significant variation in Hpa resistance between the 5 F10 families within the epiRIL (Figure 1-211 figure supplement 5; Pearson's Chi-squared test, p < 0.05; † symbols), suggesting instability of 212 the *Hpa* resistance. 213

215 *Hpa*-resistant epiRILs show genome-wide priming of defence-related genes.

To study the transcriptomic basis of the transgenerational resistance, Wt plants (line #602) and 216 4 Hpa-resistant epiRILs (#148, #193, #454 and #508), each carrying different combinations of 217 the 4 epiQTLs, were analysed by RNA sequencing at 48 and 72 hpi (Figure 2a, bottom panel). 218 Principal component analysis (PCA) of biologically replicated samples (n = 3) revealed clear 219 220 separation between all treatment/time-point/epi-genotype combinations (Figure 2b). The first PCA axis explained 31% of the variation in transcript abundance, separating samples from 221 mock- and Hpa-treated plants, whereas the second PCA axis explained 20% of the variation, 222 223 mostly separating samples from the different lines (Figure 2b). This PCA pattern indicates that the response to *Hpa* infection had a bigger effect on global gene expression than epi-genotype. 224 Moreover, samples from Hpa-inoculated epiRILs showed relatively little difference between 225 both time-points (Figure 2b), whereas samples from Hpa-inoculated Wt plants at 48 hpi 226 clustered between samples from mock-inoculated Wt plants and samples from Hpa-inoculated 227 Wt plants at 72 hpi. This pattern suggests a difference in the speed and/or intensity of the 228 transcriptional response to Hpa. To explore this possibility further, we performed three-229 factorial likelihood ratio tests (q < 0.05) to select differentially expressed genes between all 230 epigenotype/treatment/time-point combinations. This analysis identified 20,569 genes, 231 representing 61% of all annotated RNA-producing genes in the Arabidopsis genome, including 232 transposable elements, non-coding RNA genes and pseudogenes (Supplementary dataset S3). 233 Of these, 9,364 genes were induced by Hpa at 48 and/or 72 hpi in one or more lines 234 (Supplementary dataset S4). Subsequent hierarchical clustering of this gene selection revealed 235 236 a large cluster of *Hpa*-inducible transcripts displaying augmented induction in the epiRILs at the relatively early time-point of 48h after *Hpa* inoculation (Figure 2-figure supplement 1). 237

To characterize further the pathogen-inducible transcriptome of the resistant epiRILs, 238 we selected *Hpa*-inducible genes showing elevated levels of expression in the epiRILs during 239 Hpa infection. Within this gene selection, we distinguished two expression profiles. The first 240 241 group of genes had been selected for constitutively enhanced expression in the resistant epiRILs, using the following criteria (Wald tests, q < 0.05): i) Hpa-inducible in the Wt, ii) not 242 inducible by Hpa in the epiRIL and iii) displaying enhanced accumulation in mock-treated 243 epiRIL that is equal or higher than accumulation in the Hpa-inoculated Wt ('Group 1'; Figure 244 2-figure supplement 2a). The second group of genes had been selected for enhanced Hpa-245 induced expression in the epiRILs, using the following criteria (Wald tests, q < 0.05): i) Hpa-246 247 inducible in the Wt (#602), *ii*) *Hpa*-inducible in the epiRIL(s) and *iii*) displaying statistically

increased accumulation in Hpa-inoculated epiRILs compared to Hpa-inoculated Wt plants 248 ('Group 2'; Figure 2-figure supplement 2a). For each epiRIL, we identified more genes in 249 Group 2 than in Group 1 (Figure 2c; Figure 2-figure supplements 2b, 3 and 4; Supplementary 250 datasets S5 and S6). This difference was most pronounced at 48 hpi, which represents a critical 251 time-point for host defence against $Hpa^{33,34}$. Analysis of a statistical interaction between epi-252 genotype x *Hpa* treatment revealed that > 92% of all genes in Group 2 are significant for this 253 254 interaction term (Supplementary dataset S7), indicating a constitutively primed expression pattern. Visualisation of the expression profiles in heatmaps confirmed this notion, showing 255 256 that the induction of Group 2 genes by *Hpa* is strongly augmented in the resistant epiRILs compared to the Wt line (Figure 2c; Figure 2-figure supplement 4), which is consistent with 257 the definition of plant defence priming⁷. 258

259 To examine the functional contributions of the *Hpa*-inducible genes in Groups 1 and 2, we employed gene ontology (GO) term enrichment analysis. After exclusion of redundant GO 260 terms³⁶, we identified 469 GO terms, for which one or more of the sets showed statistically 261 significant enrichment. Group 2 genes at 48 hpi displayed dramatically enhanced GO term 262 enrichment compared to all other sets, which was obvious for all epiRILs (Figure 2d). This 263 enrichment was particularly pronounced for 111 GO terms relating to SA-dependent and SA-264 independent defence mechanisms (Supplementary dataset S8), which supports our phenotypic 265 characterisation of SA-dependent and SA-independent defence markers (Figure 1-figure 266 supplement 4). Collectively, these results suggest that the quantitative resistance of the epiRILs 267 is based on priming of *Hpa*-inducible defence genes. 268

269 Interestingly, compared to the other gene selections, a relatively large proportion of 270 defence-related genes in Group 2 at 48 hpi was shared between all 4 epiRILs (Figure 2-figure supplement 2b), pointing to relatively high similarity in the augmented immune response of 271 272 the epiRILs. Furthermore, only 5% of the genes in the Group 1 and 6.5% of the genes in Group 2 are physically located within the borders of the epiQTL intervals (LOD drop-off = 2). The 273 274 frequency of Group 1 and 2 genes relative to all other genes was significantly lower for the epiQTL regions compared to the entire Arabidopsis genome (14.6%; Pearson's Chi-squared 275 test, p < 0.05). Thus, the majority of *Hpa*-inducible Group 1 and 2 genes showing enhanced 276 expression in the more resistant epiRILs are (trans-)regulated by DNA methylation at the 4 277 278 epiQTLs.

280 The resistance epiQTLs do not contain defence genes that are *cis*-regulated by DNA

281 methylation, suggesting involvement of *trans*-regulatory mechanisms.

Although 92% of all genes in Group 2 were located outside the physical borders of the 4 282 epiQTL intervals (LOD-drop-off = 2), we hypothesized that a small set of defence regulatory 283 genes inside the epiQTL regions are directly (cis-)regulated by DNA methylation to mediate 284 augmented levels of defence in response to Hpa infection. Since the Group 2 genes were 285 strongly enriched with defence-related GO terms (Figure 2d), we examined whether their 286 augmented expression during Hpa infection is associated with the hypomethylated ddm1-2 287 288 haplotype. To this end, we calculated for each gene in Group 2 the ratio of normalized transcript abundance between Hpa-inoculated epiRIL and the Hpa-inoculated Wt line, which is 289 proportional to their level of augmented expression during Hpa infection. Hierarchical 290 clustering of these ratios enabled us to select for genes that exclusively show augmented 291 expression when associated with the hypomethylated *ddm1-2* haplotype of the corresponding 292 epiQTL (Figure 3a; Figure 3-figure supplement 1a). The expression ratios of 279 epiQTL-293 localised genes did not correlate with the *ddm1-2* haplotype (Figure 3a, cluster II; Figure 3-294 figure supplement 1a; Supplementary dataset S9), indicating that DNA methylation does not 295 cis-regulate their augmented Hpa-inducible expression. By contrast, 73 epiQTL-localised 296 297 genes only showed augmented expression when associated with the hypomethylated ddm1-2haplotype (Figure 3a, cluster I; Figure 3-figure supplement 1a; Supplementary dataset S10). 298 299 To confirm the hypomethylated status of these genes, we performed comprehensive bisulfite sequencing analysis of DNA methylation for the 4 epiRILs and the Wt line. DMR analysis of 300 301 the gene body (GB), 2kb promoter region (P) and 1kb downstream (D) regions confirmed that the levels of augmented gene expression of the 279 genes in cluster II do not correlate positively 302 with the extent of DNA hypomethylation (Figure 3b, Figure 3-figure supplement 1b). This 303 notion was confirmed by linear regression analysis between the augmented expression ratio 304 305 (48 hpi) and the average level of DNA hypomethylation (Figure 3-figure supplement 2), indicating that the 279 genes in cluster II are regulated indirectly (in trans) by DNA 306 methylation. By contrast, the 73 epiQTL-based genes in cluster I showed a positive correlation 307 between augmented expression ratio (48 hpi) and DNA hypomethylation, which was 308 statistically significant for each epiQTL (p < 0.05; Figure 3-figure supplement 2). These results 309 indicate that the 73 genes in cluster I are regulated locally (in cis) by DNA methylation. 310

Nearly all *cis*-regulated genes in cluster I showed a TE-like pattern of DNA methylation in the
Wt (teM; methylation at CG, CHG and CHH contexts), whereas most cluster II genes showed

either no methylation or a pattern of gene-body methylation in the Wt (gbM; methylation at 313 CG only; Figure 3b and Figure 3-figure supplement 1b). Furthermore, dividing 314 hypomethylation at gene bodies of Group 2 genes by type of DNA methylation (i.e. either teM 315 or gbM) and plotting these values against augmented expression ratio revealed a statistically 316 significant correlation between expression ratio and reduced teM ($p=1,06e^{-8}$; Figure 3-figure 317 supplement 3), whereas no such correlation was found for reduced gbM (p=0.66; Figure 3-318 figure supplement 3). These results support the growing notion that reduced teM increases gene 319 expression, whereas changes in gbM have no direct influence on gene expression³⁷. 320

