Pathogen Infection and MORC Proteins Affect Chromatin Accessibility of Transposable Elements and Expression of Their Proximal Genes in Arabidopsis

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To assess the role of MORC1 in epigenetics in relation to plant immunity, genome-wide chromatin accessibility was compared between mock- or Pseudomonas syringae pv. tomato–inoculated wild type (WT) Arabidopsis, the morc1/2 double mutant, or both. Most changes in chromatin accessibility, scored by DNase I hypersensitive sites (DHSs), were located in the promoters of genes and transposable elements (TEs). Comparisons between morc1/2 and WT receiving the same treatment revealed differential DHSs (dDHSs) predominantly associated with heterochromatic TEs. By contrast, comparisons between mock- and P. syringae pv. tomato–inoculated plants from the same genotype showed a smaller but significant population was in TEs. Moreover, many defense genes, including PR-1, PR-2, and PR-5, were proximal to P. syringae pv. tomato–induced, TE-associated dDHSs. A random subset of these defense genes showed moderately delayed or reduced expression or both in P. syringae pv. tomato–infected morc1/2 as compared with WT. MORC1 was physically bound to chromatin in a P. syringae pv. tomato infection-responsive manner at sites dispersed throughout the genome. Notably, silencing of TE-associated dDHSs proximal to these infection-induced, MORC1-interacting sites led to significant suppression of P. syringae pv. tomato–induced transcription of adjacent defense genes, including PR-1. These results provide evidence that MORC1 is associated with TEs and suggest that a subset of these TEs may help regulate their proximal defense genes.

Plants have evolved a variety of defense mechanisms to protect themselves from potentially pathogenic microorganisms. Upon infection, recognition of pathogen-associated molecular patterns (PAMPs) by extracellular surface receptors leads to the activation of PAMP-triggered immunity (PTI). While PTI is sufficient to prevent the further colonization of many microbes, evolutionary selection has led to the appearance of some pathogens containing effectors that suppress PTI. Whether such pathogens can effectively infect the host plant is then determined by whether the plant expresses a resistance (R) protein that recognizes one of these effectors (also termed avirulence [Avr] factors) and induces effector-triggered immunity (ETI) (Jones and Dangl 2006). Following the activation of either ETI or PTI, a variety of defense responses are induced, such as the accumulation of salicylic acid (SA) and the increased expression of certain pathogenesis-related (PR) genes (Vlot et al. 2009). In addition, the activation of plant immunity is associated with large-scale changes in gene expression (Moore et al. 2011; Vlot et al. 2009). This transcriptional reprogramming involves the highly coordinated action of myriad transcription factors and their associated proteins that function to recruit or modulate RNA polymerase II (RNAPII) (Moore et al. 2011). Growing evidence suggests that chromatin modification and remodeling, which regulates the accessibility of DNA to the transcriptional machinery, as well as post-transcriptional regulation of defense-associated mRNAs are essential for this phenomenon (Berr et al. 2012; Ma et al. 2011; Staiger et al. 2013).

Epigenetic gene regulation plays a critical role in cell differentiation and reprogramming. It also maintains genome integrity by silencing the expression of transposable elements (TEs) and other repeat sequences (Boyko and Kovalchuk 2011; Zaratiegui et al. 2007). Whether DNA is packaged as transcriptionally active euchromatin or transcriptionally silent heterochromatin is determined by the interplay between repressive epigenetic marks, such as DNA methylation and specific histone modifications, and chromatin remodeling complexes that position, evict, or alter the composition of nucleosomes (Boyko and Kovalchuk 2008; Ma et al. 2011). Evidence supporting a role for epigenetic regulation of plant immune responses comes from the combined demonstrations that i) loss of histone-modifying enzymes, including histone deacetylase (Kim et al. 2008; Wang et al. 2010), histone methyltransferases (Alvarez-Venegas et al. 2007; Berr et al. 2011; De-La-Peña et al. 2012; Palma et al. 2010), or histone ubiquitin-ligase (Dhawan et al. 2009), alters defense-gene expression and resistance to bacterial and fungal pathogens, ii) SA treatment leads to altered histone methylation and acetylation in key defense genes, including PR-1 (Alvarez-Venegas et al. 2007; Mosher et al. 2006), iii) mutations in several members of the SWI/SNF family of ATP-dependent
chromatin remodeling factors, including DDM1 (deficient in DNA methylation 1), ARP6 (actin-related protein 6), and PIE1 (photoperiod-independent early flowering 1), either alter defense gene expression, influence plant immunity, or both (Cheng et al. 2013; Down et al. 2012; Li et al. 2010; March-Diaz et al. 2008), and iv) mutants lacking components in the RNA-directed DNA methylation (RdDM) pathway that mediates epigenetic DNA methylation exhibit altered expression of numerous pathogen-responsive genes and enhanced resistance to virulent, avirulent, and nonpathogenic strains of *P. syringae pv. tomato* (Down et al. 2012; Yu et al. 2013).

TEs have long been considered transcriptionally inactive junk DNA (Lewin 1986; Probst and Almouzni 2011). However, several recent reports indicate that dynamic changes in TEs can occur, particularly in response to stress (Grandbastien et al. 2005; Ito et al. 2011; Tittel-Elmer et al. 2010). For example, the first comprehensive stress-induced DNA methylation map of the *Arabidopsis* genome indicated that TEs are differentially methylated in response to biotic stress or SA treatment (Down et al. 2012). Moreover, demethylation of TEs following treatment with the bacterial PAMP flag22 correlated with the downregulation of components involved in RdDM and the concomitant activation of TE expression (Yu et al. 2013). SA treatment also led to decreased methylation of some TEs and this corresponded not only with their upregulation, but also with increased expression of neighboring genes (Down et al. 2012). In addition, a wide range of TEs that are transcribed in response to stress with stress-responsive transcription, including ONSN and mPing, have been implicated in inducing neighboring genes (Ito et al. 2011; Makarevitch et al. 2015; Naito et al. 2009; Yasuda et al. 2013). Together, these observations suggest that TEs may play a role in triggering the transcriptional activation of proximal defense genes.

Efforts to elucidate the components required for resistance to *Turnip crinkle virus* (TCV) in *Arabidopsis* led us to identify the *CRT1* (compromised recognition of TCV) gene (Kang et al. 2008). Sequence analysis revealed that *CRT1* contains the ATPase and S5 domains characteristic of microchidia (MORC) proteins (Kang et al. 2010). Additional members of the *Arabidopsis* CRT1 family have been identified by other groups and variously named MORC, DMS, or CRT/CRH (Brabbs et al. 2013; Lorković et al. 2012; Moissiard et al. 2012). To avoid further confusion, we use the nomenclature proposed in a previous report and refer to CRT1 and its family members as MORC1-7 (Langen et al. 2014). Through genetic and biochemical analyses, we and others demonstrated that MORC1 is required for multiple layers of immunity, including ETI and PTL, following inoculation by a wide range of plant pathogens (Kang et al. 2008, 2010, 2012; Wang et al. 2011). In addition, MORC1 is one of very few proteins known to date that physically associates with a large number of immune components, including at least 12 R proteins and the PAMP recognition receptor FLS2 (Langen et al. 2014; Kang et al. 2008, 2010, 2012).

