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Research Report

Classification: Biological Sciences / Plant Biology

Title: Active and repressed biosynthetic gene clusters have spatially distinct chromosome states

Short title. Chromosome structure at biosynthetic gene clusters

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Author Contributions

H.W.N., A.O. designed research; H.W.N, D.D., A.R.-C., J.E.S., M.D.S., S.L.V.-F., E.W. performed research; P.F. contributed new reagents/analytic tools; H.W.N., D.D., A.R.-C., J.E.S., M.D.S., S.W.W., S.L.V.-F. analyzed data; and L.H., S.L.V.-F., A.O. and H.W.N. wrote the paper.

Abstract.

While colocalization within a bacterial operon enables co-expression of the constituent genes, the mechanistic logic of clustering of non-homologous monocistronic genes in eukaryotes is not immediately obvious. Biosynthetic gene clusters that encode pathways for specialised metabolites are an exception to the classical eukaryote rule of random gene location and provide paradigmatic exemplars with which to understand eukaryotic cluster dynamics and regulation. Here, using 3C, Hi-C and Capture Hi-C organ-specific chromosome conformation capture techniques along with high-resolution microscopy, we investigate how chromosome topology relates to transcriptional activity of clustered biosynthetic pathway genes in Arabidopsis thaliana. Our analyses reveal that biosynthetic gene clusters are embedded in local hot-spots of three-dimensional contacts that segregate cluster regions from the surrounding chromosome environment. The spatial conformation of these cluster-associated domains differs between transcriptionally active and silenced clusters. We further show that silenced clusters associate with heterochromatic chromosomal domains towards the periphery of the nucleus, while transcriptionally active clusters re-locate away from the nuclear periphery. Examination of chromosome structure at unrelated clusters in maize, rice and tomato indicates that integration of clustered pathway genes into distinct topological domains is a common feature in plant genomes. Our results shed light on the potential mechanisms that constrain co-expression within clusters of non-homologous eukaryotic genes and suggest that gene clustering in the one-dimensional chromosome is accompanied by compartmentalisation of the three-dimensional chromosome.

Significance statement

Clusters of co-expressed and co-localised biosynthetic pathway genes in plants are a paradigmatic example of the non-random organisation of the eukaryotic genome and present an ideal opportunity to understand the logic of eukaryote gene cluster regulation. Here, we carry out an in-depth analysis of the chromosomal topology of biosynthetic gene clusters and their positioning in nuclear space. We demonstrate that plant biosynthetic gene clusters reside in highly interactive domains that undergo marked changes in local conformation and nuclear positioning in cluster expressing and non-expressing organs. As such, metabolic gene clusters rank amongst the most dynamic regions in the genome of the model species *A. thaliana*. Our results shed light on the potential mechanisms that constrain co-expression within clusters of genes.

Main Text

Introduction. Gene order is a central feature that distinguishes eukaryotic genomes from their prokaryotic counterparts. Prokaryotic genomes are characterised by co-localisation of functionally related genes in operons (1). In contrast, functionally related genes in eukaryotes are commonly dispersed throughout genomes. Progress in genomics and transcriptomics have shown that gene order in eukaryotes is far from random, and that the positioning of genes affects their transcriptional activity and evolutionary retention (2-4). In addition, diverse examples of co-localised and functionally related genes ('operon-like' gene clusters) have been identified in eukaryotes that are reminiscent of gene organisation in prokaryotes (5-11). While the polycistronic transcription of bacterial operons provides an immediate and established mechanistic logic for co-expression of functionally related genes, for eukaryotic operon-like clusters of genes that are predominantly transcribed as single monocistronic units with individual promoters this mechanistic logic is not obvious (1, 3, 12).

In plants, it has recently been discovered that genomes contain regions characterised by operon-like clusters of co-localised and non-sequence-related genes involved in the biosynthesis of natural products. These clusters encode pathways for the biosynthesis of diverse molecules, ranging from medicinal alkaloids to polyketide components of wax layers and triterpenes that shape the root microbiota (13-16). The identification of this clustering phenomenon has led to the development of new genomics-driven strategies for pathway discovery (17-20). Plant biosynthetic gene clusters range in size from ~35 kb to several hundred kb. They are located in genomic regions that are prone to chromosomal rearrangement, and have arisen by recruitment of genes from elsewhere in the genome followed by neofunctionalization (14, 21-23). Individual metabolic gene clusters and their variants are usually confined to narrow taxonomic windows (14).

