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► To cite this version:

Thomas Roulé, Federico Ariel, Caroline Hartmann, Jose Gutierrez-Marcos, Nosheen Hussain, et al.. The lncRNA MARS modulates the epigenetic reprogramming of the marneral cluster in response to ABA. 2020. hal-03016338

HAL Id: hal-03016338

<https://hal.science/hal-03016338>

Preprint submitted on 20 Nov 2020

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The lncRNA *MARS* modulates the epigenetic reprogramming of the marneral cluster in response to ABA

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ABSTRACT

Clustered organization of biosynthetic non-homologous genes is emerging as a characteristic feature of plant genomes. The co-regulation of clustered genes seems to largely depend on epigenetic reprogramming and three-dimensional chromatin conformation. Here we identified the long noncoding RNA (lncRNA) *MARNERAL Silencing (MARS)*, localized inside the *Arabidopsis marneral* cluster, and which controls the local epigenetic activation of its surrounding region in response to ABA. *MARS* modulates the POLYCOMB REPRESSIVE COMPLEX 1 (PRC1) component LIKE-HETEROCHROMATIN PROTEIN 1 (LHP1) binding throughout the cluster in a dose-dependent manner, determining H3K27me3 deposition and chromatin condensation. In response to ABA, *MARS* decoys LHP1 away from the cluster and promotes the formation of a chromatin loop bringing together the *MARNERAL SYNTHASE 1 (MRN1)* locus and a distal ABA-responsive enhancer. The enrichment of co-regulated lncRNAs in clustered metabolic genes suggests that the acquisition of noncoding transcriptional units constitute an additional regulatory layer driving the evolution of biosynthetic pathways.

KEYWORDS: lncRNA, enhancer, cluster, chromatin conformation, LHP1, ABA, seed germination, epigenetics, marneral

INTRODUCTION

In eukaryotes, functionally related genes are usually scattered across the genome. However, a growing number of operon-like clustered organization of non-homologous genes participating in common metabolic pathways point at an emerging feature of animal, fungi and plant genomes¹.

In plants, synthesis of numerous secondary metabolic compounds is important for the dynamic interaction with their environment, affecting their life and survival². Terpenoids are bioactive molecules of diverse chemical structure³. In *Arabidopsis thaliana*, the biosynthesis of four triterpenes, namely thalianol⁴, tirucalla-7,24-dien-3b-ol⁵, arabidiol⁶ and marneral⁷, is governed by enzymes encoded by genes organized in clusters⁸. The thalianol and marneral related genes are located in the smallest metabolic clusters identified in plants to date, each being less than 40kb in size⁸. Both compounds are derived from 2,3-oxidosqualene and the corresponding gene clusters contain the oxidosqualene cyclases (OSCs), thalianol synthase (THAS) and marneral synthase (MRN1), respectively. The marneral cluster includes two additional protein-coding genes, *CYP705A12* and *CYP71A16*, participating in marneral oxidation⁷.

Growing evidence indicates that the co-regulation of clustered genes relies on epigenetic mechanisms. It has been shown that the deposition of the histone variant H2A.Z positively correlates with transcriptionally active clusters. Accordingly, nucleosome stability precluding gene expression is dependent on ARP6, a component of the SWR1 chromatin remodeling complex required for the deposition of H2A.Z into nucleosomes⁹. Additionally, it was shown that the thalianol and marneral clusters exhibit increased expression in the Polycomb mutant *curly leaf (clf)* with compromised H3K27me3 deposition, and reduced expression in the trithorax-group protein mutant *pickle (pkl)*, a positive regulator that counteracts H3K27me3 silencing¹⁰. Strikingly, it has been recently shown that biosynthetic gene clusters are embedded in local hot spots of three-dimensional (3D) contacts that segregate cluster regions from the surrounding chromosome environment in a tissue-dependent manner. Notably, H3K27me3 appeared as a central feature of the 3D domains at silenced clusters¹¹.

Long noncoding RNAs (lncRNAs) have emerged as important regulators of eukaryotic gene expression at different levels¹². In plants, several lncRNAs have been shown to interact with the Polycomb Repressive Complex 1 and 2 components LIKE HETEROCHROMATIN PROTEIN 1 (LHP1) and CLF, respectively, which are related to H3K27me3 distribution^{13,14}. Furthermore, it has been proposed that lncRNAs can modulate the transcriptional activity of

neighboring genes by shaping local 3D chromatin conformation^{15–17}. Here we show that the marneral cluster in *Arabidopsis* includes three noncoding transcriptional units. Among them, the lncRNA *MARS* influences the expression of marneral cluster genes in response to ABA through modification of the epigenetic landscape. *MARS* deregulation affects H3K27me3 distribution, LHP1 deposition and chromatin condensation throughout the cluster. Furthermore, an ABA responsive chromatin loop dynamically regulates *MRN1* transcriptional activation by bringing together the *MRN1* proximal promoter and an enhancer element enriched in ABA-related transcription factors (TF) binding sites. *MARS*-mediated control of the marneral cluster affects seed germination in response to ABA. The general co-regulation of genes located within lncRNA-containing clusters in *Arabidopsis* points to noncoding transcription as an important feature in coordinated transcriptional activity of clustered loci.

RESULTS

The marneral gene cluster includes three noncoding transcriptional units

The small marneral cluster includes three genes: marneral synthase (*MRN1*), *CYP705A12* and *CYP71A16* that are two P450 cytochrome-encoding genes (**Fig. 1a**), all participating in the biosynthesis and metabolism of the triterpene marneral⁷.

The advent of novel sequencing technologies has allowed the identification of an increasing number of lncRNAs throughout the *Arabidopsis* genome. According to the latest annotation (Araport 11¹⁸), three additional transcriptional units are located within the marneral cluster, between the *CYP71A16* and the *MRN1* loci. The *AT5G00580* and the pair of antisense genes *AT5G06325* and *AT5G06335* are located upstream of the *MRN1* gene at 6kpb and 3kbp, respectively (**Fig. 1a**). The 1,941bp-long *AT5G00580* locus generates four transcript isoforms ranging from 636 nt to 1,877 nt in length (**Fig. 1b and 1c**). In contrast, each of the antisense genes *AT5G06325* and *AT5G06335* are transcribed into only one RNA molecule of 509 nt and 367 nt, respectively (**Fig. 1a**). All these transcripts were classified as lncRNAs when using two coding prediction tools, CPC¹⁹ and CPC2²⁰, because of their low coding potential and their length (over 200 nt), similarly to previously characterized lncRNAs (*COLDIAIR*²¹; *APOLO*¹⁵; and *ASCO*²²) (**Fig. 1d**).

According to available transcriptomic datasets (Araport11), *AT5G00580* transcriptional accumulation positively correlates with that of marneral genes, whereas *AT5G06325* and *AT5G06335* RNAs do not (**Supplementary Fig. 1**). Notably, our analysis of

the transcriptional behavior of the noncoding gene *AT5G00580* and the marneral cluster protein-coding genes revealed a correlated expression in response to phosphate and nitrate starvation, heat stress, as well as to exogenous auxin and ABA (**Fig. 1e**). Interestingly, the *AT5G00580* lncRNA exhibited the strongest induction in response to heat stress and exogenous ABA, in comparison with *MRN1* and the two *CYP* genes (**Fig. 1e**). Altogether, our observations uncovered that the marneral cluster includes three noncoding transcriptional units, one of which is actively transcribed and co-regulated with its neighboring protein-coding genes.

The lncRNA *MARS* shapes the transcriptional response of the marneral gene cluster to ABA

It has been shown that lncRNAs can regulate the expression of their neighboring genes through epigenetic mechanisms^{14,23}. Thus, we wondered if the lncRNA derived from the *AT5G00580* locus may regulate the transcriptional activity of the protein-coding genes included in the marneral cluster. To this end, we modified the lncRNA expression without affecting the cluster DNA region using an RNAi construct targeting the first exon of *AT5G00580*, and isolated two independent lines. The RNAi line 1 impaired the transcriptional accumulation of *AT5G00580* without affecting the rest of the cluster. Interestingly, the RNAi line 2 exhibited a strong down-regulation of *AT5G00580* together with a significant basal repression of the two *CYP* genes (**Fig. 2a**). Strikingly, the response of the three protein-coding genes of the marneral cluster to exogenous ABA was significantly deregulated in both independent RNAi lines. Transcriptional levels of *MRN1* and the two *CYP* genes increased earlier in RNAi seedlings (15 min) than in the wild-type (Col-0, 30 min) (**Fig. 2b bottom panel**). In addition, the transcriptional accumulation of these genes later reached three-fold higher levels in the RNAi lines compared to Col-0 (**Fig. 2b top panel**). To further support our observations, we isolated the insertional mutant *SALK_133089* located 200 bp upstream the transcription start site (TSS) of *AT5G00580* gene. Although this insertion did not affect *AT5G00580* basal levels (**Fig. 2a**), it partially impaired its induction in response to ABA. In agreement with both RNAi lines, *MRN1* and *CYP705A12* genes were strongly induced by exogenous ABA (**Supplementary Fig. 2**), in contrast to *CYP71A16*, whose promoter region may be locally affected by the T-DNA insertion. Notably, *MRN1*, which encodes the marneral synthase, is the gene in the cluster most strongly affected by *AT5G00580* down-regulation. Collectively, our results indicate that the noncoding transcriptional activity derived from the *AT5G00580* locus represses the dynamic expression of the marneral cluster genes, mainly *MRN1*, in response to ABA. Therefore, we named the *AT5G00580*-derived noncoding transcript *MARneral Silencing (MARS)* lncRNA.

MARS affects seed germination likely through its impact on *MRN1* expression

The phytohormone ABA has been implicated in the perception and transduction of environmental signals participating also in a wide range of growth and developmental events such as seed maturation, dormancy and germination²⁴. Considering that the marneral cluster exhibited a strong *MARS*-dependent response to ABA, we wondered what the physiological impact of *MARS* deregulation was during seed germination. To this end, we assessed seed germination in Col-0 and *MARS* down-regulated lines (RNAi lines 1/2 and *SALK_133089*) with or without exogenous ABA. Notably, *MARS* silencing resulted in a delayed germination compared to Col-0 (**Supplementary Fig. 3a**), as revealed by an increase in T50 (time for 50% of germination) of nearly 5 hours (**Supplementary Fig. 3b**). Interestingly, in response to 0.5μM ABA, the germination of RNAi-*MARS* was further delayed than Col-0 with an increase of T50 of nearly 10 hours (**Supplementary Fig. 3c and 3d**). Accordingly, transgenic plants over-expressing *MRN1* also exhibited a delayed germination phenotype regardless of the treatment with ABA (**Supplementary Fig. 3**). The behavior of 35S:*MRN1* seedlings suggests that *MRN1* up-regulation in RNAi-*MARS* lines could be linked to the increased sensitivity to ABA (**Fig. 2b**). Altogether, our results indicate that *MARS* can modulate seed germination through the regulation of the expression of *MRN1*.