The demonstration that a small subpopulation of MORC1 is present in the nucleus and that nuclear MORC1 levels increase following activation of PTI or ETI in *Arabidopsis* (Kang et al. 2012) raised the possibility that this protein also has a nuclear function during plant immunity. Consistent with this hypothesis, MORC proteins from a wide variety of eukaryotes and prokaryotes have been implicated in either DNA recombination and repair, chromatin modification, or both (Iyer et al. 2008; Perry and Zhao 2003). Furthermore, MORC1 binds nucleic acids and exhibits Mn2+-dependent endonuclease activity (Kang et al. 2012) as well as ATPase activity (Kang et al. 2008). Mouse MORC1, the first MORC protein identified, is required for meiotic nuclear division (Watson et al. 1998); more recently, it was implicated in TE repression in the male gamline (Pastor et al. 2014). Suggesting a similar role for *Arabidopsis* MORC proteins, one or both MORC1 and MORC6 were identified in three independent screens for mutants defective for gene silencing (Lorković et al. 2012; Moissiard et al. 2012). The de-repression of silenced reporter genes and TEs in *morc1* and *morc6* mutants suggests that these proteins are required for epigenetic gene silencing (Brabbs et al. 2013; Lorković et al. 2012; Moissiard et al. 2012). However, it is currently unclear whether MORC1 and MORC6 fulfill their functions by modulating RdDM (Brabbs et al. 2013; Lorković et al. 2012) or by influencing chromatin structure (Moissiard et al. 2012).

In this study, we used both DNase-Seq (Hesselberth et al. 2009) and ChIP (chromatin immunoprecipitation)-Seq (Furey 2012) approaches to investigate the potential role MORC proteins play in regulating gene silencing and the mechanism through which this process impacts plant immunity. Analyses of DNase hypersensitive sites (DHSs) in mock- and *P. syringae pv. tomato*–inoculated wild type (WT) and *morc1/2* plants revealed that an enriched proportion of differential DHSs (dDHSs) are located in TEs. Strikingly, TE-associated dDHSs induced by *P. syringae pv. tomato* infection were identified near many defense-related genes. This finding, combined with the demonstration that i) MORC1 is a chromatin-interacting protein, ii) *P. syringae pv. tomato* infection increases MORC1 binding to *P. syringae pv. tomato*–induced dDHSs associated with genes and TEs, and iii) expression of many defense genes is delayed, reduced, or both in *morc1/2* mutant plants, suggests that MORC proteins modulate plant immune responses by binding TEs and thereby influencing both their expression and that of proximal genes following pathogen infection.

**RESULTS**

Pathogen infection did not dramatically alter the percentage of promoter-associated DHSs in either WT or MORC-deficient *Arabidopsis*.

To characterize changes in the chromatin landscape in response to pathogen infection, we prepared and sequenced DNase-Seq libraries from three genotypes: i) WT, ii) *morc1-2 morc2-1 morc6-3* (morc1/2) (Kang et al. 2010), a double knock-out mutant lacking MORC1 and its closest homolog MORC2, and iii) *morc1-2 morc2-1 morc6-3* (morc1/2/6) (Moissiard et al. 2014), a triple knock-out mutant in which morc6 was introduced into the morc1/2 background. Plants from each genotype were subjected to four different treatments: no treatment (naïve) or inoculation with bacterial DNA. The Pearson correlation coefficient between biological replicates for each combination ranged from 0.76 to 0.97; only four of the 36 libraries were lower than 0.85. Given that independently grown batches of plants were used for the biological triplicates, to reduce false signals, these correlation coefficient values suggest high reproducibility of the data sets (Supplementary Table S2).

A total of 29,450 DHSs were identified in the *Arabidopsis* genome (Fig. 1; Supplementary Table S3). As expected, the distribution of DHSs was biased toward euchromatin over heterochromatin, clearly displaying a strong correlation with the density of genes (Supplementary Fig. S1). Among the DHSs, approximately 65% were located within 1 kb upstream of the transcription start sites (TSSs) or 1 kb downstream of the transcription termination site (TTSs) of protein-coding genes, approximately 18% were associated with the gene body, such
as exons, introns, 5’ and 3’ untranslated regions (UTRs), approximately 11% were located in sequences 1 kb upstream or downstream of TEs, approximately 1% were detected within TEs, and the remaining approximate 5% were in intergenic regions (Fig. 1A; Supplementary Table S4). For protein-coding genes the majority of DHSs (64%) were detected within 1 kb upstream of their TSSs. Similarly, 78% of the DHSs associated with TEs were located in this region. These data suggest that the identified DHSs are heavily enriched in promoter regions, which are normally located within 1 kb upstream of the TSS. Pairwise comparisons among untreated naïve plants and mock- or pathogen-inoculated WT, morc1/2, or morc1/2/6 mutants revealed no dramatic alterations in percentage of DHS at different genomic locations (Fig. 1).

d DHSs induced by 
P. syringae pv. tomato infection or loss of MORC family members were enriched in the promoters of TEs.

To assess chromatin dynamics in response to one or both pathogen infection or MORC1/2/6 deficiency, we performed pair-wise comparisons of the DHSs identified in the different plant backgrounds in the presence or absence of pathogen infection. The dDHSs identified by this process represent genomic sites at which the level of chromatin accessibility differs within the corresponding comparison. Pathogen infection induced significant changes in chromatin accessibility, since comparisons of DHSs between mock- and pathogen-infected plants within the WT, morc1/2, or morc1/2/6 backgrounds revealed hundreds of dDHSs (Fig. 2A; Supplementary Table S5). By contrast, pairwise comparisons of DHSs in naïve versus mock or avirulent versus virulent 
P. syringae pv. tomato identified 17 or fewer dDHSs (Fig. 2A; Supplementary Table S6), indicating that mock inoculation does not substantially alter the DHS pattern present in untreated naïve plants and that the DHSs induced by virulent and avirulent 
P. syringae pv. tomato are very similar.