The genes within these biosynthetic clusters are typically co-expressed in specific plant organs and/or in response to certain environmental triggers. High transcriptional activity in metabolite producing cells is often contrasted by tight transcriptional silencing in non-producing cells (14, 19, 24, 25).

It is hypothesised that physical linkage of functionally-related genes in eukaryotes is associated with specialised regulatory processes (3, 12). Physical linkage may facilitate coordinate gene regulation through shared promoter and regulatory DNA elements as well as common epigenetic modifications of cluster-associated histones and DNA motifs (19, 20, 2631). Furthermore, it has been proposed that three-dimensional (3D) chromosome structure and localisation to specific nuclear territories are mechanisms for co-ordinate transcriptional regulation of adjacent genes. Seminal work on clusters of homologous genes in humans and animals, such as the HOX and β -globin clusters, support this hypothesis (32-34).

In plants and for eukaryotic clusters of non-homologous genes in general, however, it remains unknown how groups of neighbouring and co-expressed genes integrate into the nuclear three-dimensional environment. A recent study in the filamentous fungus *Epichloë festucae* reported the localisation of a biosynthetic gene cluster in a single topologically associated domain (TAD) and the authors suggested that activation of this cluster may be associated with a remodelling of this chromosome structure (35).

In previous studies we have shown that common signatures of chromatin marks delineate plant biosynthetic gene clusters, and also that *Arabidopsis thaliana* chromatin mutants have altered cluster transcript levels compared to the wild type (20). Furthermore in diploid oat (*Avena strigosa*), we have shown by high resolution DNA *in situ* hybridisation that expression of a biosynthetic gene cluster for the synthesis of antimicrobial defense compounds known as avenacins is associated with chromatin decondensation (25).

Here, we characterise the chromosome topology of metabolic gene clusters in plants, using previously characterised gene clusters in *A. thaliana* as our models. We have recently shown that these clustered biosynthetic pathways form a metabolic network that shapes the root microbiota (13). These gene clusters are co-ordinately transcribed in the roots but silenced in the aerial organs of the plant (20, 23, 24, 36). They therefore offer an ideal experimental system for investigating the organ-specific regulation of plant biosynthetic gene clusters.

To define the chromosome architecture of metabolic gene clusters and their integration into the nuclear environment we carried out chromosome conformation capture (3C, Hi-C and Capture Hi-C) experiments using nuclear preparations from roots and leaves of *A. thaliana* seedlings. We show that the *A. thaliana* biosynthetic gene clusters are embedded in local interactive three-dimensional chromosomal domains that adopt different structures in expressing and non-expressing organs. Comparative analysis reveals that these domains undergo some of the most drastic genome-wide changes to chromosome topology when roots and leaves are compared. We further demonstrate that these biosynthetic clusters are localised to heterochromatic areas of the genome when silenced. Incorporation and analysis of available Hi-C maps implicates histone H3 lysine 27 trimethylation (H3K27me3) as a central feature of the 3D domains at silenced clusters. Examination of chromosome structure at unrelated clusters in maize, rice and tomato indicates that integration of clustered pathway genes into distinct topologically associated domains is a widespread feature in plants.

Collectively, our work provides a high-resolution view of the nuclear organisation of biosynthetic gene clusters in plants. It demonstrates that a unique pattern of chromosomal conformations is established at clusters. Our findings also open up a novel potential route to manipulate plant specialized metabolism by interfering with higher-order regulatory mechanisms

Results and discussion.

Analysis of 3D chromosome conformation reveals organ-specific differences

To define the three-dimensional chromosome architecture at biosynthetic gene clusters in *A*. *thaliana* we set out to establish chromosome conformation capture protocols in conditions that would reflect transcriptional 'on' and 'off' states of the clusters. In earlier work we and others have shown that expression of several previously characterized biosynthetic gene clusters, amongst them the thalianol cluster, is tightly repressed in the leaves and highly expressed in the roots of young *A*. *thaliana* seedlings (20, 23, 24, 36, 37). To corroborate this, we performed whole transcriptome analysis of RNA extracted from roots and leaves of seven-day old seedlings (SI Dataset 1). We observed marked changes in the transcript levels for three distinct clusters – the thalianol, marneral and arabidiol/baruol clusters – when root and leaves were compared (Fig. 1A, B).