MARS controls the epigenetic status of the marneral locus

It has been shown that gene clusters in plants are tightly regulated by epigenetic modifications, including the repressive mark H3K27me3¹⁰. According to publicly available ChIP-Seq datasets²⁵, the marneral cluster region is highly enriched in H3K27me3 deposition in shoots, extensively coinciding with LHP1 recognition (**Supplementary Fig. 4**). Consistently, an ATAC-Seq available dataset²⁶ revealed that the marneral cluster exhibits a high chromatin condensation in shoots (**Supplementary Fig. 4**). Thus, the marneral cluster is characterized by an epigenetically silent status in aerial organs coinciding with its low expression level¹⁰.

We wondered if the transcriptional activation of the marneral cluster in response to exogenous ABA was associated with a dynamic epigenetic reprogramming. We first assessed H3K27me3 deposition across the marneral cluster, including the gene body of *MRN1*, *MARS* and the two *CYP* loci (**Fig. 3a and Supplementary Fig. 5**). Interestingly,

exogenous ABA triggered a strong reduction of H3K27me3 deposition throughout the marneral cluster (**Fig. 3a and Supplementary Fig. 5**). Strikingly, H3K27me3 basal levels were also significantly lower in RNAi-*MARS* seedlings. Remarkably, H3K27me3 deposition was even lower across the body of all genes of the cluster in response to ABA in the RNAi-*MARS* lines when compared with Col-0, in agreement with the stronger induction by ABA of this subset of genes upon *MARS* silencing (**Fig. 3a, Supplementary Fig. 5 and Fig. 2b**). Furthermore, we assessed LHP1 recognition of the marneral cluster. In agreement with previous observations (**Supplementary Fig. 4²⁵**), LHP1 distribution was high in *MRN1* promoter and more weakly across *MARS* gene body and the intergenic region between *CYP71A16* and *MARS* (**Fig. 3b and Supplementary Fig. 6**). Remarkably, LHP1 recognition was strongly impaired in response to ABA as well as in RNAi-*MARS* seedlings (**Fig. 3b and Supplementary Fig. 6**). Therefore, our results indicate that ABA triggers an epigenetic reprogramming of the marneral cluster, likely in a process involving the lncRNA *MARS*.

***MARS* is directly recognized by LHP1 and modulates local chromatin condensation**

It has been shown that the deposition of the repressive mark H3K27me3 and the concomitant recognition of the plant PRC1 component LHP1 are correlated with high chromatin condensation²⁷. Therefore, we determined by Formaldehyde-Assisted Isolation of Regulatory Elements (FAIRE) the chromatin condensation of the whole marneral cluster. In contrast to Col-0 showing a highly condensed chromatin, RNAi-*MARS* seedlings exhibit a lower chromatin condensation in control conditions. Notably, the global chromatin status of the cluster was even less condensed in RNAi-*MARS* seedlings in response to ABA (**Fig. 4a and Supplementary Fig. 7**), in agreement with a decrease of both H3K27me3 deposition and LHP1 binding (**Fig. 3b and Supplementary Fig. 6**) and the concomitant transcriptional activation of the clustered genes (**Fig. 2b**). Consistently with our observations, *lhp1* mutant seedlings also showed a global chromatin decondensation in control conditions, comparable to Col-0 in response to ABA. Notably, further chromatin decondensation triggered by ABA was completely impaired (**Fig. 4b and Supplementary Fig. 8**), supporting the role of LHP1 in the dynamic epigenetic silencing of the marneral cluster.

It has been shown that LHP1 can recognize RNAs *in vitro*¹³ and the lncRNA *APOLO* *in vivo*¹⁵. Moreover, it has been proposed that *APOLO* over-accumulation can decoy LHP1 away from target chromatin²⁸. Therefore, we wondered whether *MARS* lncRNA was able to interact with the chromatin-related protein LHP1 participating in the modulation of the local epigenetic environment. Thus, we first determined that *MARS* was enriched in the nucleus,

compared with total RNA, as the previously characterized lncRNAs *APOLO* and *ASCO* that interact respectively with nuclear epigenetic and splicing machineries, and the spliceosome structural ncRNA *U6* (**Supplementary Fig. 9a**). Then, we confirmed by RNA immunoprecipitation (RIP) that LHP1 can interact with *MARS* *in vivo*, in contrast to the *MRN1* mRNA or housekeeping RNAs taken as negative controls (**Fig. 4c**).

LHP1 binding to the marneral cluster was impaired both in response to exogenous ABA (inducing *MARS*, **Fig. 1e and Supplementary Fig. 3b**) and in RNAi-*MARS* seedlings, hinting at a stoichiometry-dependent action of *MARS* on LHP1 recognition. Therefore, we used chromatin extracts from RNAi-*MARS* line 1 seedling, with very low *MARS* transcript levels (**Fig. 2a**) to assess LHP1 recognition of the marneral cluster upon the addition of increasing concentrations of *in vitro*-transcribed *MARS* RNA. Strikingly, we found that low *MARS* RNA concentrations (between 0.01 and 0.1 μ g of RNA; **Fig. 4d and Supplementary Fig. 9b**) successfully promoted LHP1 binding to the cluster, in contrast to higher concentrations (between 1 and 10 μ g of RNA), supporting the relevance of *MARS*-LHP1 stoichiometry for LHP1-target recognition (**Fig. 4d and Supplementary Fig. 9b**). Altogether, our results suggest that the physical interaction of the nuclear-enriched lncRNA *MARS* to LHP1 modulates its binding to proximal chromatin likely participating in the modulation of the dynamic chromatin condensation of the marneral cluster.

***MARS* expression modulates an LHP1-dependent chromatin loop bringing together the *MRN1* locus and an ABA enhancer element**

It was recently observed that the spatial conformation of cluster-associated domains differs between transcriptionally active and silenced clusters. In Arabidopsis, segregating 3D contacts are distinguished among organs, in agreement with the corresponding transcriptional activity of clustered genes¹¹. Therefore, we explored whether *MARS* could participate in the dynamic regulation of the local 3D chromatin conformation modulating the transcription of the marneral cluster. According to available HiC datasets^{25,29} there is a significant interaction linking the intergenic region between *CYP71A16* and *MARS* and the *MRN1* locus (indicated as “Chromatin loop” in **Fig. 5a**). By using Chromatin Conformation Capture (3C), we determined that the formation of this chromatin loop drastically increased over 30 min with exogenous ABA and remained formed for at least 4 hours (**Fig 5b**). Thus, the formation of this chromatin loop positively correlates with the transcriptional accumulation of the marneral cluster genes in response to ABA (**Fig. 2b**).

The *MARS* locus is encompassed in the ABA-dependent chromatin loop (**Fig. 5a**). In order to determine the role of *MARS* in the modulation of local 3D chromatin conformation, we assessed the formation of the chromatin loop in RNAi-*MARS* lines. Notably, RNAi-*MARS* seedlings exhibit enhanced chromatin loop formation, which remained unchanged in response to exogenous ABA (**Fig. 5b**). Interestingly, LHP1 has been implicated in shaping local 3D conformation of target regions²⁵, suggesting that the LHP1-*MARS* module may dynamically switch the epigenetic status of the marneral cluster from a condensed-linear to a decondensed-3D structured chromatin conformation. Supporting this hypothesis, *lhp1* mutant seedlings exhibited enhanced chromatin loop formation compared to Col-0 (**Fig. 5c**). Overall, our results demonstrate that a chromatin loop within the marneral cluster is regulated by LHP1 and the interacting lncRNA *MARS*, encoded in the region encompassed by the loop.

To better understand the role of the *MARS*-dependent chromatin loop in response to ABA we looked for ABA-related *cis* regulatory sequences throughout the marneral cluster. We extracted from ChIP-seq the binding distribution of 13 ABA-related transcription factors (TFs)³⁰. Interestingly, a high enrichment in ABA TF binding sites was found at the *MARS* locus, as well as in the intergenic region between the *CYP71A16* and *MARS* loci, notably surrounding the contact point brought into close spatial proximity with the *MRN1* locus by the ABA-dependent 3D chromatin loop (**Fig. 5a**). We thus assessed the activating capacity of this region, potentially acting as a distant enhancer element of the *MRN1* proximal promoter. To this end, we made use of a *GUS*-based reporter system (as described in ³¹), fusing two regions of interest to a minimal 35S promoter. Two additional DNA regions nearby the putative enhancers were used as negative controls: one between *CYP705A12* and *CYP71A16* and the other at the 3' end of *AT5G42620* locus (**Fig. 5a** indicated in red). Among the two putative enhancers regions tested, one was able to activate *GUS* expression (Intergenic region 2, **Fig. 5a and 5d**), coinciding with the region showing a high enrichment of ABA-related TF binding sites close to the chromatin loop anchor point (**Fig. 5a**). Collectively, our results indicate that an ABA-driven chromatin loop brings into close spatial proximity the *MRN1* locus and a transcriptional activation site likely acting as an ABA enhancer element. Notably, this chromatin reorganization process depends on the LHP1-*MARS* module.

Long noncoding RNAs as emerging regulators of gene clusters

Physically linked genes organized in clusters are generally coregulated⁸. Considering that the lncRNA *MARS* is implicated in the regulation of the marneral cluster, we wondered whether the presence of noncoding transcriptional units may constitute a relevant feature of

gene cluster organization. Therefore, we collected two lists of Arabidopsis clustered genes, i.e. one of co-expressed neighboring genes¹⁰ and one of predicted metabolic gene clusters (by PlantSMASH³²). According to the latest Arabidopsis genome annotation (Araport11), among the 390 clusters of co-expressed neighboring genes, 189 (48%) contain at least one embedded lncRNA inside the cluster. Most importantly, among the 45 metabolic clusters, 28 (62%) include lncRNAs (**Fig. 6a**). Furthermore, among the clusters containing a lncRNA, a correlation analysis based on the maximum strength of co-expression between a lncRNA and any clustered gene revealed that the metabolic clusters exhibit a significantly higher correlation than co-expressed clusters (**Fig. 6b**). Altogether, our analyses suggest that lncRNA-mediated local epigenetic remodeling may constitute an emerging feature of non-homologous genes metabolic clusters in plants.

DISCUSSION

The cell nucleus is a dynamic arrangement of DNA, RNAs and proteins^{33,34}. Genome topology has emerged as an important feature in the complex network of mechanisms regulating gene activity and genome connectivity, leading to regionalized chromosomal spatial distribution and the clustering of diverse genomic regions with similar expression patterns³⁵.