The majority of in *cis*-regulated genes in cluster I genes were annotated as TEs, such as DNA 321 transposons of the CACTA family, retrotransposons of the GYPSY or COPIA families, or TE-322 related genes, encoding transposases or enzymes necessary for TE function (Supplementary 323 324 dataset S10). Only six genes were annotated as protein-coding genes, of which two shared homology to known protein-encoding genes (At2G07240, cysteine-type peptidase; 325 326 At2G07750, RNA helicase). However, none of these two genes has previously been associated with plant defence. Furthermore, analysis of the genomic context of the six protein-coding 327 genes revealed the presence of overlapping and/or nearby TEs (Figure 3-figure supplement 4), 328 suggesting that their correlation between augmented expression and DNA hypomethylation is 329 determined by association with TEs. Since TE-encoded proteins have no antimicrobial activity 330 or direct defence regulatory function, our results suggest that global defence gene priming by 331 hypomethylated epiQTLs is not based on *cis*-regulation of defence regulatory genes, but rather 332 on alternative *trans*-acting mechanisms by DNA methylation of the TE-rich epiQTL. 333

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337 Discussion

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By screening the Col-0 x ddm1-2 epiRIL population for leaf colonisation by the downy mildew pathogen *Hpa*, we have identified 4 epiQTLs that provide quantitative disease resistance (Figure 1b). The combined contribution of all 4 DMR peak markers was estimated at 60% of the total variation (Figure 1d), which is higher than previously reported variation in developmental plant traits for this population²⁴⁻²⁶. It was previously shown that half of all stably

inherited DMRs in the Col-0 x ddm1-2 epiRILs also occur in natural Arabidopsis 344 accessions^{24,38}. Considering that the epiRIL population includes heritable variation in a range 345 of ecologically important plant traits, including flowering, root growth, nutrient plasticity and 346 (a)biotic stress resistance²⁴⁻²⁶, it is tempting to speculate that variation in DDM1-dependent 347 DNA methylation contributes to natural variation and environmental adaptation of 348 Arabidopsis. Indeed, the phenotypic diversity in the Col-0 x *ddm1-2* epiRIL population closely 349 resembles that of natural Arabidopsis accessions^{39,40}. Furthermore, independent studies have 350 shown that high levels of enduring (a)biotic stress can trigger transgenerational acquired 351 resistance (TAR) in Arabidopsis^{10,41,42}. Interestingly, repeated inoculation of 2- to 5-weeks old 352 Arabidopsis seedlings with the hemi-biotrophic leaf pathogen Pseudomonas syringae pv. 353 tomato causes TAR, which is associated with reduced transcription of DDM1 gene in local 354 leaves that is maintained in the apical meristem of paternal plants (Furci and Ton, unpublished 355 results). To what extent this prolonged repression in DDM1 gene transcription causes 356 heritable reduction in DNA methylation at the epiQTLs requires further study. 357

Aller et al. (2018) have recently used the same Col-0 x ddm1-2 epiRIL population to 358 map the contribution of heritable variation in DNA methylation to the production of defence-359 related glucosinolate metabolites⁴³. Interestingly, the resistance epiQTL on chromosome I from 360 our study partially overlaps with an epiQTL that influences basal production of the aliphatic 361 glucosinolate 3-methylthiopropyl (3MTP)⁴³. Glucosinolates contribute to defence against both 362 herbivores and microbes⁴⁴. Moreover, myrosinase-dependent breakdown products of indole-363 derived 4-methoxy-indol-3-ylmethylglucosinolate have been linked to the regulation of 364 callose-mediated cell wall defence in Arabidopsis^{45,46}. However, the 3MTP-controlling 365 epiQTL identified by Aller et al. (2018) was relatively weak compared to the epiQTL 366 controlling Hpa resistance (Figure 1b), indicating that its contribution to Hpa resistance would 367 at most be marginal. Furthermore, our transcriptome analysis revealed that the largest variation 368 in gene expression between epiRILs and the Wt line comes from the transcriptional response 369 370 to *Hpa*, rather than differences in basal gene expression (Figure 2b-c). Moreover, the genes in Group 2, which displayed enhanced Hpa-induced expression in the resistant epiRILs at the 371 critical early time-point of 48 hpi, were strongly enriched with defence-related GO terms 372 (Figure 2d). The majority of these Group 2 genes showed a statistically significant interaction 373 between epi-genotype and Hpa treatment (Supplementary dataset S7), indicating that these 374 epiRILs were primed to activate defence-related genes. This notion was supported by the actual 375 376 expression profiles of Group 2 genes (Figure 2c; Figure 2-figure supplement 4), as well as the

defence phenotypes of the 8 most resistant epiRILs in the population (Figure 2a; Figure 1figure supplement 4). Furthermore, our epiRIL screen for growth phenotypes demonstrated that the resistance-controlling epiQTLs do not have a major impacts on plant growth (Figure 1b), which is consistent with previous findings that defence priming is a low-cost defence strategy⁴⁷. While we cannot exclude other mechanisms, these independent lines of evidence collectively indicate that genome-wide priming of defence genes is the most plausible mechanism by which the epiQTLs mediate quantitative disease resistance in the population.

Over recent years, various studies have established a link between DNA 384 hypomethylation and plant immune priming^{4,6,14}. However, causal evidence that heritable 385 regions of reduced DNA methylation mediate transgenerational disease resistance is lacking. 386 Our study has shown that heritable regions of hypomethylated DNA are sufficient to mediate 387 388 resistance in a genetic Wt background. Furthermore, our study is the first to link phenotypic and epigenetic variation of selected epiRILs to profiles of global gene expression, revealing 389 390 that epigenetically controlled resistance is associated with genome-wide priming of defencerelated genes (Figure 2b-d; Figure 2-figure supplement 1; Figure 2-figure supplement 4). The 391 majority of these pathogenesis-related genes showed augmented induction at 48 hpi (Figure 392 2c), which represents a critical early time-point in the interaction between Arabidopsis and 393 Hpa, during which hyphae from germinating spores start to penetrate the epidermal cell layer 394 and invade the mesophyll^{33,34}. Notably, this set of primed genes was substantially more 395 enriched in SA-dependent and SA-independent defence GO terms than the set of Hpa-inducible 396 genes that were constitutively up-regulated in *Hpa*-resistant epiRILs (Figure 2d), corroborating 397 the analysis of phenotypical defence markers (Figure 2a; Figure 1-figure supplement 4). 398

399 DNA methylation of TEs has been reported to *cis*-regulate expression of nearby genes in Arabidopsis⁴⁸⁻⁵². By contrast, our study did not find evidence that DNA methylation in the 400 401 epiQTLs cis-regulates the responsiveness of nearby of defence genes. Firstly, the majority of primed defence genes in the Hpa-resistant epiRILs were located outside the epiQTL intervals 402 403 (92%). Secondly, of all primed genes within the epiQTLs, only 73 showed augmented 404 induction that coincided with DNA hypomethylation (Figure 3a; Figure 3-figure supplement 405 1; Figure 3-figure supplement 2; Supplementary dataset S10). Of these, 67 encoded TEs or TErelated genes, while the six protein-encoding genes were closely associated with one or more 406 407 TEs and did not have functions related plant defence (Figure 3a; Figure 3-figure supplement 1; Supplementary dataset S10). Since TEs do not encode defence signalling proteins, we propose 408 that DNA hypomethylation at the TE-rich epiQTLs mediates augmented induction of defence 409

genes across the genome via trans-acting mechanisms. A recent transcriptome study of Hpa-410 infected Arabidopsis identified 166 defence-related genes that were primed in the 411 hypomethylated *nrpe1-11* mutant and/or repressed in hyper-methylated *ros1-4* mutant¹⁴. The 412 majority of these defence genes were not targeted by NRPE1- and/or ROS1-dependent DNA 413 (de)methylation, indicating that their responsiveness is *trans*-regulated by DNA methylation. 414 Although NRPE1 and ROS1 target partially different genomic loci than DDM1²⁰, this study 415 supports our hypothesis that DNA methylation controls global defence gene responsiveness 416 via trans-acting mechanisms. 417

There are various mechanisms by which DNA methylation could trans-regulate 418 defence gene expression. It is possible that transcribed TEs in the hypomethylated epiQTLs 419 generate 21-22nt or 24nt small RNAs (sRNAs) that influence distant heterochromatin 420 formation through via RDR6- and DCL3-dependent RdDM pathways⁵³. Support for *trans*-421 regulation by sRNAs came from a recent study, which reported that induction and subsequent 422 423 re-silencing of pericentromeric TEs in Arabidopsis upon Pseudomonas syringae infection is accompanied with accumulation of RdDM-related sRNAs that are complementary to TEs and 424 distal defence genes. Interestingly, while the accumulation of these sRNAs coincided with re-425 silencing of the complementary TEs, the complementary defence genes remained expressed in 426 the infected tissues⁵⁴. These findings are supported by another recent study, which 427 demonstrated that AGO1-associated small RNAs can trans-activate distant defence gene 428 expression through interaction with the SWI/SNF chromatin remodelling complex⁵⁵. Apart 429 from sRNAs, it is also possible that long intergenic noncoding RNAs (lincRNAs) from the 430 hypomethylated epiQTLs regulate pathogen-induced expression of distant defence genes. A 431 recent study revealed that pericentromeric TEs of Arabidopsis can produce DDM1-dependent 432 lincRNAs that are increased by abiotic stress exposure⁵⁶. Since lincRNAs can promote 433 euchromatin and heterochromatin formation at distant genomic loci^{57,58}, hypomethylated TEs 434 within the epiQTLs could generate priming-inducing lincRNAs. While knowledge about 435 lincRNAs in plants remains limited, like sRNAs, their activity depends on sequence 436 complementary with target loci⁵⁹. Unlike non-coding RNAs, long-range chromatin interactions 437 can *trans*-regulate gene expression independently of sequence complementarity $^{60-63}$. Previous 438 high-throughput chromosome conformation capture (Hi-C) analysis revealed that the *ddm1-2* 439 mutation has a profound impact on long-range chromatin interactions within and beyond the 440 pericentromeric regions⁶⁴. Projection of these DDM1-dependent interactions onto the 441 Arabidopsis genome shows extensive coverage of the resistance epiQTLs identified in this 442

study (Figure 3-figure supplement 5). Whether these long-range interactions contribute to *trans*-regulation of defence gene priming would require further study, including a fully
replicated Hi-C analysis of the resistant epiRILs characterised in this study.