We then performed genome-wide Hi-C analysis for DNA from both organs. We obtained 71 – 108 million valid unique paired-end reads from each library (Tables S1, S2). We corrected and normalised the derived interaction counts for experimental biases and genomic distance. The normalized counts serve as measure of interaction strength between any two chromosomal sites and were plotted as two-dimensional Hi-C maps. Visual examination showed strong interchromosomal contacts between pericentromeric blocks and between all telomers in both roots and leaves (Fig 1C). We could also readily detect a number of prominent off-diagonal punctate signals that reflect interactions of defined chromosomal loci

known as interactive heterochromatic islands (IHIs) (Figs 1C, S1) (38, 39). These features are overall consistent with those observed in the previously reported overarching conformation of *A. thaliana* chromosomes in whole seedlings (38-40).

In in-depth comparative examinations of the chromosome architecture, we observed several differences between the chromosome features observed for roots and leaves. In leaves, significantly increased interaction counts were enriched in the pericentromeric regions while in roots enhanced interactions were predominantly localised to the chromosome arms and telomeres (Fig. 1C, D, S1, SI Dataset 2). We further detected changes to the IHIs. The intensity of individual interchromosomal IHI interactions varied significantly and one previously undescribed IHI located on chromosome 1 was identified (Figs. 1D, S2,3). These findings highlight organ-specific reconfigurations of the *A. thaliana* 3D chromosome architecture. Differences in interaction intensity between different tissues have also been reported for Hi-C analyses in rice and maize (41).

In *A. thaliana*, alterations to interaction intensities of pericentromeric and telomeric regions of the genomes and IHIs have been described for mutant lines that are defective for different epigenetic pathways such as DNA methylation and histone H3 lysine 9 methylation (38). This may suggest that differences in the epigenetic environment between roots and leaves underlie the observed 3D changes. Furthermore, differences in nuclear shape and ploidy levels of individual nuclei of roots and leaves may be associated with the variations in interaction intensities (42).

Silenced metabolic gene clusters associate with heterochromatic areas within the nucleus

We then focused our analysis on the thalianol metabolic gene cluster. We have recently shown that the metabolic products derived from the thalianol cluster have important roles in shaping the root microbiome of *A. thaliana* (13). We have further established that the cluster is delineated by repressive histone H3 lysine 27 trimethylation marks (H3K27me3) and the histone variant H2A.Z involved in positive regulation (20, 37). The thalianol cluster consists of four core genes that cover approximately 33 kb and a peripheral gene that is separated from the core cluster by 10 kb (the latter 10 kb region including two unrelated intervening genes) (Fig 1A) (13, 24). All cluster genes are widely expressed in root tissues and repressed in aerial plant tissues (Fig 1B, S4).

Visual inspection of our Hi-C maps for roots and leaves show small local interactive domains encompassing the thalianol cluster and separating it from the neighbouring genomic environment (Fig 2A, S5). Re-analysis of previous whole seedling-derived conformation data shows a similar domain at the thalianol cluster (Fig S6) (31). Strikingly, the location and interaction strength of the interactive domain change between roots and leaves and two distinct interactive domains can be distinguished (Fig 2A). An area encompassing the thalianol cluster engages in very strong three-dimensional contacts in roots while in leaves the location of the interactive domain shifts downstream and covers the cluster and a region downstream of the cluster. This domain is larger than in roots and the intensity of domain-wide 3D interactions is reduced (Fig 2A). We then performed A/B compartment analyses for all chromosome arms of our Hi-C maps and analysed the compartment association of the thalianol cluster. In leaves, we found the cluster to be positioned within a B compartment, a more compact structural domain with increased intradomain contacts, and in roots, we found the cluster to be localised within an A compartment with depleted intradomain contacts and a looser structure (Fig S7).

After detecting these differences, we revisited the comparative analysis of our Hi-C libraries to identify differential interactions associated with the cluster. By calling genome-wide significant differential interactions we observed a striking pattern of both local and global changes to the three-dimensional structure of the thalianol cluster. Our comparative analysis shows that the chromosomal region encompassing the thalianol cluster ranks amongst the most differentially interacting areas of the *A. thaliana* genome in the root – leaf comparison (Fig 2B). In leaves, the cluster engages in significantly enriched interactions towards the pericentromeric areas of the chromosome (Figs 2C, S8). In contrast, root-specific interactions are significantly elevated towards the long arm of chromosome 5 outside the pericentromeric region (Fig 2C, SI Dataset 3).

We identified similar conformational features for the marneral and arabidiol/baruol gene clusters (Figs 2B, S9). These patterns reflect the different transcriptional states of the clusters. In leaves, when the biosynthetic gene clusters are silenced, the clusters are directed towards heterochromatic areas of the genome and in roots, when active, are located towards open, transcriptionally active areas of the genome. Accordingly, the genes located in regions differentially interacting with metabolic gene clusters show significantly lower expression levels in leaves compared to roots (2D).