In the last few years, noncoding transcription has been implicated in shaping 3D nuclear organization³⁶. Notably, RNase-A micro-injection into the nucleus revealed that long nuclear-retained RNAs maintained euchromatin in a biologically active decondensed state, whereas heterochromatin domains exhibited an RNA-independent structure^{37,38}. More recently, HiC analyses were performed in mammalian cells exposed or not to RNase, before and after crosslinking, or upon transcriptional inhibition³⁹. As a result, it was observed that topologically associated domains (TAD) boundaries remained mostly unaffected by RNase treatment, whereas compartmental interactions suffered a subtle disruption. In contrast, transcriptional inhibition led to weaker TAD boundaries, hinting at different roles of steady-state RNA vs. active transcription in nuclear organization³⁹.

In plants, several lncRNAs have been implicated in local chromatin conformation dynamics affecting the transcriptional activity of neighboring genes^{14,40}. Notably, the lncRNA *COLDWRAP* participates in the formation of an intragenic chromatin loop blocking the transcription of the flowering time regulator *FLOWERING LOCUS C* (*FLC*¹⁶) in response to cold, in a process involving the recruitment of PRC2 by direct interaction with the component CLF. The lncRNA *APOLO* also controls the transcriptional activity of its neighboring gene *PINOID* (*PID*) by dynamically modulating the formation of an intergenic chromatin loop

encompassing the divergent promoter of *PID* and *APOLO*¹⁵, in a process involving the PRC1 component LHP1. More recently, it was proposed that high levels of *APOLO* can decoy LHP1 away from multiple loci in *trans*, modulating the 3D conformation of target genes²⁸. In rice, the expression of the leucine-rich repeat receptor kinase clustered genes *RLKs* is modulated by the locally-encoded lncRNA *LRK ANTISENSE INTERGENIC RNA (LAIR)*. It was proposed that *LAIR* may directly recruit OsMOF (MALES ABSENT ON THE FIRST) and OsWDR5 (WD REPEAT DOMAIN 5), involved in H4K16 acetylation and chromatin remodeling⁴¹. Here we showed that the lncRNA *MARS* contributes to the co-regulation of a set of physically linked genes in *cis* in Arabidopsis. We demonstrated that the relative abundance of *in vitro*-transcribed *MARS* fine-tunes LHP1 binding to the cluster region in a stoichiometry-dependent manner, thus explaining how *MARS* affects H3K27me3 deposition and chromatin condensation. It has been shown in yeast that histone depletion boosts chromatin flexibility and facilitates chromatin loop formation on the kilobase pair scale⁴². In agreement thereof, we uncovered here the dynamic role of the LHP1-*MARS* module affecting nucleosome distribution across the marneral cluster in response to ABA, thus promoting the formation of an intra-cluster chromatin loop.

It has been recently observed that biosynthetic gene clusters are embedded in local three-dimensionally organized hot spots that segregate the region from the surrounding chromosome environment¹¹. Here we showed that active noncoding transcriptional units within the cluster may contribute to 3D conformation dynamics switching from silent to active states. Our results indicated that the *MARS*-dependent chromatin loop may bring the *MRN1* locus and a distal ABA-responsive element into close spatial proximity, likely acting as an enhancer. Notably, *MARS*-dependent LHP1 and H3K27me3 removal in Col-0, RNAi-*MARS* and the *lhp1* mutant correlated with chromatin decondensation, loop formation and increased marneral genes transcriptional activity in response to ABA. According to this model, chromatin loop conformation is related to LHP1 binding and is modulated by *MARS*. LHP1 recognition at basal *MARS* levels maintains a possibly linear conformation of the region, precluding the enhancer-*MRN1* locus interaction, whereas the positively activating chromatin loop is formed in the absence of LHP1. *MARS* transcriptional accumulation directly modulates LHP1 binding to the marneral cluster and high levels of *MARS* then titrate LHP1 away from the cluster (**Fig. 6c**; in response to ABA). Interestingly, when *MARS* levels are too low compared to basal levels (as in the RNAi lines), recruitment of LHP1 to the cluster is also impaired (**Fig. 6c**; *MARS* repression).

In mammals, growing evidence supports the role of lncRNAs in chromatin conformation determination⁴³ and enhancer activity (e.g. *PVT1*⁴⁴ and *CCAT1-L*⁴⁵). Here, we

showed that the nuclear-enriched lncRNA *MARS* brings together the *MRN1* proximal promoter and a putative enhancer element enriched in ABA-responsive TF binding sites. Interestingly, it has been shown that human lncRNAs can modulate the binding of TFs to their target chromatin (*DHFR*⁴⁶) and *PANDA*⁴⁷, whereas TFs have been implicated in chromatin loop formation in plants³⁵. Furthermore, it was shown that in addition to the TF NF-YA, the lncRNA *PANDA* interacts with the scaffold-attachment-factor A (SAFA) as well as with PRC1 and PRC2 to modulate cell senescence⁴⁸. Therefore, further research will be needed to determine what ABA-responsive TFs are in control of the marneral cluster and to elucidate how they participate in chromatin loop formation along the area, in relation with the PRC1-interacting lncRNA *MARS*.

Plants are a tremendous source of diverse chemicals which are important for their life and survival¹⁰. Marneral biosynthesis has been linked to root and leaf development, flowering time and embryogenesis². Here we found that the Arabidopsis marneral cluster is activated by the phytohormone ABA, in a lncRNA-dependent epigenetic reprogramming. *MARS* deregulation affects the cluster response to ABA, impacting seed germination. Interestingly, noncoding transcription had already been associated with seed germination. The *DELAY OF GERMINATION 1 (DOG1)* locus is a major actor regulating seed dormancy strength in Arabidopsis. Indeed, an antisense lncRNA (*asDOG1*) is able to repress *DOG1* transcription in mature plants. Notably, it was observed that *DOG1* suppression is released by shutting down antisense transcription, which is induced by ABA and drought⁴⁹.

It was proposed that the marneral cluster was founded by the duplication of ancestral genes, independent events of gene rearrangement and the recruitment of additional genes⁷. The exploration of the noncoding transcriptome in Arabidopsis recently served to identify ecotype-specific lncRNA-mediated responses to the environment⁵⁰. It was suggested that the noncoding genome may participate in multiple mechanisms involved in ecotype adaptation. Collectively, our results indicate that the acquisition of novel noncoding transcriptional units within biosynthetic gene clusters may constitute an additional regulatory layer behind their natural variation in plant evolution.

METHODS

Lead contact and materials availability

Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact, Martin Crespi (martin.crespi@ips2.universite-paris-saclay.fr).

Plant lines generated in this study are available from the Lead Contact with a completed Materials Transfer Agreement.

Lines selection and generation

All plants used in this study are in Columbia-0 background. RNAi-MARS were obtained using the pFRN binary vector⁵¹ bearing 250bp of the first exon of MARS gene (see primers in **Supplementary Table 1**), previously sub-cloned into the pENTR/D-TOPO vector. Arabidopsis plants were transformed using *Agrobacterium tumefaciens* Agl-0⁵². The T-DNA inserted line SALK_133089 was ordered to NASC (N633089).

Growth conditions

Seeds were sown in plates vertically disposed in a growing chamber in long day conditions (16 h in light 150uE; 8 h in dark; 21°C) for all the experiments. Plants were grown on solid half-strength MS medium (MS/2) supplemented with 0.7% sucrose, and without sucrose for the germination assay. For nitrate starvation assay, KNO₃ and Ca(NO₃)₂ were replaced from MS/2 by a corresponding amount of KCl and CaCl₂ respectively, 2.25 mM NH₄HCO₃ was added for nitrate-containing medium. For the phosphate starvation assay, growth medium contained 0.15 mM MgSO₄, 2.1 mM NH₄NO₃, 1.9 mM KNO₃, 0.34 mM CaCl₂, 0.5 μM KI, 10 μM FeCl₂, 10 μM H₃BO₃, 10 μM MnSO₄, 3 μM ZnSO₄, 0.1 μM CuSO₄, 0.1 μM CoCl₂, 0.1 μM Na₂MoO₄, 0.5 g.L⁻¹ sucrose supplemented with 500uM Pi for Pi containing medium versus 10uM for Pi free medium. All media were supplemented with 0.8g/L agar (Sigma-Aldrich, A1296 #BCBL6182V) and buffered at pH 5.6 with 3.4mM 2-(N-morpholino) ethane sulfonic acid. For the treatment with exogenous ABA or auxin, seedlings were sprayed with 10uM ABA and 10uM 1-Naphthaleneacetic acid (NAA), respectively. For heat stress, plates were transferred to a growth chamber at 37°C under the same lightning conditions. For nitrate and phosphate starvation assays, seedlings have been transferred at day 12 after sowing (DAS) from respectively nitrate and phosphate containing medium to nitrate and phosphate free medium. Finally, for seed germination assay, 0.5uM ABA was supplemented or not to the medium. Germination rate was evaluated twice a day. Seeds were considered germinated when the seed coats were perforated by elongating radicle. For all the experiments, samples were taken from 12 DAS starting two hours after light illumination, at different time-points, after cross-linking or not, depending on the experiment.

RT-qPCR

Total RNA was extracted from whole seedlings using TRI Reagent (Sigma-Aldrich) and treated with DNase (Fermentas) as indicated by the manufacturers. Reverse transcription was realized on 1µg total RNA using the Maxima Reverse Transcriptase (Thermo Scientific). qPCR was performed on a Light Cycler 480 with SYBR Green master I (Roche) in standard protocol (40 cycles, 60°C annealing). Primers used in this study are listed in **Supplementary Table 1**. Data were analyzed using the delta delta Ct method using *PROTEIN PHOSPHATASE 2A SUBUNIT A3 (AT1G13320)* for gene normalization⁵³ and time 0 for time-course experiment.

Chromatin Immunoprecipitation (ChIP)

ChIP was performed using anti-IgG (Millipore, Cat#12-370), anti-H3K27me3 (Millipore, Cat#07-449) and anti-LHP1 (Covalab, Pab0923-P), as previously described¹⁵, starting from two grams of seedlings crosslinked in 1% (v/v) formaldehyde. Chromatin was sonicated in a water bath Bioruptor Plus (Diagenode; 60 cycles of 30s ON and 30s OFF pulses at high intensity). ChIP was performed in an SX-8G IP-Star Compact Automated System (Diagenode). Antibody coated into Protein A Dynabeads (Invitrogen) were incubated 12 hours at 4 °C with the samples. Recovering of immunoprecipitated DNA was realized using Phenol:Chloroform:Isoamyl Alcohol (25:24:1, Sigma) followed by ethanol precipitation and analyzed by qPCR. For input samples, non-immunoprecipitated sonicated chromatin was processed in parallel.