To further investigate whether MORC proteins influence chromatin accessibility, we compared the DHSs identified in different genotypes subjected to the same treatment. Modest numbers of dDHSs, ranging from 59 to 118 (Fig. 2B), were observed in WT versus morc1/2 and WT versus morc1/2/6 comparisons in which the plants were responding to the same treatment. By contrast, the DHS comparisons between morc1/2 and morc1/2/6 identified very few dDHSs, with a maximum of five. This latter finding is in line with i) RNA-Seq analyses of morc1/2 and morc1/2/6, which revealed very little difference between these lines (Moissiard et al. 2014), and ii) our findings that morc1/2 and morc1/2/6 exhibit comparable levels of enhanced susceptibility to virulent and avirulent 
P. syringae pv. tomato (Supplementary Fig. S2). Given the small number of dDHSs observed in comparisons between i) naïve and mock,
ii) avirulent and virulent *P. syringae*, and iii) *morc1/2* and *morc1/2/Pst*, only analyses derived from combinations of mock- or *P. syringae*–inoculated WT plants (WT<sub>mock</sub> and WT<sub>Pst</sub>) and mock- or *P. syringae* pv. *tomato*–inoculated *morc1/2* (morc1<sub>Pst</sub>/2 and morc1<sub>Pst</sub>2) will be presented hereafter (Fig. 2).

Comparisons between the DHSs identified in mock- and pathogen-infected plants within the same genetic background (termed *P. syringae* pv. *tomato*–induced dDHSs) revealed a substantially larger number of dDHSs in genes than in TEs (Fig. 3A). By contrast, DHS comparisons between WT and *morc1/2* plants subjected to the same treatment (termed *morc1/2*–enhanced dDHSs) identified much lower numbers of dDHSs overall, with slightly more present in TEs than in genes. Interestingly, when the percentage of dDHSs associated with TEs (termed TE-dDHSs) was calculated for each pairwise comparison, a significantly greater number of TE-dDHSs was observed than would be statistically expected (Fig. 3B, indicated as a broken line). Approximately 21% of the *P. syringae* pv. *tomato*–induced dDHSs were associated with TEs in both WT and *morc1/2* backgrounds (Fig. 3B). Notably, over 60% of the *morc1/2*–enhanced dDHSs were associated with TEs, irrespective of infection. These striking results suggest that MORC family members participate in modulating the physical accessibility of TE-associated sequences.

To gain insights into the putative biological functions of the gene-associated dDHSs, their gene annotations were analyzed using The Arabidopsis Information Resource (TAIR) database (Fig. 3C; Supplementary Table S7). Of the *P. syringae* pv. *tomato*–induced dDHSs, a significant number were associated with ‘response to stress’ or ‘response to biotic/abiotic stress’ genes (Fig. 3C, black and dark gray bars). Due to the low number of the *morc1/2*–enhanced gene dDHSs (43 or fewer), little significance was found in their gene annotations (Fig. 3C).

Members of the MORC family modulated the DNase I accessibility of heterochromatic TEs, while *P. syringae* pv. *tomato* infection altered the accessibility of TEs distributed throughout the genome.

TEs are predominantly localized in heterochromatin; these heterochromatic TEs are subject to transcriptional gene silencing primarily via RdDM (Matzke and Mosher 2014). By contrast, the silencing of euchromatic TEs, which also involves DNA methylation, appears to be mediated by a distinct mechanism (Zemach et al. 2013). For instance, *ddm1* mutant plants mainly lose repression of heterochromatic but not euchromatic TEs. To assess the location of the TE-dDHSs identified in the four comparisons, we plotted their genomic distribution against the relative density of genes and TEs (Fig. 4). Interestingly, the *morc1/2*–enhanced TE-dDHSs were preferentially associated with heterochromatin (Fig. 4, horizontal tracks 3 and 4), whereas the *P. syringae* pv. *tomato*–induced TE-dDHSs were more evenly distributed across the genome (Fig. 4, tracks 5 and 6). There is little, if any, overlap between the TE- or gene-associated dDHSs induced by *P. syringae* pv. *tomato* inoculation and those induced by the loss of MORC family members (Supplementary Fig. S3).

DNase I-quantitative polymerase chain reaction (qPCR) analysis confirmed the chromatin accessibility of selected TE-dDHSs identified by DNase-Seq.

Nine dDHSs associated with TEs, either in heterochromatin or in euchromatin, were randomly selected and were analyzed using DNase I qPCR to further assess the reproducibility of the DNase-Seq dataset. The *P. syringae* pv. *tomato*–inducible TE-dDHS AP8539 was additionally chosen, as it mapped to a site proximal to the well-known defense gene *PR-1* (Fig. 5A). This TE-dDHS spans a large region that encompasses two dDHSs, one resides in the promoter of a TE approximately 2 kb upstream of the *PR-1* gene (designated as AP8539b) and the other is located within the *PR-1* promoter (designated as AP8539a). Thus, we monitored chromatin accessibility at both sites. Using DNA prepared from DNase I-treated nuclei, qPCR was performed for the selected TE-dDHSs. Note that, while DNase-Seq identifies the ends of DNAs that are cut with DNase I, DNase I qPCR amplifies DNAs that are not disrupted by DNase I. Consequently, less amplification via qPCR corresponds to increased genome accessibility. Comparisons between the DNase-Seq reads and DNase I qPCR analyses for *morc1/2*-enhanced TE-dDHSs (Fig. 5B) and for *P. syringae* pv. *tomato*–induced TE-dDHSs (Fig. 5C) indicated that the results obtained from both
techniques were consistent. The only exception was AP4254, which exhibited greater accessibility in morc1/2 plants than in WT plants following DNase-Seq but only a little difference between these plants following DNase I-qPCR.

Defense genes were overrepresented in the genomic regions proximal to *P. syringae* pv. *tomato*-induced TE-dDHSs and their induction by *P. syringae* pv. *tomato* was either delayed, reduced, or both in morc1/2.

Decreased methylation of some TEs in response to SA was shown to increase expression of neighboring genes (Dowen et al. 2012), suggesting that derepression of TEs might trigger the transcriptional activation of proximal genes. Therefore, an- et al. 2012), suggesting that derepression of TEs might trigger the transcriptional activation of proximal genes. Therefore, an-

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**Fig. 4.** Effects of MORC1 and infection on DNase I accessibility of heterochromatic transposable elements (TEs). The relative densities of genes and TEs are presented in the top two tracks and the genomic distribution of TE-associated differential DNase I hypersensitive sites (dDHSs) found in the four pairwise comparisons are shown in the lower four tracks. The y axis indicates the number of dDHS in a 100-kb window. For each comparison, the DHSs enhanced in the plants listed first are presented in blue, while the DHSs enhanced in the plant listed second are presented in red. Dotted green lines indicate the borders between chromosomes.

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**P. syringae** pv. *tomato* infection changed the profile of MORC1-TE interaction.