To corroborate the change in localisation we performed 3D DNA FISH analysis of the thalianol cluster. Chromocenters are preferentially associated with the nuclear periphery in *A*. *thaliana* nuclei (43, 44) and as such we analysed cluster localisation towards the periphery in nuclei of roots and leaves. We show that the cluster region strongly associates with the nuclear periphery in leaf tissues while in roots this association is significantly reduced (Fig. 2E).

Conformational switching accompanies changes in transcriptional activity at the thalianol metabolic gene cluster

To better define the changes to the 3D chromosome architecture at metabolic gene clusters we established a Capture Hi-C (CHi-C) protocol for *A. thaliana* (45, 46). We defined a set of genomic regions with sizes between 200 kb and 600 kb and designed capture probes that cover all restriction fragments within. We chose the regions based on the annotation of metabolic gene clusters and distribution along all chromosomes and central chromosome features (SI Dataset 4). We obtained libraries with similar yield of valid di-tags but with much improved sequence depth at the captured sites compared to our Hi-C library and previously published data (Tables S3, S4, S5 and S6).

Analysis of the CHi-C library enabled us to recapitulate the chromosome-wide interaction switch of the thalianol cluster (Fig S10). Furthermore, it enabled us to precisely analyse the local variations in three-dimensional chromosome structure at the thalianol cluster coinciding with the transcriptional 'on' and 'off' state. In roots, when active, the cluster is located within an interactive domain that consists of two layers of variable strength. The smaller and higher intensity domain precisely demarcates the thalianol cluster, ranging from *THAA2*, the peripheral cluster gene, to the *THAS* gene with a size of 50 kb. The larger domain extends from the non-cluster gene *At5g47910* to the thalianol cluster (Figs 3A, S11A).

In contrast to the root specific domain, the interactive domain formed in leaves, when the cluster is silenced, is larger in size and starts at the *THAA2* gene, the peripheral cluster gene, and ends at the non-clustered and non-co-expressed genes At5g48150 and At5g48160. Overall the domain is 110 kb in size and covers the thalianol cluster and a group of genes with increased expression level in roots compared to leaves (At5g48070 to At5g48140) (Fig 3B, S11A, S11B). Structural modelling of the major 3D domains associated with the thalianol cluster indicates that when active the locus assumes a compact conformation and when silent it is incorporated into a chromosomal loop (Fig S12).

Differential visualisation of CHi-C maps of root and leaves shows a striking border between the silencing and activating domains around the gene *At5g48050*, an area depleted of obvious regulatory elements (Fig 3C). Independent 3C experiments that measure contact intensity between individual restriction fragments corroborate the identified domain structure (Fig 3E).

This association with two local interactive domains, i.e. a bimodal chromosomal configuration, may allow the thalianol cluster to read regulatory information from two distinct chromosomal areas (Fig 3D). For the HoxD cluster in mice, location between two topologically associated domains (TADs) and dynamic association to either of them ensures collinear distribution of Hox transcription factors to the correct developmental body structures (47).

It is striking that the chromocenter interactions of the clusters are partly driven by contacts of the cluster downstream region that are specific for the silencing domain (Fig S10). We suggest a three-fold mechanism in which the cluster is brought in proximity to the envelope: (a) the cluster is released from an interactive domain associated with strong transcriptional activity; (b) establishment of new contacts with a region downstream of the cluster; (c) positioning towards heterochromatic chromosomal areas near the nuclear periphery.

Similar to the thalianol cluster, the 3D chromosome architecture at the marneral cluster is associated with a complex pattern of local and regional interactions that differ between roots and leaves. As seen in our Hi-C maps, a large regional interactive domain is formed between the marneral cluster and a genomic area 300 kb away from the cluster. This domain is specifically formed in leaf organs when the cluster is silenced (Figs. S13, S14A, S14B). The CHi-C analysis reveals an additional local interactive domain encompassing the marneral cluster that exhibits more pronounced contact intensities in roots, where the cluster is transcribed, compared to leaves. This suggests a similar dual conformational switch between the actively transcribed and silenced forms of the cluster, as observed for the thalianol cluster (Figs. S13, S14C, S14D). At the arabidiol/baruol cluster we observed a single local interactive domain that precisely encompasses the cluster and shows increased contact frequency in leaves compared to roots and therefore negatively correlates with transcriptional activity (Figs. S15, S16).