In-vitro transcribed *MARS* RNA was obtained from a PCR product amplified from wild-type genomic DNA using the T7 promoter on the 5' PCR primer (**Supplementary Table 1**). PCR products were verified using agarose electrophoresis, and purified using NucleoSpin kit (Macherey-Nagel). 1µg of purified DNA was used for *in-vitro* transcription following the manufacturer instructions (HiScribe T7 High Yield RNA Synthesis Kit, NEB). Purified non-crosslinked chromatin obtained from five grams of *MARS* RNAi line 1 seedlings were resuspending in 1 mL of nuclei lysis buffer and split into five tubes. An increasing quantity of *MARS* RNA was added to each tube from 0 to 10 µg RNA and incubated under soft rotation during 3 h at 4 °C. Chromatin samples were then cross-linked using 1% (v/v) of formaldehyde for five minutes. Sonication and the following ChIP steps were performed as above.

Formaldehyde-Assisted Isolation of Regulatory Elements (FAIRE)

FAIRE was performed as described by ⁵⁴. After chromatin purification as for ChIP, only 50 µl from the 500 µl of purified chromatin were used (diluted to 500 µl with 10 mM Tris-HCl pH 8). For the qPCR, the same set of primers as for ChIP were used.

Nuclear purification

Non-cross-linked seedlings were used to assess the sub-cellular localization of RNAs. To obtain the nuclear fraction, chromatin was purified as for ChIP and resuspended, after the sucrose gradient, into 1mL of TRI Reagent (Sigma-Aldrich). For the total fraction, 200 µL of cell suspension in cell resuspension solution, were collected and completed with 800 µL of TRI Reagent to follow with the RNA extraction. RNA's samples were treated by DNase, and RT was performed using random hexamers prior to qPCR analysis.

RNA immunoprecipitation (RIP)

For RIP, the *lhp1* mutant complemented with the *ProLHP1:LHP1:GFP*⁵⁵ were used after 4 hours of treatment with ABA. After crosslinking and chromatin extraction as for ChIP, ten percent of resuspended chromatin was conserved at -20 °C as the input. Chromatin was sonicated in a water bath Bioruptor Plus (Diagenode; 5 cycles of 30 s ON and 30 s OFF pulses at high intensity). Anti-LHP1 RIP was performed using the anti-GFP antibody (Abcam ab290), as previously described¹⁵. Results were expressed as the percentage of cDNA detected after IP taking the input value as 100%.

Chromosome conformation capture (3C)

3C was performed as described by ⁵⁶. Briefly, chromatin was extracted from two grams of cross-linked seedlings as for ChIP. Overnight digestion at 37 °C were performed using 400U of Hind III enzyme (NEB). Digested DNA was ligated during 5 h incubation at 16 °C with 100 U of T4 DNA ligase (NEB). DNA was recovered after reverse crosslinking and Proteinase K treatment (Invitrogen) by Phenol:Chloroform:Isoamyl Acid (25:24:1; Sigma) extraction and ethanol precipitation. Interaction frequency was calculated by qPCR using a DNA region uncut by Hind III to normalize the amount of DNA.

Transcriptional activation assay in tobacco leaves

The *GUS* reporter system for validating the activity of the putative enhancer element was adapted from ³¹. Different DNA fragments were cloned in the GreenGate system⁵⁷ fused to a minimal 35S promoter element from CAMV (synthesized by Eurofins Genomics). The

sub-unit B3 from 35S promoter element from CAMV⁵⁸ was synthesized and used as a positive control. All primers used for cloning are indicated in **Supplementary Table 1**.

A. tumefaciens-mediated transient transformation was performed on 5-week-old tobacco plants using a needle-less syringe. Together with enhancer constructs, another vector containing mCherry driven by 35S promoter was co-transfected to control the transformation efficiency. Two leaf discs were collected near the infiltration site. One, to determine the transfection efficiency by mCherry fluorescence observation under epifluorescent microscope. The second was used for GUS staining, as previously described⁵⁹. Samples were incubated 4 h in the dark at 37 °C before observation.

Identification of lncRNA loci in Arabidopsis gene clusters

The genes of co-expressed clusters were retrieved from ¹⁰. The boundaries of the gene clusters were extracted using Araport11 annotations. The boundaries of the metabolic clusters were extracted from the plantSMASH predicted clusters on Arabidopsis³². Using Araport11 GFF, the lncRNAs (genes with a locus type annotated as “long_noncoding_rna”, “novel_transcribed_region” or “other_rna”) present within the boundaries of the cluster were retrieved.

Gene expression correlation analysis

To compute the correlation of expression in different organ of Arabidopsis we used the 113 RNA-seq datasets that were used for the Araport11 annotations ([10.1111/tpj.13415](https://tpj.13415)). These datasets were generated from untreated or mock-treated wild-type Col-0 plants. After removing the adaptors with Trim Galore with default parameters, the reads were mapped on TAIR10 with STAR v2.7.2a ([10.1093/bioinformatics/bts635](https://bioinformatics.bts635)) and the parameters ‘--alignIntronMin 20 --alignIntronMax 3000’. Gene expression was then quantified with featureCounts v2.0.0 ([10.1093/bioinformatics/btt656](https://bioinformatics.btt656)) with the parameters “-B -C -p -s 0” using the GFF of Araport11. Raw counts were then normalized by median of ratios using the DESeq2 R package⁶⁰.

For the correlation of expression inside the marneral cluster, the expression of the genes of the clusters and 25kb around it (four genes upstream and two downstream) were collected for the correlation analysis. Pearson's correlations for each pair of genes were computed after log 2 transformation of the normalized counts. The correlation value and associated p-value were plotted with the corrplot R package⁶¹.

Inside each co-expressed and metabolic clusters of genes, Pearson's correlation was computed between every possible pairs of lncRNA and coding gene as for the genes inside the marneral cluster. The maximum correlation value was kept as an indication of lncRNAs correlation with the genes of the cluster.

Quantification and statistical analyses

For all the experiments, at least two independent biological samples were considered. For RT-qPCR, each sample was prepared from a pool of 5 to 10 individual seedlings. For biochemistry assays (ChIP, FAIRE, nuclear purification, RIP and 3C) two to five grams of seedlings were prepared for each independent biological sample. For validation of enhancer function, the four leaf discs were taken from four independent tobacco plants. The tests used for statistical analyses are indicated in the respective figure legends. Statistical test and associated plots have been generated using R software (v3.6.3⁶²) with the help of the tidyverse package⁶³.

ACKNOWLEDGMENTS

IPS2 benefits from the support of Saclay Plant Sciences-SPS (ANR-17-EUR-0007). We thank Jeremie Bazin and Aurélie Christ from IPS2 for the helpful discussion about results interpretation and design of the experiments. We also thank Moussa Benhamed (IPS2) for helpful advice on epigenetic regulation. We thank Olivier Martin for critical reading of the manuscript.

AUTHORS' CONTRIBUTIONS

TR, FA, TB and MC conceived and designed the experiments. TR performed the experiments. TR and TB analyzed the data. All authors discussed the results and edited the manuscript.

COMPETING FINANCIAL INTERESTS

The authors declare no competing financial interests.

REFERENCES

1. Nützmann, H.-W., Scazzocchio, C. & Osbourn, A. Metabolic Gene Clusters in Eukaryotes. *Annu.*

- 515 *Rev. Genet.* **52**, 159–183 (2018).
- 516 2. Go, Y. S. *et al.* Identification of marneral synthase, which is critical for growth and
517 development in Arabidopsis. *Plant J.* **72**, 791–804 (2012).
- 518 3. Yasumoto, S., Fukushima, E. O., Seki, H. & Muranaka, T. Novel triterpene oxidizing activity of
519 Arabidopsis thaliana CYP716A subfamily enzymes. *FEBS Lett.* **590**, 533–540 (2016).
- 520 4. Field, B. & Osbourn, A. E. Clusters in Different Plants. *Science (80-.).* **194**, 543–547 (2008).
- 521 5. Boutanaev, A. M. *et al.* Investigation of terpene diversification across multiple sequenced
522 plant genomes. *Proc. Natl. Acad. Sci. U. S. A.* **112**, E81–E88 (2015).
- 523 6. Castillo, D. A., Kolesnikova, M. D. & Matsuda, S. P. T. An effective strategy for exploring
524 unknown metabolic pathways by genome mining. *J. Am. Chem. Soc.* **135**, 5885–5894 (2013).
- 525 7. Field, B. *et al.* Formation of plant metabolic gene clusters within dynamic chromosomal
526 regions. *Proc. Natl. Acad. Sci.* **108**, 16116–16121 (2011).
- 527 8. Nützmann, H. W., Huang, A. & Osbourn, A. Plant metabolic clusters – from genetics to
528 genomics. *New Phytol.* **211**, 771–789 (2016).
- 529 9. Nützmann, H. W. & Osbourn, A. Regulation of metabolic gene clusters in Arabidopsis thaliana.
530 *New Phytol.* **205**, 503–510 (2015).
- 531 10. Yu, N. *et al.* Delineation of metabolic gene clusters in plant genomes by chromatin signatures.
532 *Nucleic Acids Res.* **44**, 2255–2265 (2016).
- 533 11. Nützmann, H. *et al.* Active and repressed biosynthetic gene clusters have spatially distinct
534 chromosome states. 1–10 (2020) doi:10.1073/pnas.1920474117.
- 535 12. Rinn, J. L. & Chang, H. Y. Long Noncoding RNAs: Molecular Modalities to Organismal
536 Functions. *Annu. Rev. Biochem.* **89**, 283–308 (2020).
- 537 13. Berry, S., Rosa, S., Howard, M., Bühler, M. & Dean, C. Disruption of an RNA-binding hinge
538 region abolishes LHP1-mediated epigenetic repression. *Genes Dev.* **31**, 2115–2120 (2017).
- 539 14. Lucero, L., Fonouni-Farde, C., Crespi, M. & Ariel, F. Long noncoding RNAs shape transcription
540 in plants. *Transcription* **00**, 1–12 (2020).
- 541 15. Ariel, F. *et al.* Noncoding transcription by alternative rna polymerases dynamically regulates
542 an auxin-driven chromatin loop. *Mol. Cell* **55**, 383–396 (2014).
- 543 16. Kim, D.-H. & Sung, S. Vernalization-triggered intragenic chromatin-loop formation by long

544 noncoding RNAs. *Dev. Cell* **176**, 100–106 (2017).

545 17. Gagliardi, D. *et al.* Dynamic regulation of chromatin topology and transcription by inverted
546 repeat-derived small RNAs in sunflower. *Proc. Natl. Acad. Sci. U. S. A.* **116**, 17578–17583
547 (2019).

548 18. Cheng, C. Y. *et al.* Araport11: a complete reannotation of the Arabidopsis thaliana reference
549 genome. *Plant J.* **89**, 789–804 (2017).

550 19. Kong, L. *et al.* CPC: Assess the protein-coding potential of transcripts using sequence features
551 and support vector machine. *Nucleic Acids Res.* **35**, 345–349 (2007).

552 20. Kang, Y. J. *et al.* CPC2: A fast and accurate coding potential calculator based on sequence
553 intrinsic features. *Nucleic Acids Res.* **45**, W12–W16 (2017).