Although we previously showed that MORC1 binds DNA with little sequence specificity in vitro (Kang et al. 2012), it is plausible that some of the morc1/2-enhanced TE-dDHSs are due to the loss of protection by bound MORC proteins. To address this possibility, chromatin immuno-precipitation followed by Illumina-based DNA sequencing (ChIP-Seq) was used to assess the interaction between Myc-tagged MORC1 and chromatin. These analyses were performed using an anti-Myc antibody and mock- or *P. syringae* pv. *tomato*-inoculated plants from a transgenic Arabidopsis morc1/2 line carrying a Myc-tagged MORC1 transgene expressed under its own promoter (Myc-gMORC1). This transgene was previously shown to complement morc1/2 (Kang et al. 2012). To control for nonspecific background binding, ChIP-Seq was performed in parallel on mock- and *P. syringae* pv. *tomato*-inoculated WT plants. Note that each sample was prepared in three independent biological triplicates.

Analysis of the ChIP-Seq peaks from mock- and *P. syringae* pv. *tomato*–inoculated Myc-gMORC1 plants (Fig. 8A, the third and fourth tracks; Supplementary Table S10) revealed that MORC1 binds sites distributed throughout the genome, although it shows a strong preference for heterochromatic TEs. To find genomic regions exhibiting altered levels of MORC binding after pathogen infection, the intensity of ChIP-Seq peaks from mock- and *P. syringae* pv. *tomato*–inoculated plants was compared. Differential ChIP-Seq peaks (dChIP peaks) exhibiting increased or decreased (Fig. 8A, induced and suppressed, blue lines and red lines, respectively, in the fifth track) intensity after *P. syringae* pv. *tomato* infection as compared with mock infection were not primarily associated with heterochromatic sites but, instead, were dispersed throughout the genome (Fig. 8A; Supplementary Table S11). Given that *P. syringae* pv. *tomato*–induced and morc1/2-enhanced dDHSs are enriched in TEs (Fig. 3), we assessed whether the genomic regions exhibiting altered interaction with MORC1 after pathogen infection correspond to TEs. Indeed, over 70% of the dChIP peaks that showed either increased or decreased MORC1 binding were TE-associated (Supplementary Fig. S5). Together, these results suggest that MORC1 is a chromatin-interacting protein that displays a significant preference for TEs and exhibits altered binding affinity after pathogen infection. Interestingly, the MORC1-associated peaks identified by ChIP-Seq were relatively broad, averaging around 8.5 kb. This contrasts with the typically narrow peaks associated with transcription
factors and suggests that MORC1 exhibits little sequence preference but, instead, may recognize higher-order chromatin structure.

Whether the MORC1-chromatin interaction sites overlap one or both the morc1/2-enhanced or *P. syringae pv. tomato*-induced dDHSs was then assessed. All of the gene- and TE-associated dDHSs were checked for overlaps with the dChIP peaks; however, dDHSs located in the intergenic regions were excluded. The background level of overlap (Fig. 8B, red line) was determined by calculating the percentage of total gene- or TE-dDHSs in each group that overlap with *P. syringae pv. tomato*–suppressed or *P. syringae pv. tomato*–induced dDHS peaks. Interestingly, dChIP peaks that were reduced after *P. syringae pv. tomato* infection correlated most strongly with morc1/2-enhanced TE-dDHSs, although a lower but statistically significant percentage overlapped with gene-associated dDHSs detected in morc1/2 versus WT plants after *P. syringae pv. tomato* infection. In comparison, the dChIP peaks that were enhanced by *P. syringae pv. tomato* infection showed a specific overlap with *P. syringae pv. tomato*–induced but not morc1/2-enhanced dDHSs located in genes. A low but statistically significant overlap between *P. syringae pv. tomato*–induced and morc1/2-enhanced TE-dDHSs also was observed. Given the connection between

Fig. 5. Quantitative polymerase chain reaction (qPCR) analysis of selected differential DNase I hypersensitive sites (dDHSs) identified by DNase-Seq. A, Genomic locations of nine randomly selected transposable element (TE)-associated dDHSs as well as TE-associated dDHSs near the defense gene PR-1 (AP8539). dDHSs located in heterochromatic and euchromatic positions are indicated by blue and red arrows, respectively. A dotted green vertical line denotes the border between chromosomes. B and C, Schematic presentation of *Arabidopsis* genes and TEs, with their corresponding genome coordinates in kilobases, as well as DNase-Seq read densities of TE-associated dDHSs (indicated as red boxes) in mock- and *Pseudomonas syringae pv. tomato* (Pst)-inoculated wild type (WTmock and WT*Pst*) and mock- and *Pseudomonas syringae pv. tomato*-inoculated morc1/2 (morc1/2mock and morc1/2*Pst*) and qPCR analysis. B, Selected morc1/2-induced dDHSs. C, Selected *P. syringae pv. tomato* infection-induced dDHSs. The relative amount of each dDHS was determined through qPCR analysis using *Tip41-like* as a reference gene (represented in the y axis). Note that DNase I-qPCR amplifies DNA that is not disrupted by DNase I; thus, lower levels of amplification indicate increased genome accessibility. Two repeats were performed for each dDHS. Statistical significance was determined using a Student’s t test; * indicates $P < 0.05$ and # indicates $P < 0.01$. 
MORC1 and TEs, we assessed whether cryptic TEs are present in the gene-d DHSs that overlap chromatin sites displaying increased MORC1 binding after *P. syringae* pv. *tomato* infection. Interestingly, cryptic TEs were identified in all of these gene-d DHSs (Supplementary Fig. S6), suggesting that MORC1 interacts with these previously unidentified TEs. In summary, our findings suggest that *P. syringae* pv. *tomato* infection leads to reduced MORC1 binding at d DHSs that are preferentially associated with heterochromatic TEs while, in contrast, infection enhances MORC1 binding at *P. syringae* pv. *tomato*-induced d DHSs located in a small population of euchromatic and heterochromatic TEs as well as in genes, likely via unannotated cryptic TEs.

**Silencing of *P. syringae* pv. *tomato*-induced TE-d DHSs proximal to a MORC1 binding region compromised induction of adjacent defense genes.**

Although MORC1/2 physically associates with some *P. syringae* pv. *tomato*-induced TE-d DHSs (Fig. 8B), none of the TE-d DHSs neighboring the defense genes monitored in Figure 7 directly overlap *P. syringae* pv. *tomato*-induced dCHiP peaks. However, a majority of these TE-d DHSs are within 250 kb of a *P. syringae* pv. *tomato*-induced dCHiP peak (Supplementary Table S12). Based on these results, we hypothesized that, following *P. syringae* pv. *tomato*-induced binding of MORC1 at adjacent sites, these TE-d DHSs are local enhancers that upregulate the expression of their neighboring defense genes. Suppression of these enhancers by RNAi-mediated silencing would, therefore, be expected to interfere with proximal gene induction. To test this hypothesis, transgenic *Arabidopsis* lines expressing a hairpin construct (Wesley et al. 2001) targeting each of seven TE-d DHSs shown in Figure 7 were generated. Five of these TE-d DHSs were located within 250 kb of one or more infection-induced dCHiP peak; the other two d DHSs that were not proximal to a dCHiP peak were chosen as controls. Silencing of the targeted regions was assessed via *McrBC*-qPCR, which involves qPCR of DNA digested with or without *McrBC*, an enzyme that specifically cuts methylated DNA. This quantitative analysis to test whether DNA methylation is induced by a hairpin RNA verified that all seven lines have significant DNA methylation in the intended regions (Fig. 9A). Silencing the putative TE-associated enhancers proximal to a *P. syringae* pv. *tomato*-induced dCHiP peak significantly reduced the induction of four of the five neighboring defense genes, including *PR-1*, *At2g06050*, *At4g19230*, and *At2g27690* but did not impact *At3g44300* (Fig. 9B). By contrast, silencing the TE-d DHSs lacking an adjacent *P. syringae* pv. *tomato*-induced dCHiP peak had little impact on the induction of defense genes *At2g19230* and *At4g16760* (Fig. 9B). These results support the possibility that these MORC1-associated TEs are local enhancers that induce adjacent defense gene expression.