In conclusion, our study has shown that heritable DNA hypomethylation at selected 446 pericentromeric regions controls quantitative disease resistance in Arabidopsis, which is 447 associated with genome-wide priming of defence-related genes. This transgenerational 448 resistance is not associated with reductions in plant growth (Figure 1b), nor does it negatively 449 450 affect resistance to other types of (a)biotic stresses tested in this study (Figure 1-figure supplement 3). However, whether this form of epigenetically controlled resistance can be 451 452 exploited in crops depends on a variety of factors, including the stability of the disease resistance and potential non-target effects. For instance, our experiments with Arabidopsis 453 454 revealed that the resistance has limited stability and can erode over one more generation in some epiRILs (Figure 1-figure supplement 5). Furthermore, the genomes of most crop species 455 456 contain substantially higher numbers of TEs, rendering predictions about the applicability and potentially undesirable side effects on growth and seed production uncertain. Future research 457 will have to point out whether introgression of hypomethylated pericentromeric loci into the 458 background of elite crop varieties allows for selection of meta-stable quantitative disease 459 resistance without side-effects on agronomically important traits. 460

461

462

463 Methods

464

465 **Plant material and growth conditions.**

Epigenetic recombinant inbred lines (epiRILs) seeds of Arabidopsis (Arabidopsis thaliana, 466 accession Col-0) were purchased from Versailles Arabidopsis Stock Centre, INRA, France 467 (http://publiclines.versailles.inra.fr/epirils/index). The epiRIL screen included siblings of the 468 F4 ddm1-2 parental plant of the epiRIL population (IBENS, France). Arabidopsis seeds were 469 stratified in water at 4°C in the dark for three-five days. For pathogen bioassays, seeds were 470 sown in a sand:compost mixture (1:3) and grown at short-day conditions for three weeks (8.5 471 h light/15.5 h dark, 21°C, 80% relative humidity, ~125 µmol s⁻¹ m⁻¹ light intensity). To test 472 transgenerational inheritance and stability of Hpa resistance in the 8 most resistant epiRILs 473 474 (Figure 1-figure supplement 5), 5 individual F9 plants were cultivated for 4 weeks at short-day

- 475 conditions and then moved to long-day conditions to initiate flowering (16 h light/8 h dark,
- 476 21°C, 80% relative humidity, ~125 μ mol s⁻¹ m⁻¹ light intensity). Seeds of the 40 F10 families
- 477 were collected for analysis of *Hpa* resistance (see below).
- 478

479 Screen for variation in disease resistance and seedling growth.

Three week-old seedlings were spray-inoculated with a suspension of asexual conidia from 480 Hyaloperonospora arabidopsidis strain WACO9 (Hpa) at a density of 10⁵ spores/ml. Hpa 481 colonization was quantified at six days post inoculation (dpi) by microscopic scoring of leaves, 482 as described previously¹⁴. Briefly, trypan blue-stained leaves were analysed with a 483 484 stereomicroscope (LAB-30, Optika Microscopes) and assigned to 4 Hpa colonisation classes: class I, no hyphal colonization; class II, < 50% leaf area colonized by pathogen hyphae without 485 486 formation of conidiophores; class III, $\leq 75\%$ leaf area colonized by hyphae, presence of conidiophores; class IV, > 75% leaf area colonized by the pathogen, abundant conidiophores 487 and sexual oospores (Figure 1-figure supplement 1). At least 100 leaves per (epi)genotype were 488 analysed, not including the cotyledons. Statistically significant differences in frequency 489 distribution of Hpa colonisation classes between lines were determined by Pearson's Chi-490 squared tests, using R (v.3.5.1). Growth analysis of the epiRIL population was based on digital 491 photos (Canon 500D, 15MP) of three week-old plants, which were taken on the day of Hpa 492 inoculation. Digital image analysis of total green leaf area (GLA) was performed using Adobe 493 Photoshop 6.0. Green pixels corresponding to GLA were selected and converted into mm² after 494 495 colour range adjustment, using the magic wand tool.

496

497 Mapping of epigenetic quantitative trait loci (epiQTLs).

498 Mapping of epiQTLs was performed using the '*scanone*' function of the R/qtl package for R⁶⁵ 499 (Haley-Knott regression, step size: 2cM), combining experimental phenotypical data with the 500 recombination map of differentially methylated regions (DMR) generated previously²³. For 501 analysis of *Hpa* resistance, the categorical scoring of *Hpa* resistance was first converted into a 502 numeric resistance index (RI), using the following formula:

503
$$RI = (f_{class I} * 4) + (f_{class II} * 3) + (f_{class III} * 2) + (f_{class IV} * 1),$$

where f = relative frequency of *Hpa* colonization class of each line, multiplied by an arbitrary weight value ranging from 4 for the most resistant category (class I) to 1 for most susceptible 506 category (class IV). Mapping of epiQTLs controlling plant growth was based on average GLA 507 values of each line before *Hpa* infection. A logarithm of odds (LOD) threshold of significance 508 for each trait was determined on the basis of 1,000 permutations for each dataset ($\alpha = 0.05$). 509 The proportion of phenotypic variance $R^2(G)$ explained by the DMR markers with the highest 510 LOD score (peak markers) of all 4 epiQTLs was calculated with the following formula²⁵:

511
$$R^{2}(G) = 1 - \frac{n-1}{n-(k+1)} \frac{\sum_{i=1}^{n} (y_{i} - [\widehat{\beta}_{0} + \sum_{j=1}^{k} \beta_{j} g_{ij}])^{2}}{\sum_{i=1}^{n} (y_{i} - \overline{y})^{2}}$$

where n = number of lines analysed, k = number of DMR markers tested; $\beta_0 =$ intercept of the multiple regression model; $\beta_j =$ QTL effect for each QTL j (slopes for each marker in the multiple regression model); $g_{ij} =$ (epi) genotype of the j^{th} marker for each individual i (coded as '1' for ddm1-2 epialleles and '-1' for WT epialleles); $y_i =$ phenotypic value of individual i; $\overline{y} =$ mean of phenotypic values. The contribution of each individual QTLj ($R^2(g)$)was calculated, using the following formula:

,

518
$$R^{2}(g) = 1 - \frac{n-1}{n-(k+1)} \frac{\hat{\beta}_{j}^{2} \sum_{i}^{n} (g_{ij} - \bar{g}_{j})^{2}}{\sum_{i}^{n} (y_{i} - \bar{y})^{2}}$$

as described by ²⁵, where *n*= number of lines analysed, *k*= number of markers tested; β_j = QTL effect for each QTL*j* (slopes for each peak marker in the multiple regression model); g_{ij} = (epi)genotype of the *j*th marker for each individual *i* (coded as '1' for *ddm1-2* epialleles and '-1' for WT epialleles); \bar{g}_j = average of the (epi)genotypes values for the *j*th marker. Covariance was calculated by subtracting the sum of the individual contributions of each QTL*j* on phenotypical variance (i.e. $R^2(g_{QTL1}) + R^2(g_{QTL2}) + R^2(g_{QTL4}) + R^2(g_{QTL5})$) from the phenotypical variance explained by the full model (i.e. $R^2(G)$).

526

527 Analysis of shared transposition events.

TE-tracker software was used to interrogate available Illumina whole-genome sequencing data
from 122 epiRILs for the presence of >2 shared transposition evens (STEs) within the epiQTLs
intervals²⁸. STEs were analysed for statistically significant linkage with resistance phenotypes
(RIs), using the same linear regression model as described above for DMR linkage analysis.

532

533 *Plectosphaerella cucumerina* pathoassays.