Of note, the establishment of dynamic local 3D domains associated with clusters of coregulated genes may not be restricted to our target clusters. We observed strong interactions between a cluster of homologous ribulose biphosphate carboxylases genes encoding small subunits of the Rubisco enzyme and an adjacent region (48). These interactions are specific for roots where the cluster is silenced and are not detectable in leaves when the cluster is expressed (Figs. S17, S18). Future single-cell Hi-C analysis may further refine the correlation between chromosomal conformation of clusters and different expression states.

Cluster-associated silencing domains are lost in H3K27me3 mutant

Our earlier work and studies in filamentous fungi have suggested that biosynthetic gene clusters are delineated by conserved chromatin modifications (20, 28, 49, 50). Amongst them is H3K27me3, a well-described histone mark primarily associated with gene silencing (51, 52). Peaks of H3K27me3 are detectable at all three metabolic gene clusters investigated here and we have previously shown that cluster expression levels are elevated in *A. thaliana* mutant lines with reduced H3K27me3 levels (Figs. S18, S19, S20) (20, 52). Recent Hi-C analyses of *A. thaliana* chromosomes have described small interactive domains and chromatin loops that are enriched for H3K27me3 marks (38, 53). Therefore, we decided to re-analyse available Hi-C maps of H3K27me3 mutants and monitor the cluster-associated interactive domains (38).

For the thalianol, marneral and arabidiol/baruol clusters we found a significant reduction in the interaction strength of the associated interactive domains in H3K27me3 depleted chromatin suggesting a role for this chromatin mark in their formation (Figs. 4A, B, C, S22A, B, C). In animals, a spatial domain that connects different gene clusters in 3D space has been shown to be constrained by H3K27me3 histone modifications (32, 34). Similarly, we detected high interaction counts between the thalianol and marneral gene clusters, which are separated by 2.4 Mb on chromosome 5 (Fig S23A). The interaction intensity was strongly elevated in wild type versus H3K27me3 levels (Fig S23B). In embryonic cells of mice, H3K27me3 labelled three-dimensional domains are formed within central active nuclear regions in contrast to the peripheral localisation of the H3K27me3 marked interactive domains identified here (34). In *Neurospora crassa*, loss of H3K27me3 marks in chromatin mutants leads to relocalisation of subtelomeric regions towards the interior of the nucleus and upregulation of target genes (54).

Interestingly, at the cluster of homologous ribulose biphosphate carboxylases genes we did not observe strong H3K27me3 markings in non-expressing root tissues (Fig. S24). In contrast, we identified significant cluster-associated H3K4me3 markings, well-described histone marks associated with gene activation, in the expressing leave tissues (Fig. S24). We did not, however, detect strong enrichment of H3K4me3 markings at biosynthetic gene clusters (Figs. S19, S20, S21).

Metabolic gene clusters reside in local interactive domains in diverse plant species

Next, we asked whether biosynthetic gene clusters are similarly located within distinct interactive domains in other plant species. To address this question, we analysed available Hi-C maps of tomato, maize and rice (41, 56). Each of these species contains at least one well-described metabolic gene cluster (Fig S25) (14). Chromosome topology in tomato, maize and rice is characterised by a more pronounced structuring of chromosomes into TADs compared to chromosome topology in A. thaliana (41, 56, 57). TADs are chromosomal regions with extensive internal chromatin contacts and limited interactions with adjacent regions (58, 59). As such, they resemble the cluster-associated interactive domains in A. thaliana described here. We therefore identified TADs in the respective genomes and analysed their association with the maize DIMBOA and rice momilactone and phytocassane clusters, as well as the tomatine biosynthetic gene cluster in tomato (SI Dataset 5) (17, 60-62). We observed that each cluster is positioned within a defined TAD that encompasses all individual cluster genes (Fig 5A, B, C, D). As seen for the biosynthetic gene clusters in A. thaliana and the ergot alkaloid EAS cluster of E. festucae cluster-associated TADs include additional genes outside the respective clusters (Fig 5A, B, C, D) (35). We did not observe obvious long-range contacts with other regions of the genome for the investigated clusters. TADs in maize, rice and tomato are suggested to be separated by different expression and epigenetic states but are not associated with co-expression of the genes located within the same domain (41). All four clusters investigated here show strong co-expression pattern and for clusters in rice and maize we have previously reported an enrichment for H3K27me3 chromatin modifications (20, 63). Analysis of gene expression datasets associated with the investigated Hi-C maps show high expression levels for the tomatine cluster, medium expression levels for the DIMBOA cluster and very low expression levels for both rice cluster (Fig. S26A, B, C, D). This suggests that formation of cluster-associated domains is not restricted to specific expression states. Future comparative studies analysing chromosome architecture under conditions with variable cluster expression states in maize, rice and tomato should shed light on the structural flexibility of the cluster-encompassing TADs in these other species.