**DISCUSSION**

To address the relationship between the roles MORC proteins play in gene silencing and plant immunity, we mapped the genomic location of DHSs in mock- and pathogen-inoculated WT and MORC-deficient plants. In a previous study, approximately 45% of the DHSs in the *Arabidopsis* genome were mapped to putative promoter regions (Zhang et al. 2012). Underscoring the correlation between DHSs and transcriptionally active chromatin, these sites were depleted of nucleosomes and were tightly associated with RNAPII binding sites. Furthermore, the prevalence of DHSs was dramatically reduced in transcriptionally silent pericentromeric regions that contain highly methylated DNA. Similar to these findings, the majority of DHSs we identified were located within 1 kb upstream of the TSS of protein-coding genes. Analysis of the TE-DHSs also revealed that the majority also were located within 1 kb of the TSS. Neither *P. syringae* pv. *tomato* infection nor loss of MORC family members substantially altered the genomic distribution of DHSs. Given that DNase I sensitivity is influenced by DNA methylation levels (Zhang et al. 2012) and that genome-wide DNA methylation levels are not appreciably altered in *morc1* or *morc6* mutants as compared with WT plants (Moissiard et al. 2012) or in *P. syringae* pv. *tomato*– versus mock-inoculated *Arabidopsis* (Dowen et al. 2012), our DHS results are consistent with these methylation studies.

Although global DHS distribution was not affected by *P. syringae* pv. *tomato* inoculation or the loss of MORC family members, pairwise comparisons of the DHSs detected in mock- or *P. syringae* pv. *tomato*–inoculated WT or *morc1/2* plants revealed notable differences. To our surprise, however, little difference was observed in d DHSs between virulent and avirulent *P. syringae* pv. *tomato* (Fig. 2A), while considerably more d DHSs were found for *P. syringae* pv. *tomato* versus mock than for avirulent *P. syringae* pv. *tomato* versus mock, suggesting that infection with virulent *P. syringae* pv. *tomato* alters genome accessibility more than avirulent *P. syringae* pv. *tomato* infection. However, since host responses to avirulent and virulent infection are often quantitative (Tao et al. 2003) and the analysis was done with only one time point, additional time points will likely be required to assess the quantitative nature of avirulent versus virulent infection-induced genome accessibility. It is noteworthy that a large number of d DHSs found both in a *P. syringae* pv. *tomato* versus mock and in an avirulent *P. syringae* pv. *tomato*
pv. tomato versus mock comparisons, when the DHSs were compared between these virulent and avirulent P. syringae pv. tomato, the differences in their DHSs were marginal.

The number of morc1/2-enhanced dDHSs was much lower than that induced by P. syringae pv. tomato infection (Fig. 3). Of these morc1/2-enhanced dDHSs, fewer than 40% were located in genes. Instead, morc1/2-enhanced dDHSs were highly enriched in TEs, particularly those located in heterochromatic regions (Fig. 4). Analysis of the genomic regions identified by ChIP-Seq confirmed that MORC1 preferentially binds heterochromatic and TE-associated regions (Fig. 8A). Furthermore, since morc1/2-enhanced TE-dDHSs overlap the genomic regions corresponding to P. syringae pv. tomato—suppressed dChIP peaks at a rate substantially greater than that of the expected background, P. syringae pv. tomato infection appears to lead to lowered MORC1/2 binding at heterochromatic TEs (Fig. 8B). It is interesting to note that a majority of the TEs transcriptionally activated in morc1, morc6, and morc1/6 (Moissiard et al. 2012) are physically located close (less than 160 kb) to morc1/2-enhanced TE-dDHSs. Together, these results are consistent with the reported association of MORC1 and MORC6 with heterochromatin (Moissiard et al. 2012) and the derepression of multiple families of TEs and endogenous genes preferentially associated with heterochromatin in one or more morc1, morc2, or morc6 mutants (Moissiard et al. 2012; Brabbs et al. 2013; Moissiard et al. 2014).

Given the considerable interaction between MORC1 and heterochromatin (Fig. 8A), it seems surprising that dramatic differences in chromatin accessibility, as determined by the number of dDHSs, were not observed in comparisons between morc1/2 and WT plants receiving the same mock or P. syringae pv. tomato treatment (Figs. 2 and 3). One possible explanation...
for the discrepancy in these findings is that the development of DHSs in the morc1/2 background may be partially suppressed by one or more functionally redundant members of the MORC family. Indeed, MORC3, the next closest homolog of MORC1 after MORC2, appears to be functionally redundant with MORC1 to some degree, since it restored TCV coat protein–induced cell death in morc1-1 plants expressing an inducible coat protein transgene (Kang et al. 2008) (Supplementary Fig. S7). Unfortunately, we cannot test whether MORC3 also suppresses DHS development in the morc1-2 background, because the morc3-1 knock-out mutation (SALK_000009) is lethal in the homozygous state (Kang et al. 2010).

An alternative possibility is that MORC1/2 does not directly bind DNA but, instead, influences chromatin accessibility by directly or indirectly interacting with other proteins. For example, MORC family members may influence DNase I hypersensitivity in heterochromatic regions by interacting with proteins involved in RdDM. Indeed, the combined observations that i) derepression of reporter genes in morc6 mutants correlates with a decrease in their DNA and histone methylation levels (Brabbs et al. 2013; Lorković et al. 2012) and ii) MORC1, MORC2, MORC6, singly or in combination, interact with several proteins involved in the RdDM pathway, including for the discrepancy in these findings is that the development of DHSs in the morc1/2 background may be partially suppressed by one or more functionally redundant members of the MORC family. Indeed, MORC3, the next closest homolog of MORC1 after MORC2, appears to be functionally redundant with MORC1 to some degree, since it restored TCV coat protein–induced cell death in morc1-1 plants expressing an inducible coat protein transgene (Kang et al. 2008) (Supplementary Fig. S7). Unfortunately, we cannot test whether MORC3 also suppresses DHS development in the morc1-2 background, because the morc3-1 knock-out mutation (SALK_000009) is lethal in the homozygous state (Kang et al. 2010).