Plectosphaerella cucumerina (Pc, strain BMM⁶⁶) was grown from frozen agar plugs (-80° C) 534 on potato dextrose agar (PDA; Difco, UK). Inoculated plates were maintained at room 535 temperature in the dark for at least two weeks. Spores were gently scraped from water-536 inundated plates, after which spore densities were adjusted to 10^6 spores/ml using a 537 hemocytometer (Improved Neubauer, Hawksley, UK). Four fully expanded leaves of similar 538 age from five weeks-old plants were inoculated by applying 5µl droplets, minimizing 539 variability due to age-related resistance. After inoculation, plants were kept at 100% RH until 540 scoring of lesion diameters. Average lesion diameters at nine dpi were based on 4 leaves per 541 542 plant from 12 plants per (epi)genotype (n=40-48), using a precision caliper (Traceable, Fischer Scientific). Statistically significant differences in necrotic lesions diameter (asterisks) were 543 quantified by two-tailed Student's t-test (p < 0.05) in pairwise comparisons with Wt line (#602), 544 using R (v3.5.1). 545

546

547 Salt stress tolerance assays.

Seeds were sterilised by exposure for 4 hours (h) to chlorine vapours from a 200ml bleach 548 solution containing 10% v/v hydrochloric acid (37% v/v HCl, Fischer Scientific, 7732-18-5). 549 Seeds were air-dried for one hour in a sterile laminar flow cabinet and plated on half strength 550 MS plates (Duchefa, M0221; +0.05% w/v MES, +1% w/v sucrose, pH 5.7), containing 551 increasing concentrations of NaCl (0mM, 50mM, 75mM and 100mM; Fischer Scientific, 7647-552 14-5). Plates were stratified for 4 days in the dark at 4°C and transferred to short-day growth 553 conditions (8.5h light/15.5h dark, 21°C, 80% RH, light intensity 100-140 µmol s⁻¹ m⁻¹). Salt 554 555 tolerance was expressed as percentage of seeds producing fully expanded cotyledons by six 556 days after stratification. Germination percentages of epi-genotypes were calculated from >50 seeds per treatment. Statistically significant differences in germination rates (asterisks) were 557 558 quantified by Fisher's exact test (p < 0.05) in pairwise comparisons with Wt line (#602) at each salt concentration, using R (v3.5.1). 559

560

561 Quantification of callose effectiveness against *Hpa* infection.

Seedlings were collected at three dpi and cleared for >24 h in 100% ethanol. One day prior to analysis, samples were incubated for 30 min in 0.07 M phosphate buffer (pH 9), followed by 15 min incubation in a 4:1 mixture (v/v) of 0.05% w/v aniline blue (Sigma-Aldrich, 415049) in 0.07M phosphate buffer (pH 9) and 0.025% w/v calcofluor white (Fluorescent brightener

28, Sigma-Aldrich, F3543) in 0.1M Tris-HCL (pH 7.5). After initial staining, samples were 566 incubated overnight in 0.5% w/v aniline blue (Sigma-Aldrich, 415049) in 0.07M phosphate 567 buffer (pH 9) and scored with an epifluorescence microscope (Olympus BX 51) fitted with 568 blue filter (XF02-2; excitation 330nm, emission 400nm). Germinated conidia (germ tubes) 569 were divided between in two classes: non-arrested and arrested by callose. In each assay, 10 570 571 leaves from different plants for each (epi)genotype were analysed, amounting to >150 conidiacallose interactions. Statistically significant differences in resistance efficiency of callose 572 (asterisks) were analysed using Pearson's Chi-squared tests (p < 0.05) in pairwise comparisons 573 574 with Wt line (#602), using R (v3.5.1).

575

576 Reverse-transcriptase quantitative polymerase chain reactions (RT-qPCR).

Three biologically replicated samples for each genotype/treatment/time-point combination 577 578 were collected at 48 and 72 hpi, each consisting of six to 12 leaves collected from different plants per pot. Samples were snap-frozen in liquid nitrogen and ground to a fine powder, using 579 a tissue lyser (QIAGEN TissueLyser). Total RNA was extracted using a guanidinium 580 thiocyanate-phenol-chloroform extraction isolation protocol. Frozen powder was vortexed for 581 30 seconds in 1ml Extraction buffer: 1M guanidine thiocyanate (Amresco, 0380), 1M 582 ammonium thiocyanate (Sigma-Aldrich, 1762-95-4), 0.1M sodium acetate (Fisher Scientific, 583 127-09-3), 38% v/v AquaPhenol (MP Biomedicals, 108-95-2) and 5% v/v glycerol (Fisher 584 Scientific, 56-81-5). Samples were incubated at room temperature (RT) for one min and then 585 centrifuged for five min at 16,500 g. The supernatant was then transferred to a new tube, mixed 586 with 200µl chloroform and vortexed for 10-15 sec. After centrifuging for five min (16,500 g), 587 588 the aqueous phase was transferred to new tubes, gently mixed by inversion with 350µl 0.8M sodium citrate (Sigma-Aldrich, 6132-04-3) and 350µl isopropanol (Fischer Chemicals, 67-63-589 590 0) and left at RT for 10 min for RNA precipitation. Samples were centrifuged for 15 min at 16,500 g (4°C), after which pellets were washed twice in 1ml 70% ethanol, centrifuged at 591 592 16,500 g for 1 min, and air-dried before dissolving in 50µl nuclease-free water. Total RNA was quantified, using a Nanodrop 8000 Spectrophotometer (Thermo Scientific). RNA extracts were 593 594 treated with DNaseI, using the RQ1 RNase-Free DNase kit (Promega, M6101). First-strand cDNA synthesis was performed from 1µg RNA, using SuperScript III Reverse Transcriptase 595 596 (Invitrogen, 18080093) according to the supplier's recommendations. The qPCR reactions were carried out with a Rotor-Gene Q real-time PCR cycler (Qiagen) and the Rotor-Gene 597 SYBR Green PCR Kit (Qiagen, 204074). Relative PR1 gene expression was calculated, using 598

Livak's ΔΔCT method⁶⁷ with correction for average PCR efficiencies for each primer pair across experiment samples. Gene expression was normalised against average expression values of At1G13440 (GAPDH), At5G25760 (UBC) and At2G28390 (SAND family protein)⁶⁸. Reactions were performed using previously described primer sequences¹⁴. Statistically significant differences in relative expression (asterisks) were quantified by two-tailed Student's t-test (p < 0.05) in pairwise comparisons with *Hpa*-treated Wt line (#602).

605

606 Transcriptome analysis.

Samples for RNA sequencing were collected at 48 and 72 hpi of three week-old plants. Every 607 608 epi-genotype/treatment/time-point combination was based on three biologically replicated samples, each consisting of 6-12 shoots from different plants. Initial RNA extraction was 609 610 performed as described for RT-qPCR reactions. Prior to library preparation, RNA concentration and integrity were measured, using 2100 Bioanalyzer (Agilent) with provided 611 reagents kits and according to manufacturer's instructions. All RNA samples had RNA 612 integrity numbers (RIN) > 7.5. Sequencing libraries were prepared from total RNA, using the 613 TruSeq Stranded Total RNA kit and Ribo-Zero Plant leaf kit (Illumina, RS-122-2401), 614 according to the manufacturer's instructions. Sequencing runs were performed on a HiSeq1500 615 platform (Illumina), generating paired-end reads of 125 bp and an average quality score (Q30) 616 > 93%. Each sample generated around 35 million paired reads. 617

Read quality was assessed by FastQC software⁶⁹. Read length and distribution were 618 optimized and adapter sequences were trimmed, using Trimmomatic software⁷⁰. Reads were 619 aligned and mapped to the Arabidopsis genome (TAIR10 annotation), using splice site-guided 620 HISAT2 alignment software (John Hopkins University, second iteration of ⁷¹). For all samples, 621 more than 95% of reads could successfully be mapped once or more onto the Arabidopsis 622 genome. Number of reads per gene were quantified with the Python package $HTseq^{72}$. 623 Differential expression analysis was performed using the DESeq2 R package, which applies a 624 625 negative binomial generalized linear model to estimate mean and dispersion of gene read counts from the average expression strength between samples⁷³. Prior to principal component 626 627 analysis (PCA) by the *plotPCA* function, gene read counts were subjected to regularized logarithmic transformation, using the *rlog* function⁷³. Likelihood ratio tests of variance within 628 a three-factorial linear model for epigenotype, treatment, time-point and interactions thereof 629 were used to identify genes showing differences in expression across one or more factors⁷³. 630

Differentially expressed genes (DEGs) were subjected to hierarchical clustering (Ward 631 method) and presented as a heat map, using the *pheatmap* R package⁷⁴. For each gene, *rlog*-632 normalized read counts of each sample were subtracted from the mean of all samples, and 633 divided by the standard deviation to facilitate heatmap visualization (z-score). To identify 634 DEGs between two treatment/time-point/epi-genotype combinations, pair-wise comparisons 635 (Wald test; q < 0.05) were performed with the DEGs selection obtained by the *lrt* test, using the 636 selection criteria illustrated in Figure 2-figure supplement 2a. All Hpa-inducible genes in the 637 Wt and/or epiRILs were selected for elevated expression in the more resistant epiRILs during 638 639 Hpa infection. Subsequently, these genes were divided between two groups based on their expression profile. Group 1 genes were selected for constitutively enhanced expression in the 640 epiRIL(s) relative to the Wt (Figure 2-figure supplements 2 and 3); Group 2 genes were 641 selected for enhanced levels of Hpa-induced expression in the epiRIL(s) relative to the Wt 642 (Figure 2-figure supplements 2 and 4). To determine the number of Group 2 genes that show a 643 statistically significant interaction between epigenotype x Hpa treatment (, all 16,009 genes 644 significant for this interaction were selected from the three-factorial linear model, using the 645 contrast function, and cross-referenced against Group 2 genes. 646

Gene ontology (GO) term enrichment analysis was performed with the Plant GSEA 647 toolkit⁷⁵. GO terms were checked for significant enrichment against the whole genome 648 background, using a hypergeometric test and Benjamini-Hochberg false discovery rate 649 correction (q < 0.05). Lists of enriched GO terms in each treatment were analysed by the GO 650 Trimming 2.0 algorithm³⁶ to remove redundancy of terms, applying a soft trimming threshold 651 of 0.40. The output list from GO Trimming 2.0 was run through GOSlim Viewer (AgBase) to 652 reduce GO terms according to GO slim ontologies (GO consortium). Enrichment was 653 quantified as the percentage of GO term-annotated genes within a certain selection relative to 654 655 the total number of Arabidopsis genes in that GO term.