554 21. Heo, J. B. & Sung, S. Vernalization-mediated epigenetic silencing by a long intronic noncoding
555 RNA. *Science (80-.).* **331**, 76–79 (2011).

556 22. Bardou, F. *et al.* Long Noncoding RNA Modulates Alternative Splicing Regulators in
557 Arabidopsis. *Dev. Cell* **30**, 166–176 (2014).

558 23. Jarroux, J., Morillon, A. & Pinskaya, M. Long Non Coding RNA Biology. *Adv. Exp. Med. Biol.*
559 **1008**, 1–46 (2017).

560 24. Vishwakarma, K. *et al.* Absciscic acid signaling and abiotic stress tolerance in plants: A review
561 on current knowledge and future prospects. *Front. Plant Sci.* **8**, 1–12 (2017).

562 25. Veluchamy, A. *et al.* LHP1 Regulates H3K27me3 Spreading and Shapes the Three-Dimensional
563 Conformation of the Arabidopsis Genome. *PLoS One* **11**, 1–25 (2016).

564 26. Sijacic, P., Bajic, M., McKinney, E. C., Meagher, R. B. & Deal, R. B. Changes in chromatin
565 accessibility between Arabidopsis stem cells and mesophyll cells illuminate cell type-specific
566 transcription factor networks. *Plant J.* **94**, 215–231 (2018).

567 27. Yang, X., Tong, A., Yan, B. & Wang, X. Governing the silencing state of chromatin: The roles of
568 polycomb repressive complex 1 in arabidopsis. *Plant Cell Physiol.* **58**, 198–206 (2017).

569 28. Ariel, F. *et al.* R-Loop Mediated trans Action of the APOLO Long Noncoding RNA. *Mol. Cell* **77**,
570 1–11 (2020).

571 29. Liu, C. *et al.* Genome-wide analysis of chromatin packing in Arabidopsis thaliana at single-gene
572 resolution. *Genome Res.* **26**, 1057–1068 (2016).

- 573 30. Song, L. *et al.* A transcription factor hierarchy defines an environmental stress response
574 network. *Science* (80-.). **354**, 598 (2016).
- 575 31. Yan, W. *et al.* Dynamic control of enhancer activity drives stage-specific gene expression
576 during flower morphogenesis. *Nat. Commun.* **10**, 1–16 (2019).
- 577 32. Kautsar, S. A., Suarez Duran, H. G., Blin, K., Osbourn, A. & Medema, M. H. PlantISMASH:
578 Automated identification, annotation and expression analysis of plant biosynthetic gene
579 clusters. *Nucleic Acids Res.* **45**, W55–W63 (2017).
- 580 33. Cavalli, G. & Misteli, T. Functional implications of genome topology. *Nat. Struct. Mol. Biol.* **20**,
581 290–299 (2013).
- 582 34. Gibcus, J. H. & Dekker, J. The Hierarchy of the 3D Genome. *Mol. Cell* **49**, 773–782 (2013).
- 583 35. Rodriguez-Granados, N. Y. *et al.* Put your 3D glasses on: Plant chromatin is on show. *J. Exp.*
584 *Bot.* **67**, 3205–3221 (2016).
- 585 36. Quinodoz, S. & Guttman, M. Long non-coding RNAs: An emerging link between gene
586 regulation and nuclear organization. *Trends Cell Biol.* **24**, 651–663 (2014).
- 587 37. Caudron-Herger, M. *et al.* Coding RNAs with a non-coding function: Maintenance of open
588 chromatin structure. *Nucleus* **2**, (2011).
- 589 38. Caudron-Herger, M. & Rippe, K. Nuclear architecture by RNA. *Curr. Opin. Genet. Dev.* **22**, 179–
590 187 (2012).
- 591 39. Barutcu, A. R., Blencowe, B. J. & Rinn, J. L. Differential contribution of steady-state RNA and
592 active transcription in chromatin organization . *EMBO Rep.* **20**, 1–13 (2019).
- 593 40. Gagliardi, D. & Manavella, P. A. Short-range regulatory chromatin loops in plants. *New Phytol.*
594 1–6 (2020) doi:10.1111/nph.16632.
- 595 41. Wang, Y. *et al.* Overexpressing lncRNA LAIR increases grain yield and regulates neighbouring
596 gene cluster expression in rice. *Nat. Commun.* **9**, 1–9 (2018).
- 597 42. Diesinger, P. M., Kunkel, S., Langowski, J. & Heermann, D. W. Histone depletion facilitates
598 chromatin loops on the kilobasepair scale. *Biophys. J.* **99**, 2995–3001 (2010).
- 599 43. Gil, N. & Ulitsky, I. Regulation of gene expression by cis-acting long non-coding RNAs. *Nat.*
600 *Rev. Genet.* **21**, 102–117 (2020).
- 601 44. Cho, S. W. *et al.* Promoter of lncRNA Gene PVT1 Is a Tumor-Suppressor DNA Boundary

602 Element. *Cell* **173**, 1398–1412.e22 (2018).

603 45. Xiang, J. F. *et al.* Human colorectal cancer-specific CCAT1-L lncRNA regulates long-range
604 chromatin interactions at the MYC locus. *Cell Res.* **24**, 513–531 (2014).

605 46. Martianov, I., Ramadass, A., Serra Barros, A., Chow, N. & Akoulitchiev, A. Repression of the
606 human dihydrofolate reductase gene by a non-coding interfering transcript. *Nature* **445**, 666–
607 670 (2007).

608 47. Hung, T. *et al.* Extensive and coordinated transcription of noncoding RNAs within cell-cycle
609 promoters. *Nat. Genet.* **43**, 621–629 (2011).

610 48. Puvvula, P. K. *et al.* Long noncoding RNA PANDA and scaffold-attachment-factor SAFA control
611 senescence entry and exit. *Nat. Commun.* **5**, (2014).

612 49. Yatusovich, R. *et al.* Antisense transcription represses Arabidopsis seed dormancy QTL DOG 1
613 to regulate drought tolerance. *EMBO Rep.* **18**, 2186–2196 (2017).

614 50. Blein, T. *et al.* Landscape of the non-coding transcriptome response of two Arabidopsis
615 ecotypes to phosphate starvation. *Plant Physiol.* **183**, pp.00446.2020 (2020).

616 51. Ariel, F. *et al.* Two direct targets of cytokinin signaling regulate symbiotic nodulation in
617 medicago truncatula. *Plant Cell* **24**, 3838–3852 (2012).

618 52. Clough, S. J. & Bent, A. F. Floral dip: A simplified method for Agrobacterium-mediated
619 transformation of Arabidopsis thaliana. *Plant J.* **16**, 735–743 (1998).

620 53. Czechowski, T., Stitt, M., Altmann, T., Udvardi, M. K. & Scheible, W. Genome-Wide
621 Identification and Testing of Superior Reference Genes for Transcript Normalization in
622 Arabidopsis. *Plant Physiol.* **139**, 5–17 (2005).

623 54. Simon, J. M., Giresi, P. G., Davis, I. J. & Lieb, J. D. Using formaldehyde-assisted isolation of
624 regulatory elements (FAIRE) to isolate active regulatory DNA. *Nat. Protoc.* **7**, 256–267 (2012).

625 55. Nakahigashi, K., Jasencakova, Z., Schubert, I. & Goto, K. The Arabidopsis HETEROCHROMATIN
626 PROTEIN1 homolog (TERMINAL FLOWER2) silences genes within the euchromatic region but
627 not genes positioned in heterochromatin. *Plant Cell Physiol.* **46**, 1747–1756 (2005).

628 56. Louwers, M., Splinter, E., van Driel, R., de Laat, W. & Stam, M. Studying physical chromatin
629 interactions in plants using Chromosome Conformation Capture (3C). *Nat. Protoc.* **4**, 1216–
630 1229 (2009).

- 631 57. Lampropoulos, A. *et al.* GreenGate - A novel, versatile, and efficient cloning system for plant
632 transgenesis. *PLoS One* **8**, (2013).
- 633 58. Moreno-risueno, M. A. *et al.* NIH Public Access. **329**, 1306–1311 (2010).
- 634 59. Jefferson, R. A., Kavanagh, T. A. & Bevan, M. W. GUS fusions: ,B-glucuronidase as a sensitive
635 and versatile gene fusion marker in higher plants. *EMBO J.* **6**, 3901–3907 (1987).
- 636 60. Love, M. I., Huber, W. & Anders, S. Moderated estimation of fold change and dispersion for
637 RNA-seq data with DESeq2. *Genome Biol.* **15**, 1–21 (2014).
- 638 61. Taiyun Wei and Viliam Simko (2017). R package "corrplot": Visualization of a Correlation
639 Matrix (Version 0.84). Available online at <https://github.com/taiyun/corrplot>
- 640 62. R Core Team (2018). R: A language and environment for statistical computing. R Foundation
641 for Statistical Computing, Vienna, Austria. Available online at <https://www.R-project.org/>.
- 642 63. Wickham, H. *et al.* Welcome to the Tidyverse. *J. Open Source Softw.* **4**, 1686 (2019).