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Fig. 8. MORC1 is physically associated with infection-induced transposable elements–differential DNaSe I hypersensitive sites (TE-dDHSSs). Myc-gMORC1 plants and wild type (WT) were mock–or P. syringae pv. tomato (Pst)-infected (5 × 10^7 CFU/ml) for 1 day. Chromatin immunoprecipitation (ChIP) was performed using an anti-Myc antibody, and the recovered DNAs were sequenced, using Illumina Hi-Seq 2500. SICER (Zang et al. 2009) was used to identify peaks that are associated with MORC1 in three independent biological replicates; WT was used as a background control. A, The relative densities of genes and TE are presented in the top two tracks. The genome position of ChIP-Seq peaks (P < 0.01) in mock- and P. syringae pv. tomato–infected Myc-gMORC1 plants as compared with the WT background control are presented in the third and fourth tracks. The y axis indicates the number of ChIP peaks in a 100-kb window. ChIP-Seq peaks (false discovery rate < 0.001) for which levels are substantially different in mock- versus P. syringae pv. tomato–inoculated plants (termed differential ChIP peaks) are presented in the fifth track. B, Percentage of dDHSs peaks overlapping with ChIP peaks were calculated to show association of MORC1 with dDHSs. To better display TE-association with MORC1, gene- and TE-dDHSSs were separately analyzed. A background level of overlap between ChIP and dDHS peaks was calculated using total gene- (n = 24,340) and TE-DHSs (n = 3,520). Statistically significant overrepresentation was determined between indicated groups and all DHSs using one sample χ^2 test; # indicates P < 0.01.

Fig. 9. Silencing of Pseudomonas syringae pv. tomato–induced transposable elements–differential DNase I hypersensitive sites (TE-dDHSSs) proximal to a MORC1 binding site compromised induction of adjacent defense genes. A, RNA interference–mediated silencing of selected TE-dDHSSs in transgenic Arabidopsis plants carrying a hairpin construct (hp) was verified via McrBC-qPCR (quantitative polymerase chain reaction); wild type (WT) plants were used as a control. The relative amount of McrBC-digested DNA in comparison with undigested DNA was analyzed in three replicates and is presented in the y axis. Note that higher levels of amplification indicate decreased DNA methylation. Several independent lines were generated for each hairpin construct. B, P. syringae pv. tomato–induced expression of defense genes proximal to silenced TE-associated dDHSSs was analyzed via quantitative reverse transcription (qRT)-PCR. Each panel corresponds to a representative transgenic line (hp) in which the TE-dDHSS adjacent to the indicated defense gene has been silenced by a hairpin construct; an empty vector (EV)-transformed line was used as a control. Leaves were inoculated with buffer (mock) or 10^6 CFU of P. syringae pv. tomato per milliliter. Untreated (naïve) control leaves were harvested at 0 h postinoculation. RNAs prepared from WT and morc1/2 plants at the indicated time points after inoculation were used for qRT-PCR. The relative amount of amplification indicates decreased DNA methylation. Several independent lines were generated for each hairpin construct. B, P. syringae pv. tomato–induced expression of defense genes proximal to silenced TE-associated dDHSSs was analyzed via quantitative reverse transcription (qRT)-PCR. Each panel corresponds to a representative transgenic line (hp) in which the TE-dDHSS adjacent to the indicated defense gene has been silenced by a hairpin construct; an empty vector (EV)-transformed line was used as a control. Leaves were inoculated with buffer (mock) or 10^6 CFU of P. syringae pv. tomato per milliliter. Untreated (naïve) control leaves were harvested at 0 h postinoculation. RNAs prepared from WT and morc1/2 plants at the indicated time points after inoculation were used for qRT-PCR analysis with primers specific for indicated genes. The TIP41-like gene was used as a reference gene for qRT-PCR and the mean ± standard error (n ≥ 3). A minimum of one more independent line was tested and found to have an expression pattern comparable to those presented here. Statistical significance from WT was determined using t test; * indicates P < 0.05 and # indicates P < 0.01.
DMS3 (defective in meristem silencing 3) (Lorković et al. 2012) and the SET domain-containing proteins SUVH9 and SUVH2 (Liu et al. 2014), suggest that MORC family members influence RdDM, which in turn could affect DNA methylation. The relationship between MORC proteins and RdDM, however, is currently unclear. Notably, older morc6 plants develop stochastic, cell-autonomous silencing of a green fluorescent protein (GFP) reporter gene that is consistently expressed in younger plants (Brabbs et al. 2013). Since this silencing of GFP expression was associated with the reappearance of DNA methylation in the reporter gene sequence, it was proposed that MORC6 promotes but is not obligated required for RdDM (Brabbs et al. 2013). Consistent with this proposition, analyses of morel and morc6 mutants failed to detect a correlation between changes in genome-wide methylation levels and the activation of a silenced reporter gene (Moissiard et al. 2012, 2014). Instead, since pericentromeric heterochromatin was decondensed in these mutants, it was proposed that MORC1 and MORC6 enforce the higher order compaction of methylated, silenced chromatin; such a function would, presumably, also modulate the DNase I sensitivity of these sequences. The very broad MORC1-associated peaks identified by our ChIP-Seq analysis also support this possibility.

In comparison with the morc1/2-enhanced dDHSs, those induced by *P. syringae* pv. *tomato* infection were primarily located in protein-encoding genes, particularly those associated with biotic or abiotic stresses. They were also enriched in TEs; these TE-dDHSs were spread throughout the genome and shared little overlap with those enhanced by morel/2 (Fig. 4). Strikingly, the 5′ flanking regions of a wide range of defense genes, including PR-1, PR-2, and PR-3, contained *P. syringae* pv. *tomato*-induced TE-dDHSs. Since the expression kinetics of these PR genes as well as 12 additional, randomly selected defense genes was generally modestly delayed or weaker in 12 additional, randomly selected defense genes was generally modestly delayed or weaker in *P. syringae* pv. *tomato*-inoculated morel/2 as compared with WT plants (Fig. 7), MORC1/2 appears to promote their expression. Consistent with this possibility, MORC1 association with *P. syringae* pv. *tomato*-induced TE-dDHSs was enhanced after *P. syringae* pv. *tomato* infection. Note that all of the *P. syringae* pv. *tomato*-induced gene dDHSs were found to be associated with unannotated TEs, essentially making them TE-dDHSs (Fig. 8B). It is interesting to note that infection-induced transcription of PR-1 and other defense genes was generally suppressed by RNAi-silencing of the neighboring (although in some cases distant) TE-dDHS or but only if the TE-dDHS was adjacent to a *P. syringae* pv. *tomato*-induced dChIP peak. Given that enhancers can be located as far as 1 Mb from the genes they regulate (Krivega and Dean 2012; Yang and Corces 2012), our silencing results suggest that i) at least some *P. syringae* pv. *tomato*-induced TE-dDHSs serve as enhancers of neighboring defense genes and ii) these putative enhancers are activated by *P. syringae* pv. *tomato*-induced MORC1/2 binding to sites that, while nearby, do not necessarily overlap with the dDHS. This latter conclusion may explain why the overlap frequency between dDHSs and ChIP peaks is relatively low (Fig. 8B). Moreover, this long-distance effect of MORC1 recruitment on defense genes underscores upcoming challenges in characterizing chromatin-interacting proteins such as MORC1 and its homologs in the modulation of associated genes. It also remains to be seen why MORC1/2 influenced defense genes in a limited fashion rather than globally and how this observation is related to the earlier report showing the stochastic silencing phenotype of a morc6 mutant (Brabbs et al. 2013).