656

657 Methylome analysis.

For each line, three independent biological replicates were collected, consisting of pooled
leaves from six plants of the same developmental stage. High quality genomic DNA was
extracted from leaves of five week-old plants, using the GenEluteTM Plant Genomic DNA
Miniprep Kit (Sigma-Aldrich). Bisulfite sequencing was performed by GATC Biotech (UK).
After quality trimming of read sequences, adapter sequences were removed, and reads were

filtered by Cutadapt (version 1.9; Pair end-mode; phred score = 20, min.length = 40). Reads were mapped to an index genome, using of BS-Seeker2 (version 2.0.10, mismatch = 0.05, maximum insert size =1000 bp). Bowtie2 (version 2.2.2) was used for alignment of reads, as described previously⁷⁶. Differential methylation for promoter regions (-2kb), gene bodies, and downstream regions (+1kb) relative to the Wt was called using methylkit (version 1.0.0; minimum coverage = 5x, q = 0.05). Differentially methylated states were visualised as a heat map, using the '*pheatmap*' R package (version 1.0.8)⁷⁴.

- To differentiate Wt methylation states of all epiQTL-based genes in Group 2 (see above), gene 670 bodies of all nuclear genes were categorised between un-methylated, gene body methylated 671 (gbM; CG context only) or TE-like methylated (teM; CHG and/or CHH with or without CG). 672 For each gene containing 20 or more cytosines, methylated and un-methylated cytosine base 673 674 calls in each context were extracted from the sequence read alignments. Positions with less than 4x coverage were ignored. Methylation patterns were categorised as TE-like if methylated 675 676 read calls relative to un-methylated read calls in CHG and/or CHH contexts showed a statistically significant increase over average methylation rates of all genes across the genome 677 in the respective context, using the "binom.test" function in R (FDR-adjusted p < 0.01). The 678 remaining genes were classified either as gbM if the same test revealed a statistically significant 679 increase in CG context, or as un-methylated if no statistically significant increase in DNA 680 methylation could be detected in any sequence context. 681
- 682

683 Correlation analysis between gene expression and DNA methylation.

Correlations between augmented expression ratio of Group 2 genes (see Transcriptome 684 685 analysis) and DNA hypomethylation (CG), were determined by plotting augmented gene ratios at 48 hpi against average hypomethylation compared to Wt (%) across promoter region, gene 686 687 body, and downstream region (see Methylome analysis). To determine which type of DNA hypomethylation correlates with augmented expression in the epiRILs, hypomethylation at 688 gene bodies of Group 2 genes were divided between teM and gbM and plotted against the 689 corresponding expression ratios at 48 hpi. If hypomethylation occurred at CG context only, 690 691 genes were classified as being reduced in gene body methylation (gbM); if hypomethylation occurred all three sequence contexts (CG, CHG, CHH), genes were classified as being reduced 692 in TE methylation (teM). Values of gbM hypomethylation were expressed as percentage 693 reduction in GC methylation relative to the Wt; values of teM hypomethylation were expressed 694

as percentage reduction in all sequence contexts. Linear regression analyses were performedusing R software (v.3.5.1).

697

698 Hi-C analysis.

HiC sequence libraries SRR1504819 and SRR1504824⁶⁴ were downloaded from NCBI SRA. 699 Sequences were pre-processed and aligned to the TAIR10 Arabidopsis nuclear genome 700 sequence⁷⁷, using HiCUP (0.5.9)⁷⁸ and Bowtie2⁷⁹ (2.2.6). Alignments were filtered and de-701 duplicated as part of the processing by HiCUP, before being further processed in HOMER⁸⁰ 702 (4.9.1) at 5kb resolution. Differential interactions were assessed reciprocally, using each 703 sample as background (analyzeHiC-ped). Interactions were determined to be potentially 704 dependent on genotype if the absolute z-score of the primary versus the secondary experiment 705 was more than 1. Visualisations were generated using $Circos^{81}$ (0.69-5), based on bundled links 706 (-max gap 10001). 707

708

709 Data availability.

Transcriptome sequencing and bisulfite sequencing reads are available from the European
Nucleotide Archive (ENA) under accession code PRJEB26953.

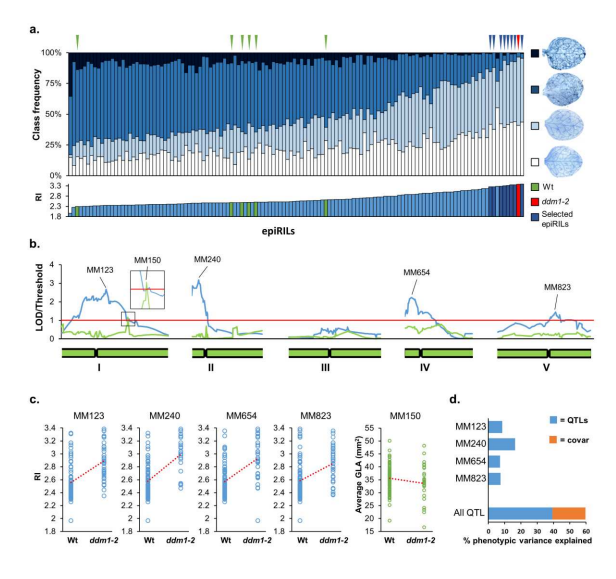
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713 Acknowledgements

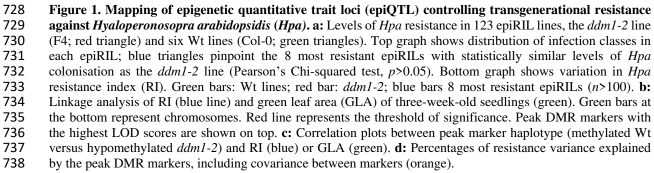
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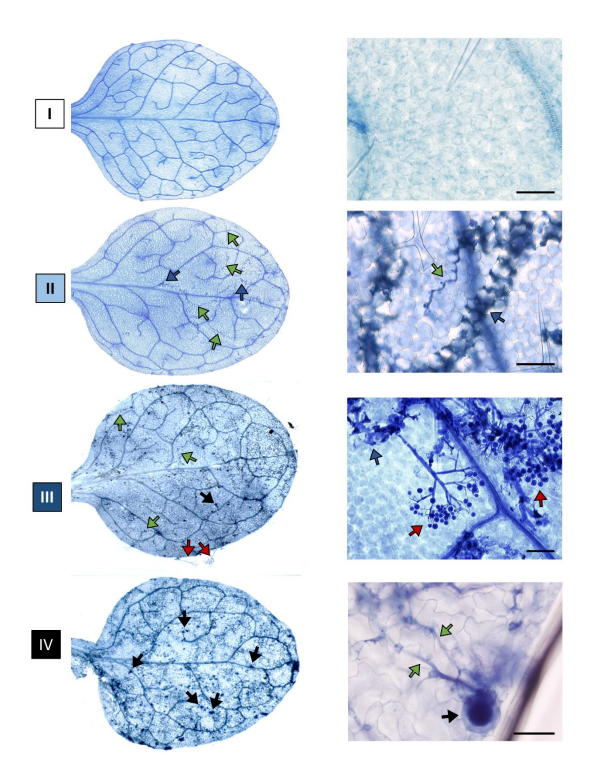
We thank David Pardo and Ana Lopez for technical assistance in the lab. We thank the La 715 Trobe University's Genomics Platform for technical support. The research was supported by 716 a consolidator grant from the European Research Council (ERC; no. 309944 "Prime-A-Plant") 717 to J.T., a Research Leadership Award from the Leverhulme Trust (no. RL-2012-042) to J.T. 718 and a BBSRC-IPA grant to J.T. (BB/P006698/1). Work in V. C. group was supported by the 719 (ANR-09-BLAN-0237 720 Agence Nationale de la Recherche EPIMOBILE). FJ acknowledges support from the Technical University of Munich - Institute for Advanced 721 Study funded by the German Excellent Initiative and the European Seventh Framework 722 723 Programme under grant agreement no. 291763. FJ is also supported by the SFB/Sonderforschungsbereich924 of the Deutsche Forschungsgemeinschaft (DFG). 724













740 Figure 1-figure supplement 1: Representative examples of infection classes used for quantification of Hpa 741 resistance. Shown are trypan blue-stained Arabidopsis leaves at six days after spray-inoculation with Hpa. White 742 (class I), absent or minimal colonisation; light blue (class II), $\leq 50\%$ leaf area colonised by the pathogen; dark 743 blue (class III), $\leq 75\%$ leaf area colonised by the pathogen, presence of conidiophores; black (class IV), > 75\% 744 leaf area colonised by the pathogen, conidiophores and abundant sexual spores. Green arrows indicate colonisation 745 by pathogen hyphae; blue arrows indicate hyphae surrounded by trailing necrosis, red arrows indicate 746 conidiophores, black arrows indicate sexual oospores. Insets on the right show higher magnifications of 747 colonisation markers. Scale bar = $50\mu m$.