Conclusions.

In summary, we report that metabolic gene clusters reside in defined local interactive domains in plant genomes. In *A. thaliana*, the structure of these domains is flexible and changes its configuration between organs that express or do not express the cluster in question.

Local interactive domains surrounding clusters of genes may insulate these clusters from their chromosomal neighbourhood. This may prevent the spreading of repressive and active chromatin environments at gene clusters into nearby chromosome areas (64, 65). Cluster-associated interactive domains may further constitute local microenvironments that support tight co-regulation of gene expression. Local domains with increased internal contacts have been observed for neighbouring, functionally unrelated genes with shared transcriptional and epigenetic states in different eukaryotic species (66, 67). The formation of these domains may indicate a general principle in eukaryotic genome organisation and may provide a structural platform for evolution of functionally related and co-ordinately regulated gene clusters.

Structural flexibility of 3D domains at biosynthetic gene clusters is accompanied by repositioning of clusters inside the nuclear space. Peripheral localisation of clusters is observed in non-expressing organs and interior localisation in expressing organs (Fig S27). As such, these cluster regions are amongst the most dynamic regions in the *A. thaliana* genome. The relocation of clusters towards heterochromatic areas of the nucleus may be important in the efficient silencing of these clusters. It has been shown that mis-expression of thalianol cluster genes leads to severe developmental defects in the plant and so tight coordinate regulation of the thalianol cluster genes is likely to be critical for survival (24). The co-localisation of pathway genes in operon-like gene clusters may directly support the formation of single 3D domains. Such conformation may in turn facilitate co-ordinate engagement of these genes in nuclear re-positioning and transcriptional co-regulation as compared to a scenario in which genes are dispersed in different chromosomal locations and embedded in separate 3D domains.

Furthermore, we show that loss of the histone mark H3K27me3 is associated with alterations in the interaction intensity of the cluster-associated chromosome architecture, supporting the

important role for this chromatin modification in shaping chromosome structure in *A*. *thaliana* (38, 68).

Our results reveal the complex chromosomal architecture surrounding metabolic gene clusters and shed light on the potential mechanisms that constrain co-expression within clusters of eukaryotic genes. We show that clustering of genes on the linear eukaryotic chromosome is accompanied by compartmentalisation of the three-dimensional chromosome. Furthermore, we provide evidence that the spatial organisation of plant chromosomes is plastic. These advances will provide the basis for future studies to better understand the role of chromosome organisation in defining gene cluster structure, expression and evolution.

Materials and Methods

Arabidopsis thaliana plants used in this study were of the Col-0 wild type. For all experiments, *A. thaliana* seeds were surface sterilised and grown vertically on petri dishes containing Murashige and Skoog plant salt medium supplemented with 0.5 % phytagel and 0.75 % sucrose (69). Plants were grown at 22 °C with a 16 h light/8 h dark photoperiod for seven days. Triplicate 3C, duplicate Hi-C and CHi-C and triplicate RNA-seq experiments were performed as described in SI Materials and Methods. DNA FISH experiments were essentially performed as described before and are outlined SI Materials and Methods (70, 71). All Hi-C, CHi-C and RNA-seq data were deposited to the National Center for Biotechnology Information (NCBI) Gene Expression Omnibus (GEO) database (accession no. PRJNA576277) (72). *A. thaliana* wild type and H3K27me3 mutant Hi-C datasets (accession no. SRP043612) and ChIPseq datasets (GSE108960, GSE108960) as well as tomato, rice and maize Hi-C datasets (PRJNA486213 and PRJNA391551) were previously reported (38, 41, 55, 73-76).