Analyses of the genomic regions exhibiting differentially increased or decreased MORC1 binding following *P. syringae* pv. *tomato* infection indicated that they were strongly associated with TEs dispersed over the entire genome (Fig. 8A). This finding, combined with the discovery that cryptic TEs are present in most of the gene-associated dDHSs exhibiting increased MORC1 binding after *P. syringae* pv. *tomato* infection, suggests that MORC1 may regulate gene expression by binding the super structure associated with TEs or TE-like sequences. A long list of studies also have suggested a link between TEs and gene regulation. In humans, TEs and repeat DNA are associated with cell-specific transcription (Thurman et al. 2012). Furthermore, a substantial portion of regulatory elements in the promoters of human and plant genes are derived from TEs or pseudo or former TEs (Girard and Freeling 1999; Jordan et al. 2003; Martienssen et al. 1990). In *Arabidopsis*, TE density is enriched in the approximately 2 kb upstream of the TSS and near the 3′ end of genes whose induction by SA or *P. syringae* pv. *tomato* treatment is associated with the appearance of differentially methylated regions (Dowen et al. 2012). Analysis of these *P. syringae* pv. *tomato*– or SA-inducible genes containing differentially methylated regions revealed that many exhibit functions associated with plant immunity. A recent study also indicated that DNA demethylation of TEs is important for the activation of some defense genes (Le et al. 2014). Our finding that *P. syringae* pv. *tomato*–induced TE-dDHSs are associated with a wide range of defense genes further argues that TEs are important regulatory elements for controlling transcription and that their modulation by RdDM components, including MORC1, plays an important role in the activation of defense responses after pathogen attack.

In summary, we demonstrate that *P. syringae* pv. *tomato* infection primarily suppresses MORC1 binding at dDHSs associated with heterochromatic TEs but enhances its binding at infection-induced dDHSs in genes and TEs. These results, combined with MORC1’s previously demonstrated involvement in heterochromatin condensation and gene silencing (Brabbs et al. 2013; Lorković et al. 2012; Moissiard et al. 2012), and our finding that defense gene expression is attenuated in the morc1/2 mutant, suggest that MORC1 plays important roles in both gene silencing and gene induction. We propose that MORC1 mediates these divergent effects via its interaction with different chromatin-binding proteins. In this scenario, the *P. syringae* pv. *tomato*–induced loss of MORC1 at heterochromatic TEs would disrupt a complex involved in gene silencing, thereby leading to activation of TE expression after pathogen infection. By contrast, the *P. syringae* pv. *tomato*–induced addition of MORC1 to a protein complex present at TEs would temporarily relieve silencing, thereby promoting robust expression of proximal genes. In support of this model, growing evidence suggests that epigenetic regulation is often context specific (Sarris et al. 2014). For instance, enhancer of Zeste 2 homolog (EZH2), which is a core enzymatic subunit of the polycomb repressive complex 2 (PRC2) in human cells, exerts opposing effects on gene expression. As a part of PRC2, EZH2 is involved in silencing a wide range of genes; however, when EZH2 is posttranslationally modified, it functions as a coactivator for androgen receptor–target genes (Xu et al. 2012). Further characterization of MORC1’s interacting proteins, including epigenetic factors such as DMS3 (Lorković et al. 2012), SUVH9, and SUVH2 (Liu et al. 2014), will likely provide important insights into the mechanisms through which MORC1 impacts gene silencing, defense gene induction, and plant immunity.

**MATERIALS AND METHODS**

**Plant material, generation of transgenic lines, bacterial infiltration, and analysis of antibacterial resistance in *Arabidopsis***

Plants were grown in soil at 22°C, 60% relative humidity, and a 16-h light period. Plants 3.5 weeks old were syringe-infiltrated with *P. syringae* pv. *tomato* (5 × 10⁶ CFU/ml) or 10 mM MgCl₂ (mock) and treated leaves were harvested 1 day postinoculation for DNase-Seq, DNase-qPCR, and ChIP-Seq.
described below or at the indicated times for RNA analysis. Transgenic *Arabidopsis* lines were generated using a floral dipping method as previously described (Kang et al. 2008). To analyze antibacterial resistance in *Arabidopsis*, leaves of *Arabidopsis* plants were infiltrated with *P. syringae pv. tomato* with or without AvrRpt2 at a rate of 10^6 CFU/ml in 10 mm MgCl2, using a needleless syringe. Infected leaves were harvested at the given time points after the infiltration, and then, were used for bacterial titer determination, as previously described (Kang et al. 2008).

**Preparation for DNase-Seq and DNase-qPCR.**

Around 4.5 g of leaf tissue was harvested in a 50 ml tube and was incubated in diethyl ether for 3 min, followed by washing three times with cold water. The tissue was homogenized in 5 ml of the homogenization buffer at 4°C, using a T10 Ultra-Turrax homogenizer (IKA). Nuclei were enriched as previously described (Manzara and Gruissem 1995). In addition, the nucleus pellet was washed five times with the homogenization buffer to remove chloroplasts and *P. syringae pv. tomato*, followed by a Percoll gradient purification as described (Henfrey and Slater 1988).

Prepared nuclei were subject to DNase I digestion as described (Hesselberth et al. 2009), with the following modification: After concentration of nuclei was calculated using a hemocytometer under the microscope, 8 x 10^5 nuclei were incubated with one unit of DNase I (Roche) at 37°C for 10 min, which gave DNA cleavage comparable to that in the yeast DNase-Seq experiment (Hesselberth et al. 2009). DNA was size-fractionated, using a sucrose gradient as described (Hesselberth et al. 2009), to obtain DNA ranging from 100 to 700 bp. DNA was further purified using a Qiagen PCR purification kit (Roche). DNA quality was checked on a 1% agarose gel stained with SYBR green (Life Technologies) and scanned with a Typhoon Trio imager (GE Healthcare).