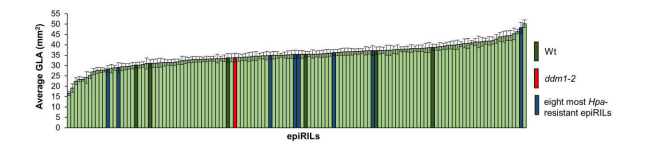
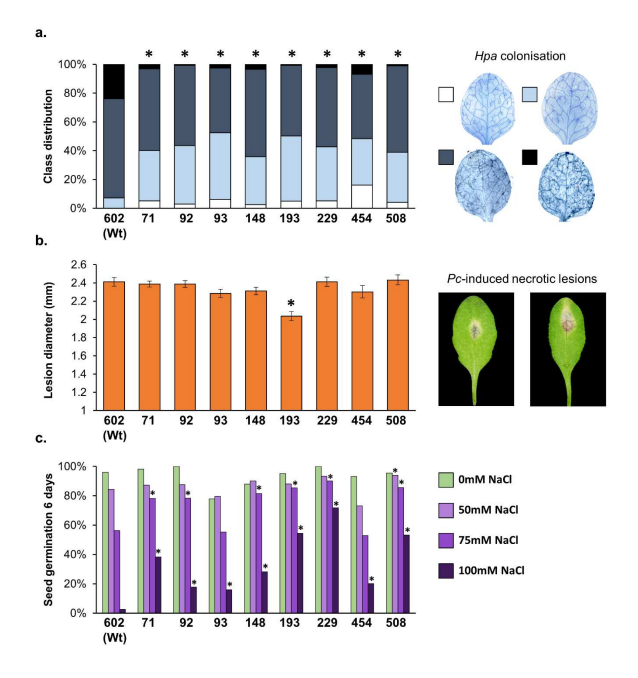
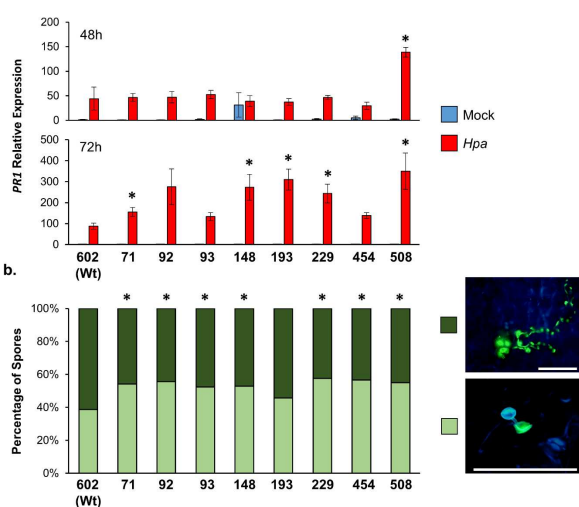


Figure 1-figure supplement 2. Average green leaf area (GLA) of the 123 epiRILs (light green), the *ddm1-2* line
 (F4; red) and six Wt lines (Col-0; dark green). Shown are average GLA values of three-week-old plants (±SEM).



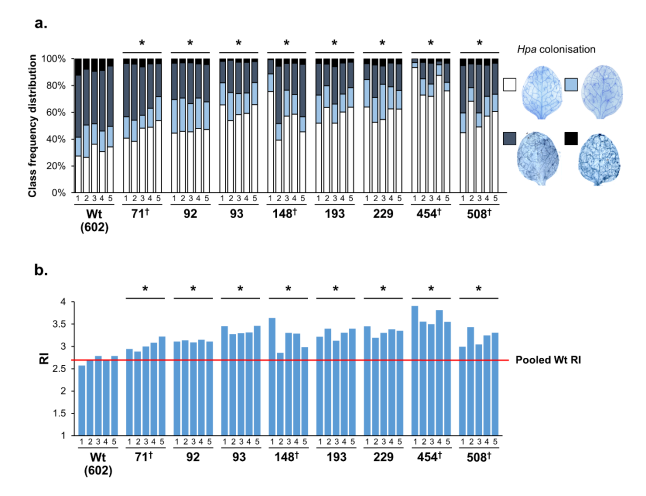
753 Figure 1-figure supplement 3. Resistance phenotypes of the 8 most Hpa-resistant epiRILs against different 754 (a)biotic stresses. a: Confirmation of resistance against biotrophic *Hpa*. Shown are levels of infection at six days 755 post inoculation (dpi) of three-week-old plants. Trypan blue-stained leaves were analysed by microscopy and 756 assigned to 4 *Hpa* infection classes (insets on the right; see Figure 1-figure supplement 1 for further details). 757 Statistically significant differences in class distribution (asterisks) were analysed using Pearson's Chi-squared 758 tests (p < 0.05) in pairwise comparisons with Wt line (#602); n > 80. **b**: Quantification of resistance against 759 necrotrophic *Pletosphaerella cucumerina* (Pc). Shown are average lesion diameters (\pm SEM) at nine days after 760 droplet inoculation with Pc spores onto similarly aged leaves of five-week-old plants. Insets show representative 761 examples of necrotic lesions by Pc. Statistically significant differences in necrotic lesions diameter (asterisks) 762 were quantified by two-tailed Student's t-test (p < 0.05) in pairwise comparisons with Wt line (#602); n=40-48. c. 763 Quantification of salt (NaCl) tolerance. Shown are percentages of seedlings developing full cotyledons after six 764 days of growth on agar with increasing NaCl concentrations. Statistically significant differences in germination 765 rates (asterisks) were quantified by Fisher's exact test (p < 0.05) in pairwise comparisons with Wt line (#602) at 766 each salt concentration; n > 50.



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Figure 1-figure supplement 4. Defence marker phenotypes of the 8 most Hpa-resistant lines. a: relative 768 769 expression of SA-dependent PR1 at 48 and 72 hpi with Hpa (red) or water (blue). Shown are mean relative 770 expression values (±SEM). Statistically significant differences in relative expression (asterisks) were quantified 771 by two-tailed Student's t-test (p < 0.05) in pairwise comparisons with Hpa-treated Wt line (#602); n=3 b: 772 Resistance efficiency of callose deposition in Hpa-inoculated plants. Shown are percentages of arrested (light) 773 and non-arrested (dark) germ tubes at 48 hpi. Insets show representative examples of aniline-blue/calcofluor-774 stained leaves by epi-fluorescence microscopy (bars = $100\mu m$; yellow indicates callose; blue indicates Hpa). 775 Statistically significant differences in resistance efficiency of callose (asterisks) were analysed using Pearson's 776 Chi-squared tests (p < 0.05) in pairwise comparisons with Wt line (#602); n > 150.

a.



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Five individual F9 plants from the 8 most resistant epiRILs and the Wt line (#602) were self-pollinated to

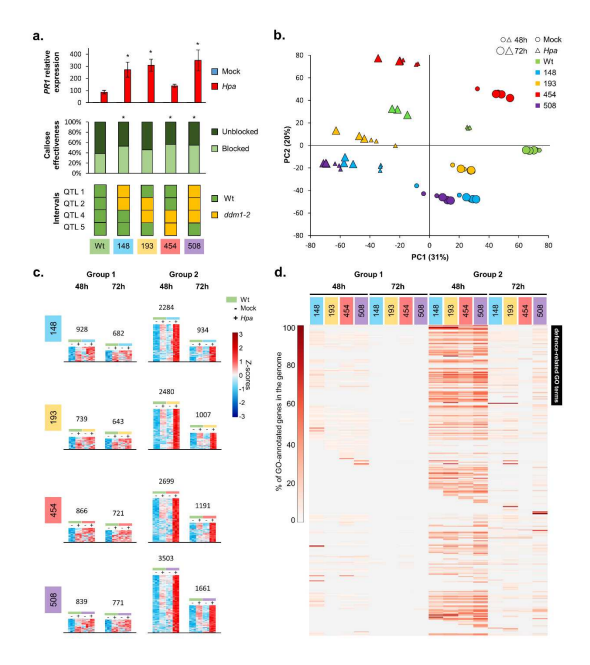
generate 40 F10 families. Plants of each F10 family were analysed for *Hpa* colonisation at 6 dpi. a: Shown are
 frequency distributions of leaves across 4 *Hpa* colonisation classes (insets on the right; see Figure 1-figure

supplement 1 for further details). **b:** Resistance index (RI) values of the F10 families. The red line indicates the

average RI value of the Wt (#602). Asterisks at the top of each graph indicate statistically significant differences
 in class distribution between pooled F10 families of the epiRIL relative to pooled F10 families of the Wt line

(Pearson's Chi-squared test; p < 0.05). Crosses (†) at the bottom of each graph indicate statistically significant

differences between F10 families within each epiRIL (Pearson's Chi-squared test; p < 0.05).