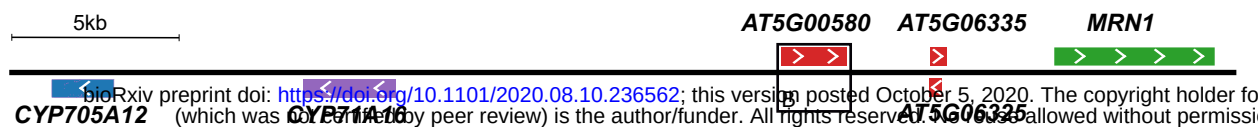
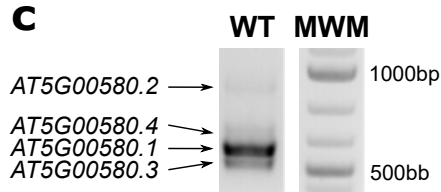
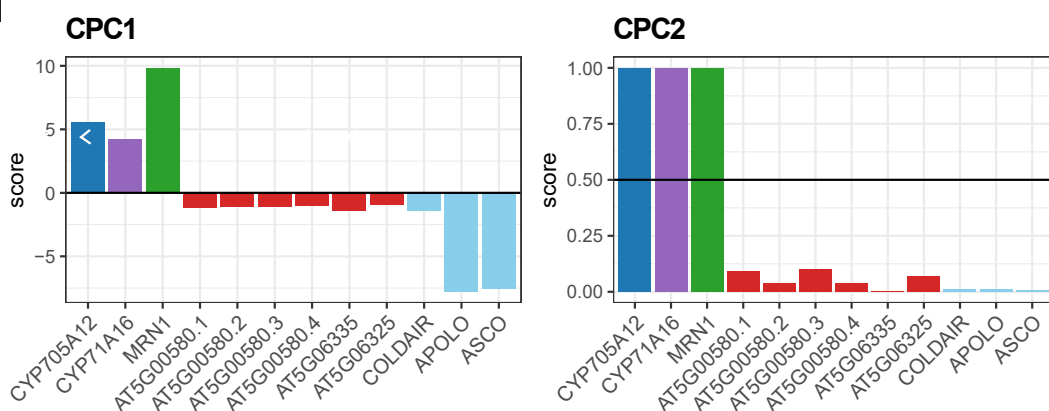
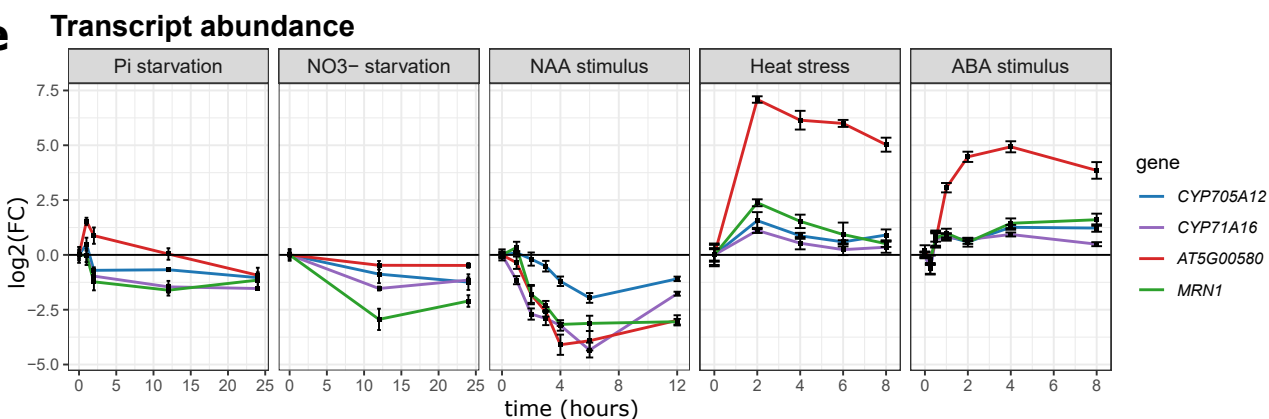
a**b****c****d****e**

Fig. 1, *AT5G00580* is a lncRNA transcribed from the marneral cluster locus and its expression correlates with its neighboring genes

a, Schematic illustration of the marneral cluster. Genes are indicated with plain rectangles and white arrows indicate the sense of transcription. The square indicates the region displayed in (B).

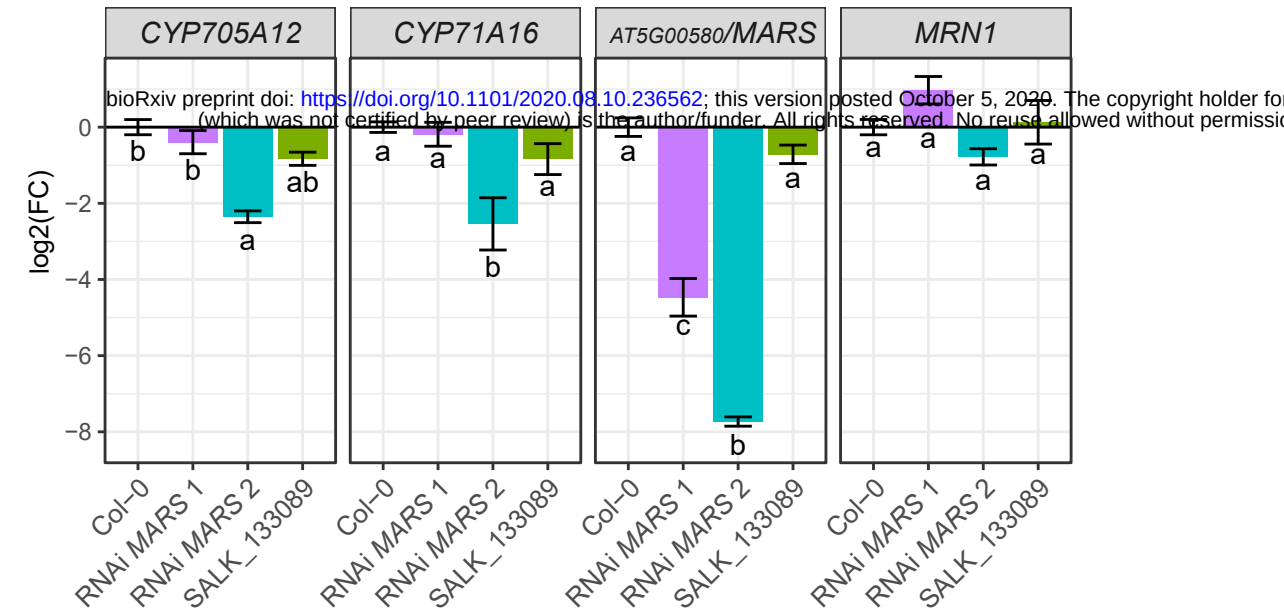
b, Schematic illustration of the different isoforms of *AT5G00580* transcripts. First line corresponds to *AT5G00580* genomic region whereas the other lines present the various isoforms. For each isoform, exons are indicated with rectangles and introns with solid lines. Black arrows indicate the primers used for the cDNA amplification in (C).

c, cDNA amplification of *AT5G00580* isoforms. The position of the primers used for the amplification are displayed on B. MWM stands for Molecular Weight Marker (GeneRuler 1 kb Plus DNA Ladder, Thermo Scientific).

d, Coding potential of the transcripts located in the marneral cluster genomic region. Scores were calculated using CPC1 (left) and CPC2 (right) programs^{19,20}. For each algorithm, the threshold between coding and non-coding genes is displayed with a horizontal solid black line. Coding genes are situated above the threshold, whereas non coding genes are situated below. *COLDAIR*, *APOLO* and *ASCO* are used as positive controls of non-coding transcripts.

e, Dynamic transcriptional levels of co-regulated genes of the marneral cluster in response to phosphate and nitrate starvation, heat stress, and exogenous ABA and auxin. Gene expression data are represented as the mean ± standard error (n ≥ 3) of the log2 fold change compared to time 0h.

a Transcript abundance



b Transcript abundance

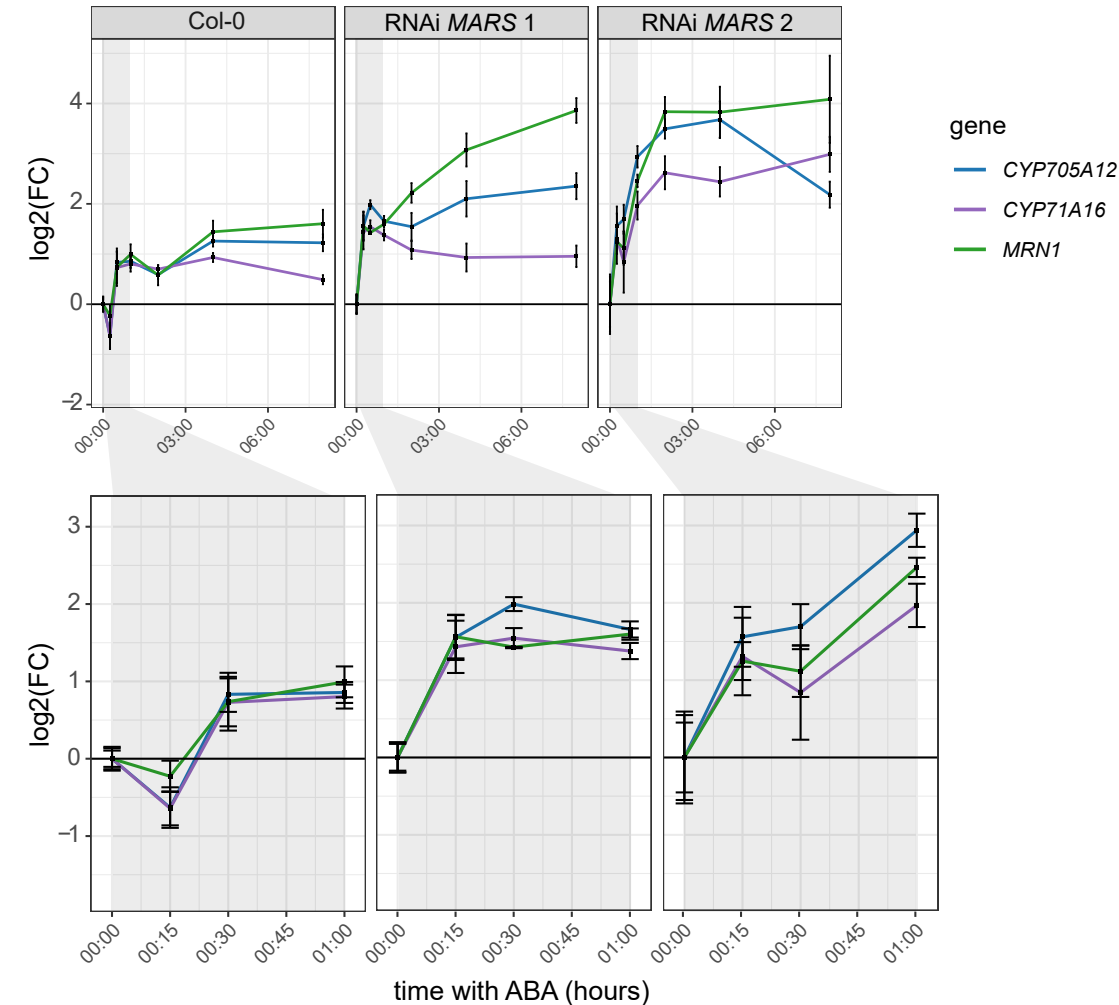
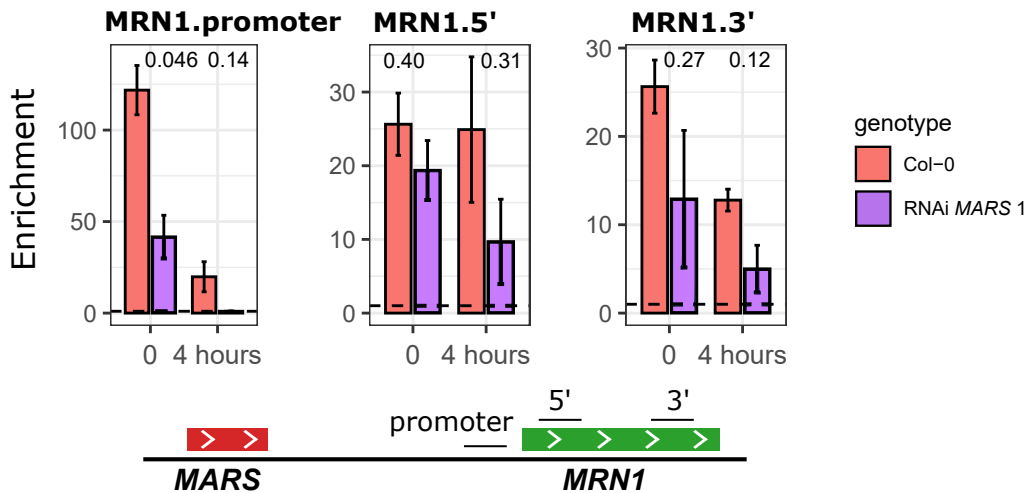