**Construction and sequencing of DNase I libraries.**

Each combination of genotype and treatment was prepared in biological triplicates, resulting in a total of 36 independent libraries. DNase I libraries were prepared using a Genomic DNA sample prep kit (Illumina) according to the manufacturer’s protocol. The libraries were sequenced on an Illumina Hiseq 2500 system at Cornell University Life Sciences Core Laboratories Center with single-end, 100 bp mode.

**DNase-Seq data and gene ontology analysis.**

DNase-Seq reads were first aligned to the *Arabidopsis* chloroplast genome using Bowtie (Langmead et al. 2009), allowing up to two mismatches, and those aligned were discarded. The resulting plast genome using Bowtie (Langmead et al. 2009), allowing up to two mismatches, and those aligned were discarded. The resulting plast genome using Bowtie (Langmead et al. 2009), allowing up to two mismatches, and only the best hits were kept. Only reads uniquely mapped (having one single best hit) to the genome were kept. The alignments of genomic features were i) TSS upstream, ii) TE upstream, iii) UTR, iv) exon or intron, v) TE, vi) TTS downstream, vii) TE downstream, and viii) intergenic. Then for each DNase-Seq library, the number of reads mapped to each of the identified DHSs was counted and normalized to reads per million mapped reads. The raw count information was fed to DESeq (Anders and Huber 2010) to identify differentially accessible DHSs upon infection by *P. syringae pv. tomato* in WT and MORC-deficient mutants with a cutoff of corrected P value < 0.05 and fold change >2.

To evaluate enrichment in the gene ontology categories, known or deduced biological functions of the genes associated with dDHs were annotated with TAIR (Berardini et al. 2004) as of January 2015. The broad biological categories ‘other biological processes,’ ‘other cellular processes,’ and ‘other metabolic processes’ were not presented in Figures 3 and 6. The raw P values were calculated using the hypergeometric distribution, which were then adjusted for multiple testings using the Benjamini-Hochberg procedure (Benjamini and Hochberg 1995).

**ChIP.**

ChIP was performed as described (Tai et al. 2005) with the following changes: 4.5 g of leaf tissue or sample was cross-linked for 60 min and nuclei were prepared as described above, except that the Percoll gradient step was skipped. Magnetic beads conjugated with protein A/G (Thermo Fisher Scientific) and α-myc (Abcam) antibodies were used to immunoprecipitate Myc-tagged MORC1 and its interacting chromatin. DNA was purified using the Qiagen PCR purification kit. ChIP-Seq libraries were generated using the NEBNext ChIP-Seq library prep reagent kit (New England Biolabs) according to the manufacturer’s instructions. The libraries were sequenced using an Illumina Hiseq 2500 system at the Genome Sequencing and Analysis Facility of The University of Texas at Austin.

**ChIP-Seq data analysis.**

ChIP-Seq raw reads were first processed to remove adapter and low-quality sequences using Trimmomatic (Bolger et al. 2014). The statistical elements of this analysis are shown in Supplementary Table S13 and Supplementary Table S14. For each library, identical reads were then collapsed into a single unique read. The resulting ChIP-Seq reads were aligned to the *Arabidopsis* genome (TAIR10 release) using Bowtie (Langmead et al. 2009), allowing up to one mismatch, and only the best hits were kept and only reads uniquely mapped (having one single best hit) to the genome were kept. The alignments from different biological replicates were combined using SAMtools (Li et al. 2009), and then were converted into browse extensible data (BED) format using BEDTools (Quinlan and Hall 2010). The read mapping coordinates in BED format were fed into SICER (Zang et al. 2009) (redundancy threshold at 1, window size at 600, fragment size at 150, effective genome fraction at 0.96, and gap size at 1,800,) to identify wide peaks from the two pairs of libraries, i.e., i) Myc-gMORC1 mock and WT mock and ii) Myc-gMORC1 Pst and WT Pst. SICER were run at the following parameters: redundancy threshold at 1, window size at 600, fragment size at 150, effective genome fraction at 0.96, and gap size at 1,800. SICER first identified significant peaks in each of the two pairs and, then, merged the two sets of peaks. Then for each merged peak, its level in *P. syringae pv. tomato* was compared with that in mock to determine the significance of changes. Significantly differential peaks were identified when the fold change of peak levels between *P. syringae pv. tomato* and mock should be larger than 1.1 and the false discovery rate should be less than 0.001.

Then for each ChIP-Seq peak, the number of mapped reads in each sample was counted and normalized to reads per kilobase per million mapped reads. Raw counts were then fed to edgeR (Robinson et al. 2010) to identify differential peaks between WT and Myc-gMORC1 plants under mock or *P. syringae pv. tomato* inoculation. ChIP-Seq peaks with fold changes of peak levels >1.1 and P value <0.01 were identified as significant and MORC1-associated.
qPCR. qPCR was used to quantify the DNA templates prepared from DNase I and ChIP experiments. Maxima SYBR green qPCR master mix (Thermo Fisher Scientific) was used, with initial incubation at 50°C for 2 min and at 95°C for 10 min, followed by 40 cycles of 95°C for 25 s and 60°C for 1 min. The primer set used for qPCR was listed in Supplementary Table S15. Level of target DNA was calculated from the difference of threshold cycle values between reference and target genes (Schmittgen and Livak 2008). The TIP41-like gene (Czechowski et al. 2005) was used as the reference for DNase I-qPCR.

Northern blot and qRT-PCR analysis. RNA was extracted using TRIzol (Life Technologies) following the manufacturer’s protocol and was quantified by NanoDrop 1000 (Thermo Fisher Scientific). Northern analysis was performed with the PR-1 probe as described (Kang and Klessig 2005) because PR-1 gene has no introns. Therefore, the PCR product made off the cDNA is not distinguishable from the PCR product produced off any small amount of contaminating genomic DNA, which could obscure real but subtle differences in expression levels (Fig. 7A). qRT-PCR analysis was performed as described (Kang et al. 2010); the TIP41-like gene (Czechowski et al. 2005) was used as a reference gene.

McrBC-qPCR. Overnight digestion of 500 ng of genomic DNA was performed with and without McrBC as suggested by the manufacturer (New England Biolabs). The digested sample was diluted 10-fold and was used for qPCR, which is described above.

DNA constructs for complementation and RNAi silencing. Constructs complementing morc1-1 were previously described (Kang et al. 2008). NheI-digested bacterial artificial chromosome DNA F23E13 carrying genomic MORC3 was cloned into pART27. This construct was used for the complementation of morc1-1. Harpin constructs for RNAi silencing were generated as described (Kang et al. 2003), using pHamibil (Wesley et al. 2001). The binary vector pART27 carrying the Nod1 cassette from pHamibil was then transformed into Arabidopsis.

Accession number. Raw DNase-Seq and ChIP-Seq reads have been deposited in the National Center for Biotechnology Information sequence read archive under accession number SRP055733.

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LITERATURE CITED