788 Figure 2. The defence-related transcriptome of *Hpa*-resistant epiRILs. a: Defence marker phenotypes and 789 epiQTL haplotypes of 4 Hpa-resistant epiRILs and the Wt (#602), which were analysed by RNA sequencing. Top 790 graph: relative expression of SA-dependent PR1 at 72 hours after inoculation (hpi) with Hpa (red) or water (blue). 791 Middle graph: resistance efficiency of callose deposition in Hpa-inoculated plants. Shown are percentages of 792 arrested (light) and non-arrested (dark) germ tubes at 48 hpi. Bottom panel: epiQTL haplotypes of selected lines. 793 Green: methylated Wt haplotype; yellow: hypomethylated ddm1-2 haplotype. Asterisks indicate statistically 794 significant differences to the Wt. (see Figure 1-figure supplement 4 for statistical information). b: Principal 795 component analysis of 27,641 genes at 48 (small symbols) and 72 (large symbols) hpi with Hpa (triangles) or 796 water (Mock; circles). Colours indicate different lines. c: Numbers and expression profiles of Hpa-inducible genes 797 that show constitutively enhanced expression (Group 1) or augmented levels of *Hpa*-induced expression (Group 798 2) in the Hpa-resistant epiRILs at 48 or 72 hpi. Heatmaps show normalised standard deviations from the mean (z-799 scores) for each gene (rows), using *rlog*-transformed read counts (see Figure 2-figure supplements 3 and 4 for 800 better detail) d: GO term enrichment of primed and constitutively up-regulated genes. Shown are 469 GO terms 801 (rows), for which one or more epiRIL(s) displayed a statistically significant enrichment in one or more categories 802 (Hypergeometric test, followed by Benjamini-Hochberg FDR correction; q < 0.05). Heatmap-projected values for 803 each GO term (rows) represent percentage of GO-annotated genes in each category relative to all GO-annotated 804 genes in the Arabidopsis genome (TAIR10). Black bar on the top right indicates 111 defence-related GO terms.

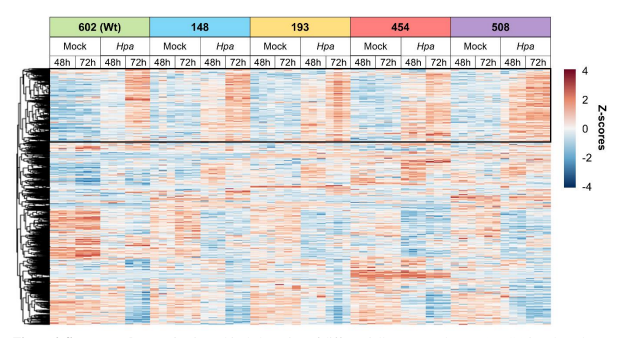


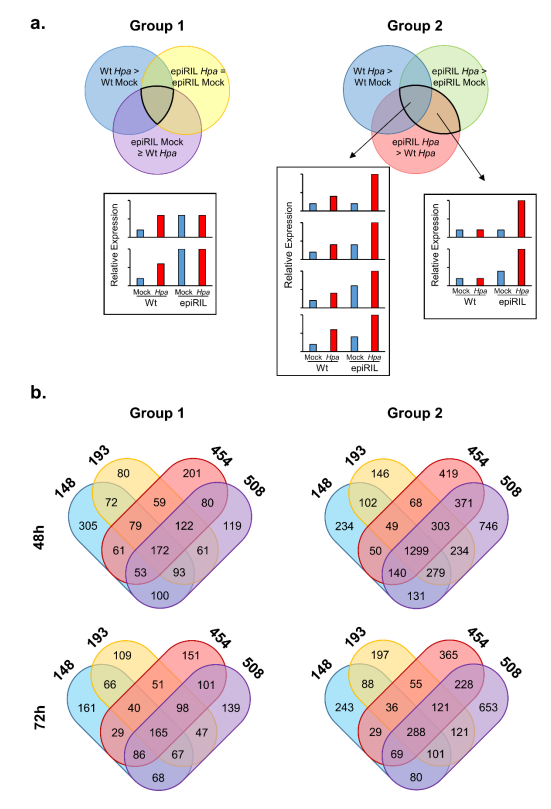
Figure 2-figure supplement 1. Hierarchical clustering of differentially expressed genes (DEGs) in selected *Hpa* resistant epiRLs and the Wt at 48 and 72 hpi (Ward method). The heatmap shows normalised standard deviations

808 from the mean (z-scores) for each DEG (rows), using *rlog*-transformed read counts. Columns represent three

809 biological replicates for each line-treatment-timepoint combination. The black square at the top of the heatmap

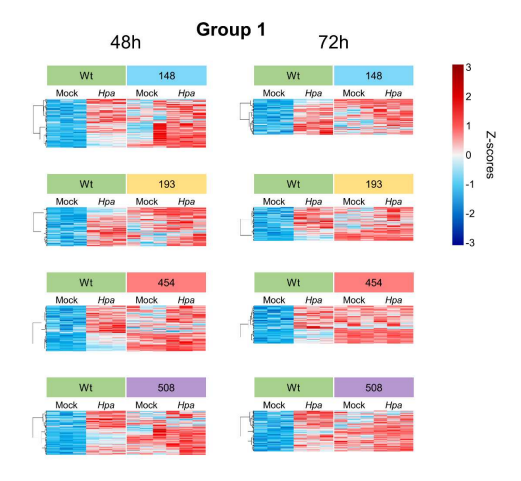
810 indicates a cluster with genes that show augmented induction in one or more resistant epiRILs at 48 h after *Hpa*

811 inoculation.



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Figure 2-figure supplement 2. Selection of *Hpa*-inducible genes that show constitutively enhanced expression (Group 1) or enhanced *Hpa*-induced expression in the resistant epiRILs (Group 2). a. Description of each circle within the Venn diagrams indicates the statistical criteria used to obtain each selection (Wald test, q < 0.05). Overlapping areas in Venn diagrams (highlighted by black lines) indicate combinations of criteria used to select differently regulated genes. Panels below show schematic examples illustrating expression profiles of selected genes. b. Number of Group 1 and Group 2 genes in *Hpa*-resistant epiRILs. Venn diagrams show numbers of genes that are unique or shared between epiRILs for each gene selection and time-point.



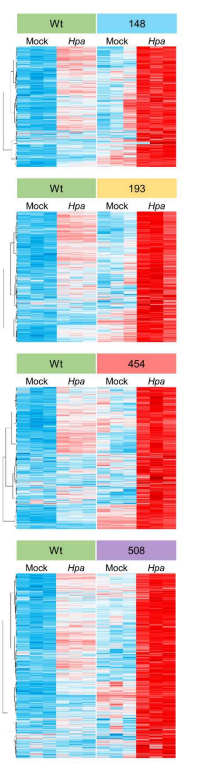
822 Figure 2-figure supplement 3. Transcript profiles of *Hpa*-inducible genes showing constitutively enhanced

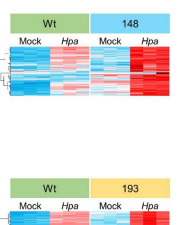
expression in the *Hpa*-resistant epiRILs (Group 1). Heatmaps show normalised standard deviations from the
 mean (z-scores) for each gene (rows) at 48 and 72 hpi, using *rlog*-transformed read counts. Columns represent

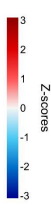
825 three biological replicates for each line-treatment combination. Expression profiles were subjected to hierarchical

826 clustering by gene (Ward method).









	Wt		454	
	Mock	Нра	Mock	Hpa
4	-			
ł				

V	Wt		508	
Mock	Нра	Mock	Hpa	

Figure 2-figure supplement 4. Transcript profiles of *Hpa*-inducible genes showing enhanced levels of *Hpa* induced expression in the *Hpa*-resistant epiRILs (Group 2). Heatmaps show normalised standard deviations
 from the mean (z-scores) for each gene (rows) at 48 and 72 hpi, using *rlog*-transformed read counts. Columns
 represent three biological replicates for each line-treatment combination. Expression profiles were subjected to
 hierarchical clustering by gene (Ward method).

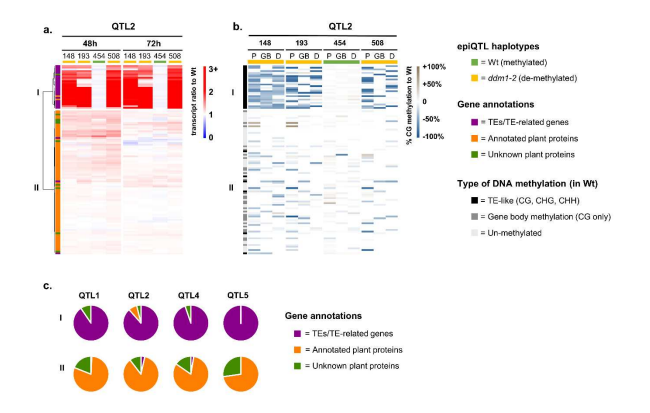


Figure 3. Relationship between augmentation of pathogen-induced expression and DNA methylation for epiQTL-localised genes. a: Expression profiles of epiQTL-based genes showing elevated levels of *Hpa*-induced expression in one or more epiRIL(s) (Group 2). Shown are genes located in the epiQTL interval of chromosome II (epiQTL2; LOD drop-off = 2; see Figure 3-figure supplement 1a for other the epiQTLs). Heatmap shows gene expression ratios between *Hpa*-inoculated epiRILs and the Wt, representing augmented expression levels during pathogen attack. Hierarchical clustering yielded two distinctly regulated gene clusters (I and II). Coloured bars on the top indicate epiQTL2 haplotypes. Green: methylated Wt haplotype. Yellow: hypomethylated *ddm1-2*

haplotype. b: Levels of CG DNA methylation of the same genes in the epiQTL2 interval (see Figure 3-figure
supplement 1b for other epiQTLs). Heatmap shows percentages of hypomethylation (blue) or hyper-methylation
(brown) relative to the Wt for 2kb promoter regions (P), gene bodies (GB) and 1kb downstream regions (D). c:

844 Distribution of gene annotations of distinctly regulated gene clusters for each epiQTL.

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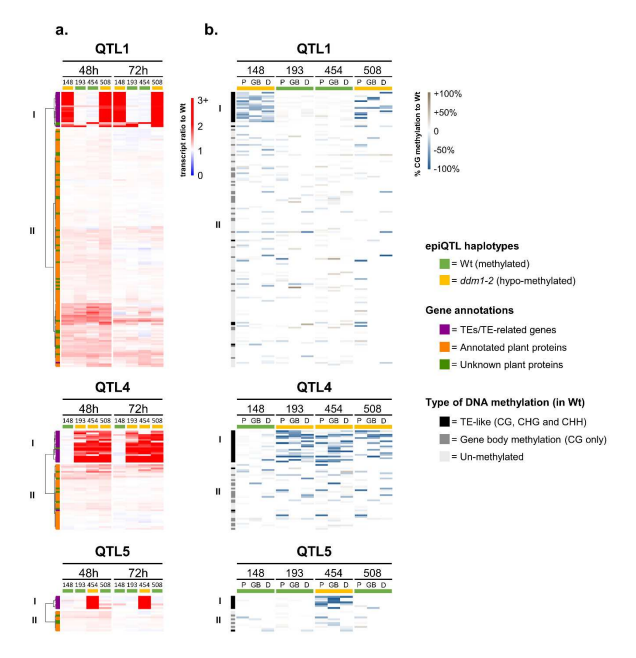
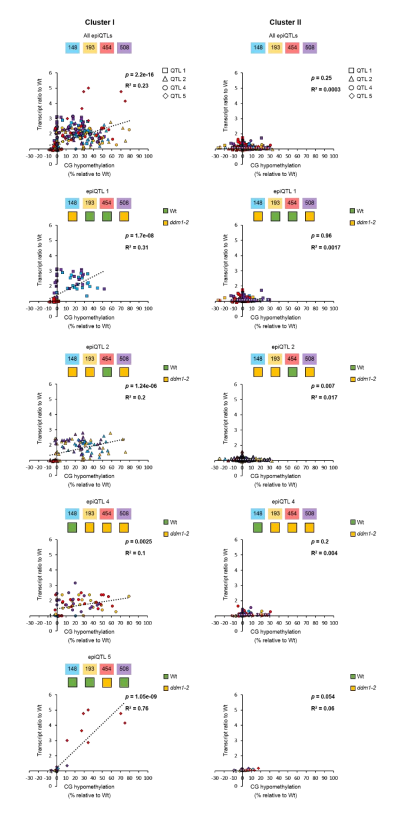
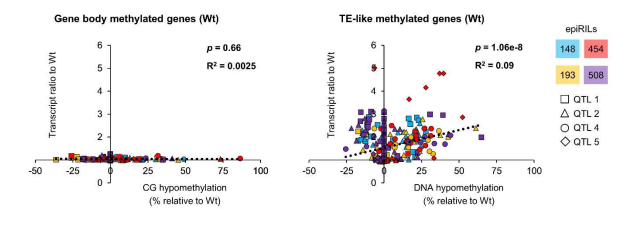


Figure 3-figure supplement 1. Relationship between augmentation of pathogen-induced expression and
DNA methylation for epiQTL-localised genes. a: Expression profiles of epiQTL-based genes with elevated
levels of *Hpa*-induced expression in one or more epiRILs (Group 2). Shown are genes located in the epiQTL
intervals (LOD drop-off = 2) of chromosomes I (epiQTL1), chromosome IV (epiQTL4) and chromosome V
(epiQTL5). b: Levels of DNA methylation of the same genes in epiQTL1, epiQTL4 and epiQTL5. For details,
see legend to Figure 3.



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858 Figure 3-figure supplement 2. Correlation analysis between augmented gene transcription and DNA 859 hypomethylation. Augmented gene transcription was defined as the ratio between the Hpa-inoculated epiRIL 860 and the Hpa-inoculated Wt at 48 hpi (Figure 3a and Figure 3-figure supplement 1a). DNA hypomethylation values were averaged across promoter regions, gene bodies and downstream regions. Scatter plots show transcript ratios 861 862 against the hypomethylation for each gene in expression clusters I and II of Group 2 (Figure 3b and Figure 3-863 figure supplement 1b), which were selected by hierarchical clustering of augmented expression profiles in the 864 epiRILs during Hpa infection. Significant positive correlations (Pearson linear regression; p < 0.05) indicate cis-865 regulation by DNA methylation.



867

868 Figure 3-figure supplement 3. Correlation analysis between augmented gene transcription and type of DNA 869 hypomethylation. Scatter plots show augmented transcript ratios against hypomethylation for all epiQTL-based genes in Group 2 (Figure 3 and Figure 3-figure supplement 1). Augmented transcription was defined as the ratio 870 871 between the Hpa-inoculated epiRIL and the Hpa-inoculated Wt at 48 hpi (Figure 3a and Figure 3-figure 872 supplement 1a). Hypomethylation values at gene bodies in the epiRILs were divided according to the type DNA 873 methylation. If hypomethylation occurred at CG context only, genes were classified as being reduced in gene 874 body methylation (gbM); if hypomethylation occurred all three sequence contexts (CG, CHG, CHH), genes were 875 classified as being reduced in TE methylation (teM). Values of gbM hypomethylation are expressed as percentage 876 reduction in GC methylation relative to the Wt; values of teM hypomethylation are expressed as percentage reduction in all sequence contexts. Statistically significant correlations (Pearson linear regression; p < 0.05) 877 878 indicate cis-regulation by DNA methylation.

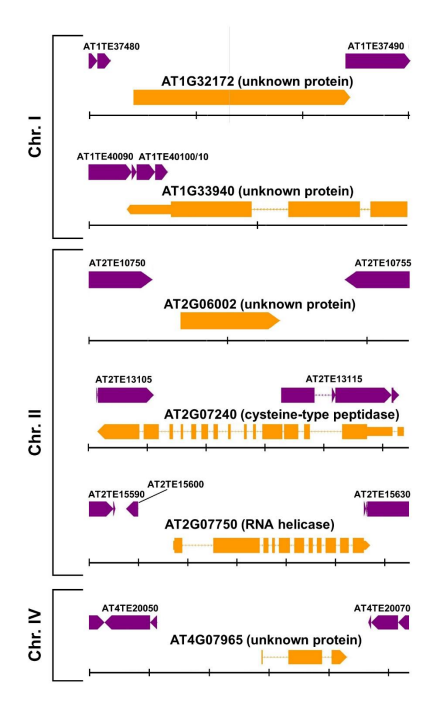


Figure 3-figure supplement 4. Genomic contexts of six plant protein-encoding genes in the epiQTL
 intervals, whose transcriptional priming coincides with reduced DNA methylation. Orange bars indicate gene
 models; superimposed purple bars indicate associated transposable elements (TEs). Large blocks represent exons;
 lines between blocks represent introns; smaller blocks at the 3' and 5' ends represent un-translated regions. Units
 of the back scale correspond to 1Kb.

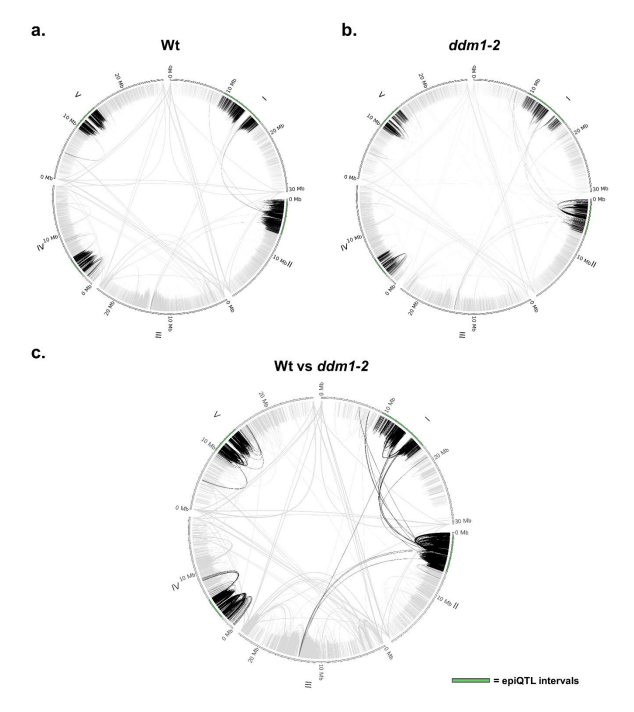


Figure 3-figure supplement 5. Genome-wide chromatin interactions in Wt and ddm1-2 **Arabidopsis.** Circular diagrams show all five Arabidopsis chromosomes. The 4 epiQTL regions are highlighted in green. Chromatin interactions are indicated by lines. Gray lines: interactions outside the epiQTLs. Black lines: interactions with the epiQTLs. Presented results are based on Hi-C data from Feng et al. $(2014)^{62}$ **a.** Genome-wide chromatin interactions in the Wt (Col-0). **b.** Genome-wide chromatin interactions in the ddm1-2 mutant. **c.** DDM1dependent chromatin interactions that are altered in the ddm1-2 mutant compared to the Wt plants.

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