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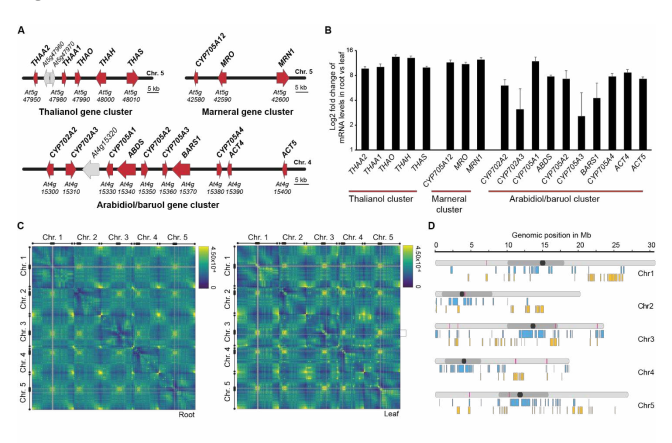
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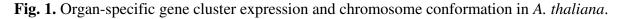
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Figures





(A) The thalianol, marneral and baruol/arabidiol gene clusters. Red arrows, cluster genes.Grey arrows, uncharacterised genes.

(B) Relative quantification of mRNA levels of thalianol, marneral and arabidiol/baruol cluster genes in the roots and leaves of 7-day old seedlings as assessed by RNA-seq analysis of three biological replicates. Error bars indicate standard error of logFC.

(C) Two-dimensional Hi-C interaction maps of the 3D conformation of *A. thaliana* chromosomes in roots and leaves. Chromosomes are labelled from left to right and top to bottom. Centromeric and pericentromeric regions are marked with black rounded boxes. Telomeres are marked with circles. Yellow to blue colouring indicates strong to weak interaction tendency. Genomic bin size: 25 kb.

(D) Location of differentially interacting regions on *A.thaliana* chromosomes. Distribution of the top 10% regions engaging in most differential interactions throughout the genome, are shown (p-value < 0.01). Regions with increased interaction tendency in leaves and roots are

shown in blue and yellow, respectively. Note: regions may interact with different chromosomal sites in root and leaf and may thus show both blue and yellow markings. Light grey, chromosome arms; dark grey, pericentromeric region; black, centromeric region; pink, IHIs.

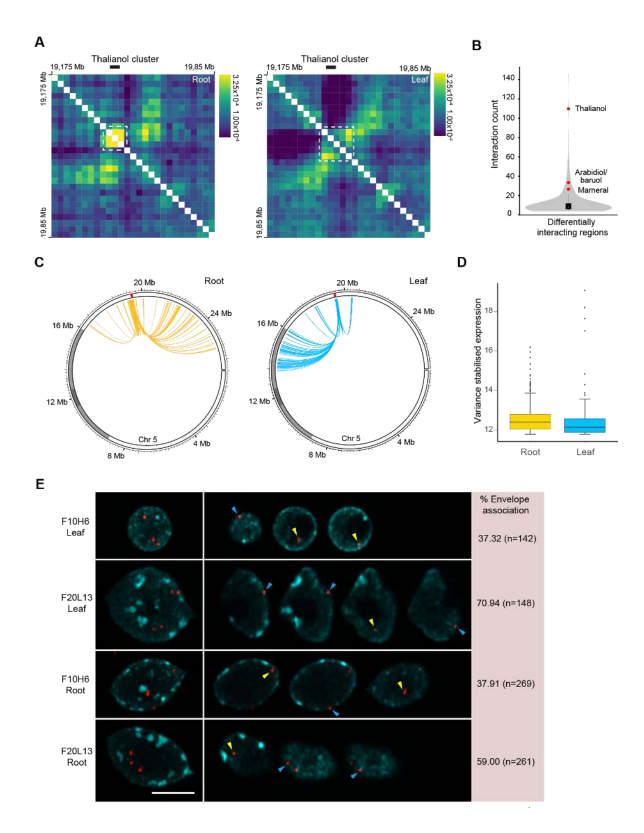


Fig. 2. Integration of the thalianol gene cluster in the 3D nuclear space

(A) Detailed two-dimensional Hi-C maps of the genomic region surrounding the thalianol cluster. White boxes indicate cluster-associated local interactive domains in roots (left) and leaves (right). Chromosome co-ordinates are labelled from left to right and top to bottom.

Yellow to blue colouring indicates strong to weak interaction tendency. Genomic bin size: 25 kb.

(B) Violin plot of frequency and range of interaction counts of differentially interacting regions. The thalianol, arabidiol/baruol and marneral cluster-associated interactive domains are indicated by the red dots.

(C) Circos plots showing intrachromosomal differential genomic interactions for the thalianol cluster in leaves and roots. Chromosome arms are shown in white, pericentromers in light grey, centromeres in dark grey and the thalianol cluster in red. Each arc/connection represents a significantly enriched interaction (p-value < 0.01).