Fig. 2, *MARS* transcriptional activity modulates the response to ABA of the marneral cluster

a, Transcript levels of the marneral cluster genes in control conditions in RNAi lines targeting *AT5G00580/MARS* and *SALK_133089* line. Transcriptional levels are represented as the mean $\pm$ standard error (n = 3) of the log₂ fold change compared to Col-0. Letters indicate statistic group determined by one-way analysis of variance (ANOVA) followed by Tukey's post-hoc test. For each gene, each letter indicates statistical difference between genotypes (p $\leq$ 0.05).

b, Transcript levels of the genes of the marneral cluster in response to ABA treatment in RNAi lines targeting *AT5G00580/MARS*. Gene expression data are represented as the mean $\pm$ standard error (n = 3) of the log₂ fold change compared to time 0h.

a H3K27me3



b LHP1

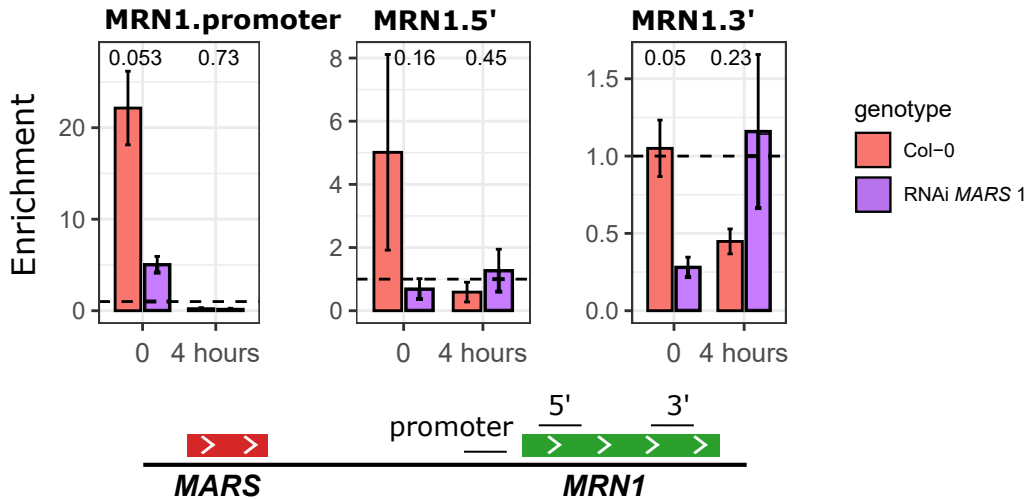
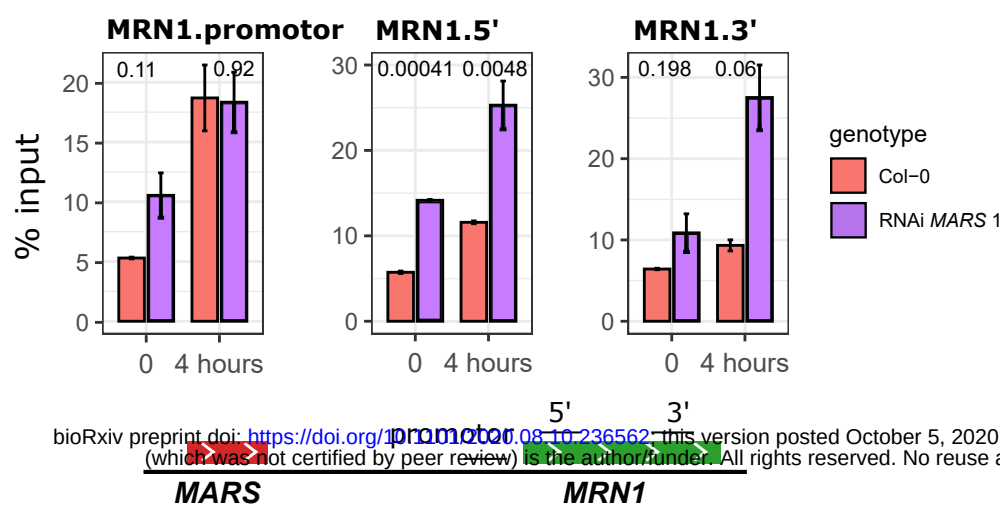


Fig. 3, *MARS* modulates the epigenetic landscape of *MRN1* locus

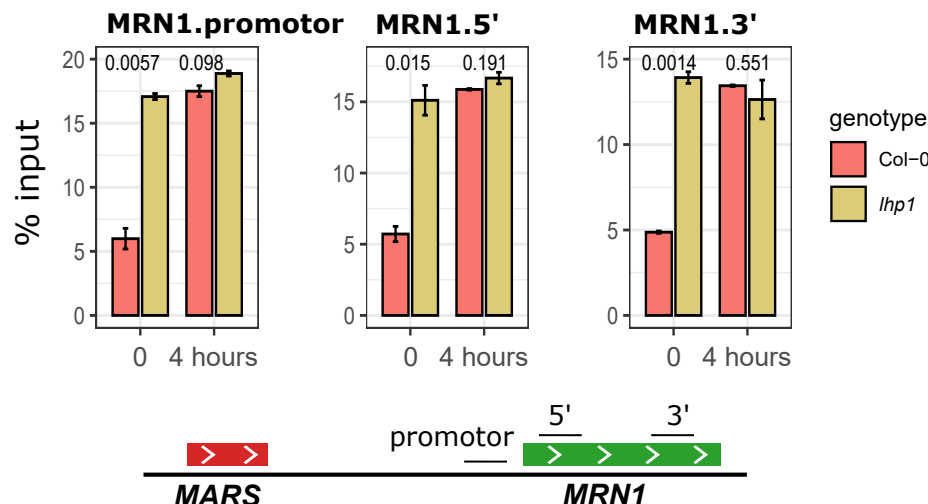
a, H3K27me3 deposition over the *MRN1* gene in Col-0 and RNAi-*MARS* seedlings under control conditions and in response to ABA. Higher values of ChIP-qPCR indicate more H3K27me3.

b, LHP1 binding to the *MRN1* gene in Col-0 and RNAi-*MARS* seedlings in the same conditions as in (A). Higher values of ChIP-qPCR indicate more LHP1 deposition. In (A) and (B), values under the dotted line are considered as not enriched. Results are represented as the mean $\pm$ standard error ($n = 2$) of the H3K27me3/IgG or LHP1/IgG ratio. Numbers are p-value of the difference between the two genotypes determined by Student t-test.

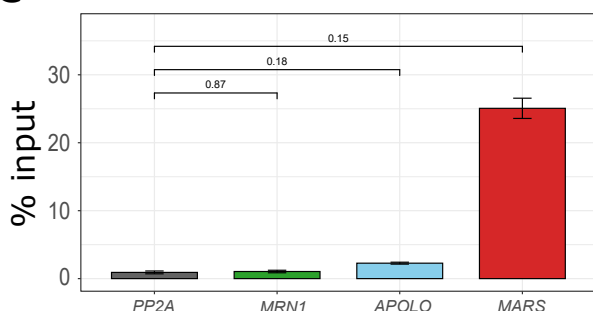
a FAIRE



b FAIRE



c LHP1 RIP



d LHP1 ChIP, *MRN1*.promoter

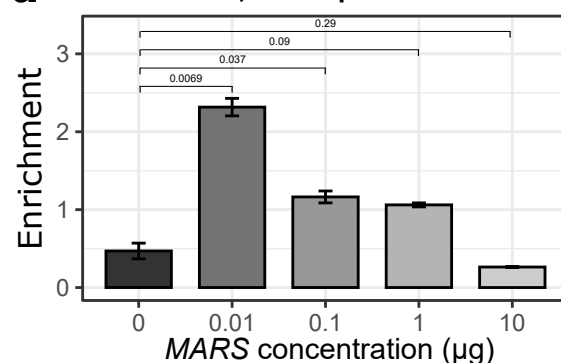


Fig. 4, *MARS* influences chromatin condensation of *MRN1* gene through its interaction with LHP1 protein

a, Chromatin condensation in *MRN1* gene of Col-0 and RNAi-*MARS* seedlings in control conditions and in response to ABA, determined by Formaldehyde Assisted Isolation of Regulatory Element (FAIRE)-qPCR.

b, Evolution of the chromatin condensation in *MRN1* gene of Col-0 and *lhp1* mutant subjected to ABA treatment determined by Formaldehyde Assisted Isolation of Regulatory Element (FAIRE) qPCR.

In **a** and **b**, results are expressed as the mean $\pm$ standard error ($n = 3$) of the percentage of input (signal measured before isolation of decondensed region of chromatin, free of nucleosomes). Lower value indicates more condensed chromatin. Numbers are p-value of the difference between the two genotypes determined by Student t-test.

c, LHP1-*MARS* interaction was assessed by RNA immunoprecipitation (RIP) using *LHP1-GFP* seedlings. Negative controls include a housekeeping gene (*PP2A*) and *MRN1* mRNA. The interaction between *APOLO* and LHP1 was taken as a positive control¹⁵. Results are expressed as the mean $\pm$ standard error ($n = 4$) of the percentage of input (signal measured before immunoprecipitation).

d, LHP1 binding to the *MRN1* promoter region in chromatin from RNAi-*MARS* seedlings upon increasing amounts of *in-vitro* transcribed *MARS* RNA. After incubation (see Methods), the samples were crosslinked for LHP1 ChIP-qPCR. Higher values indicate more LHP1-DNA interaction. Results are expressed as the mean $\pm$ standard error ($n = 2$) of the LHP1/IgG ratio.

In **c** and **d** numbers are p-value of the difference between the different corresponding genes determined by Student t-test.

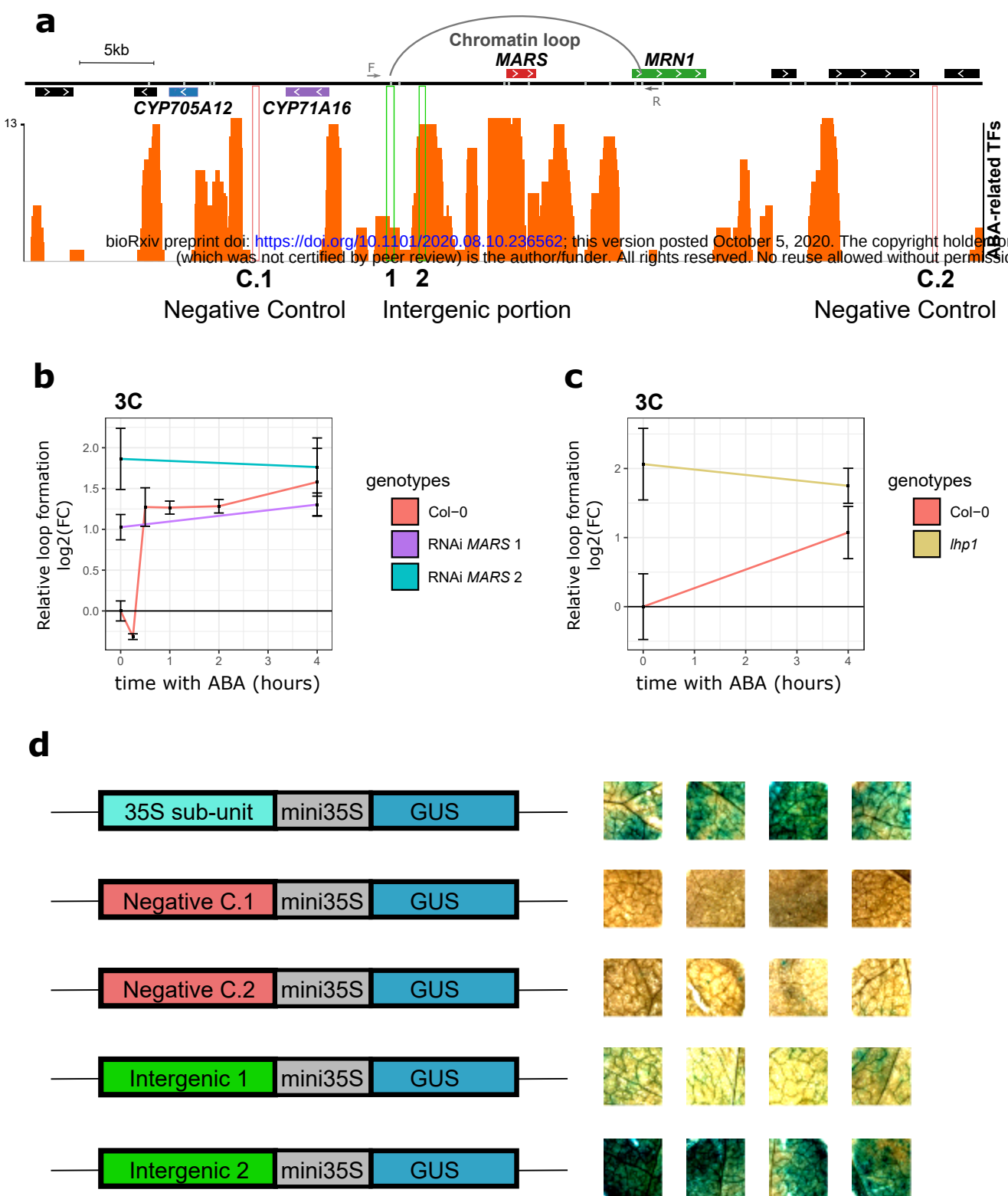


Fig. 5, An LHP1-dependent chromatin loop brings together the *MRN1* locus and a putative enhancer element in response to ABA

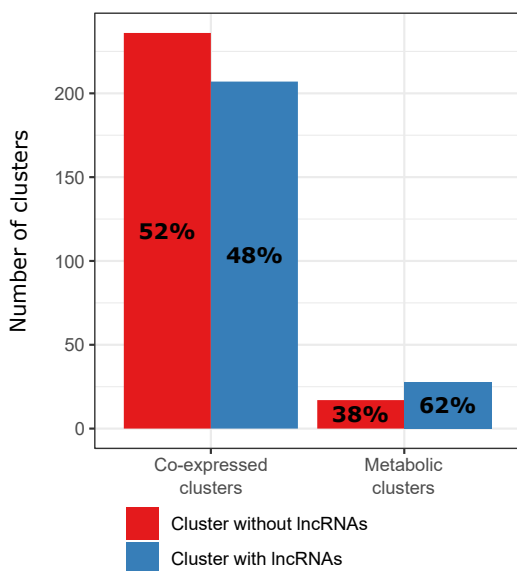
a, Schematic illustration of the loop linking the *MRN1* locus with the intergenic region between *CYP71A16* and *MARS*. Forward (F) and Reverse (R) oligonucleotides used for 3C-qPCR (in B–C) are indicated with arrows. The orange track shows the number of different ABA-related transcription factor binding sites (HB6, HB7, GBF2, GBF3, MYB3, MYB44, NF-YC2, NF-YB2, ANAC102, ANAC032, ABF1, ABF3, ABF4, RD26, ZAT6, FBH3, DREBA2A, AT5G04760, HAT22 and HSFA6A) found along the marneral cluster³⁰. Green and red rectangles indicate the putative enhancer region and the negative controls, respectively, tested for the GUS-based reporter system in **d**.

b, Relative chromatin loop formation in response to ABA in Col-0 and RNAi-*MARS* seedlings. Results are expressed as the mean $\pm$ standard error ($n = 2$) from 3C-qPCR using primer F and R shown on **a**, compared to time 0h.

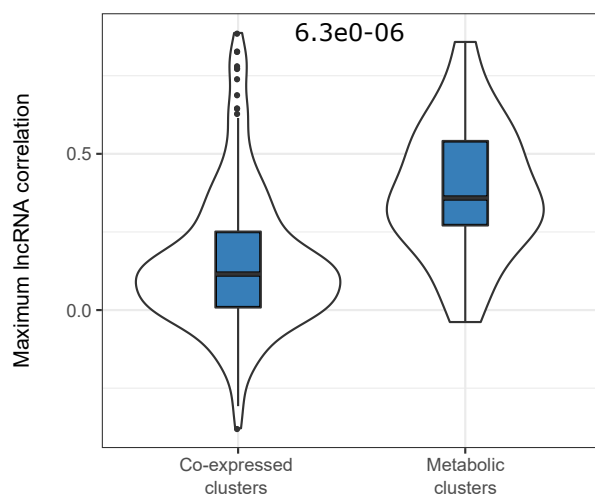
c, Relative chromatin loop formation in response to ABA treatment in Col-0 and *lhp1* mutant. Data are represented as the mean $\pm$ standard error ($n = 3$) from 3C-qPCR using primer F and R shown on **a**, compared to time 0h.

d, Constructs used for GUS-based reporter system are illustrated on the left. Corresponding transformed tobacco leaf discs are on the right ($n = 4$). First line represents the positive control in which 35S sub-unit controls *GUS* expression. The second and third lines show two independent negative controls in which *GUS* gene is driven by a genomic region that does not contain ABA-related binding sites as indicated in **a**. In the remaining lines, the transcriptional activity is assessed for the two intergenic regions indicated in **a**.

a



b



c

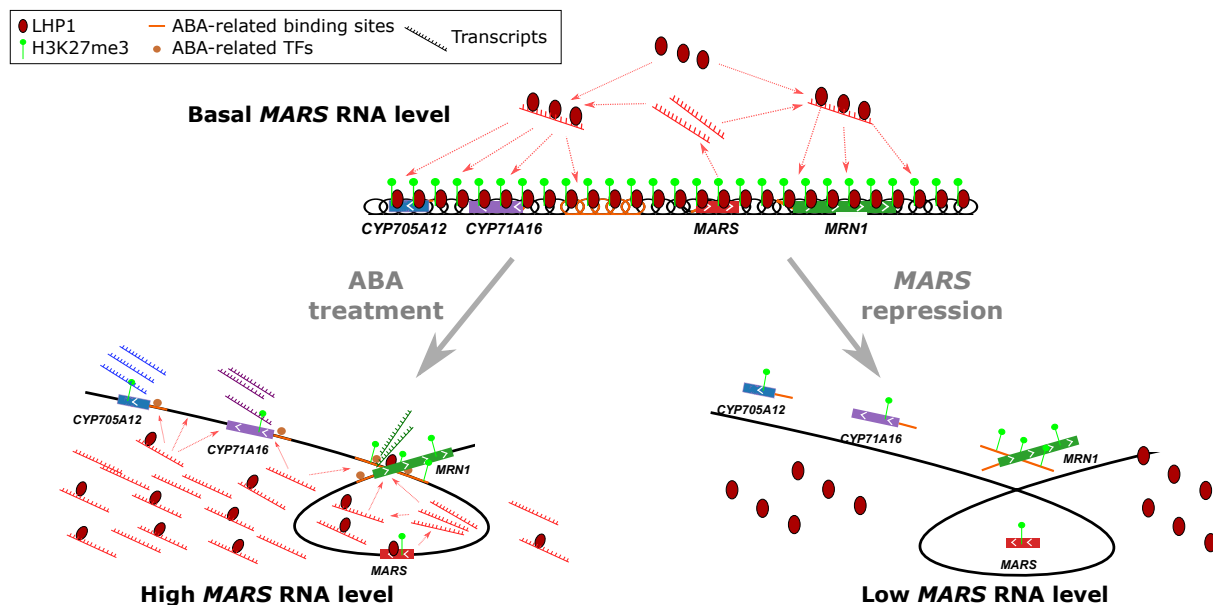


Fig. 6, Regulation of metabolic clusters in plants by IncRNA

a, The proportion of metabolic clusters including IncRNA loci is higher than for other clusters. The co-expressed clusters were predicted in ¹⁰ and correspond to co-expressed neighboring genes. The metabolic clusters are co-expressed neighboring genes involved in the biosynthesis of a particular secondary metabolite predicted by plantSMASH³².

b, Maximum level of correlation between a IncRNA and any clustered gene calculated in each cluster. The number shown above indicates the p-value of the difference between the two type of clusters determined by Student t-test.

c, The IncRNA *MARS* regulates the expression of the marneral cluster genes through epigenetic reprogramming and chromatin conformation. In control conditions (upper panel) the chromatin of the marneral cluster is enriched in H3K27me3 and LHP1, which results in a condensed and possibly linear chromatin conformation. In response to ABA (bottom left panel) *MARS* over-accumulated transcripts titrate LHP1 away from the cluster. The decrease of LHP1 deposition diminishes H3K27me3 distribution, relaxes the chromatin and as a consequence allows the formation of a chromatin loop that brings together the enhancer element and *MRN1* proximal promoter, leading to transcriptional activation. When *MARS* is repressed, LHP1 recruitment to the cluster is impaired, thus leading to a similar chromatin state: decrease in H3K27me3 mark, chromatin decondensation and increase in chromatin loop conformation. Under this chromatin state, the clustered genes become highly responsive to the ABA treatment.