(D) Box plots showing normalised expression levels of genes within chromosomal regions of differential interaction with the thalianol, marneral and arabidiol/baruol clusters. Variance stabilized gene expression values are shown for differentially interacting regions in roots and leaves. Gene expression levels are significantly higher in roots compared to leaves ($p = 1.188 \times 10^{-7}$, Wilcoxon rank-sum test).

(E) 3D DNA FISH analysis of nuclear envelope association of the thalianol cluster. Representative images and envelope association percentage for bacterial artificial chromosomes BAC F20L13 (covering thalianol cluster) and a control, BAC F10H6 (covering an area of open chromatin). The nuclei in the left panels are maximum intensity projections of a *z*-stack through each nucleus. On the right, each locus is shown in a single optical section through the nucleus in the *xy*-direction. Note, the higher envelope association rate of F20L13 compared to F10H6 in leaves ($p = 1.11 \times 10^{-8}$, two-sided Fisher's exact test) and the significantly lower association of F20L13 in roots compared to leaves (p = 0.019, two-sided Fisher's exact test). Scale bar 5 µm. Blue arrow, foci associated with the nuclear envelope; yellow arrows, foci not associated with nuclear envelope. n, number of foci counted.

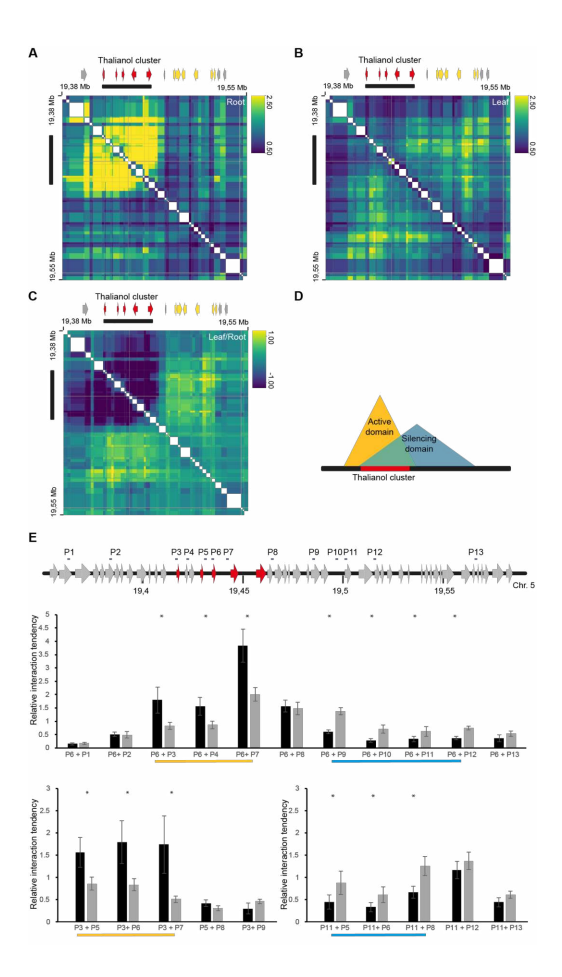


Fig 3. Local interactive domains at the thalianol cluster.

(A, B) Two-dimensional CHi-C interaction map of the thalianol cluster region in roots (A) and leaves (B). Yellow to blue colouring indicates strong to weak interaction tendency.Genomic bin size: restriction fragment.

(C) Differential two-dimensional CHi-C interaction map of the thalianol cluster region in leaves vs roots. Co-ordinates are labelled from left to right and top to bottom. Cluster genes are shown in red, and additional genes with increased transcript levels in root vs leaves in orange. Genes at the borders of the cluster-associated domain are in grey (from left to right – *At5g47910, At5g48050, At5g48150, At5g48160*). Yellow to blue colouring indicates strong to weak interaction tendency. Genomic bin size: restriction fragment.

(D) Model of interactive domains at the thalianol cluster. A strong interactive domain is formed at the thalianol cluster during active transcription and a weaker but larger domain is formed during transcriptional repression.

(E) 3C analysis of the extended thalianol cluster region. Top, the chromosomal area around the thalianol cluster. Cluster genes are shown in red and non-cluster genes in grey. The histograms below display the 3C interaction profile of different sites in the extended cluster region. Interaction tendencies in roots and leaves are shown in black and grey bars, respectively. Significantly increased interactions in roots are underlined in yellow and significantly increased interactions in leaves are underlined in blue. Asterisk indicates significant difference between interaction tendencies (Student's T-test, p < 0.01).

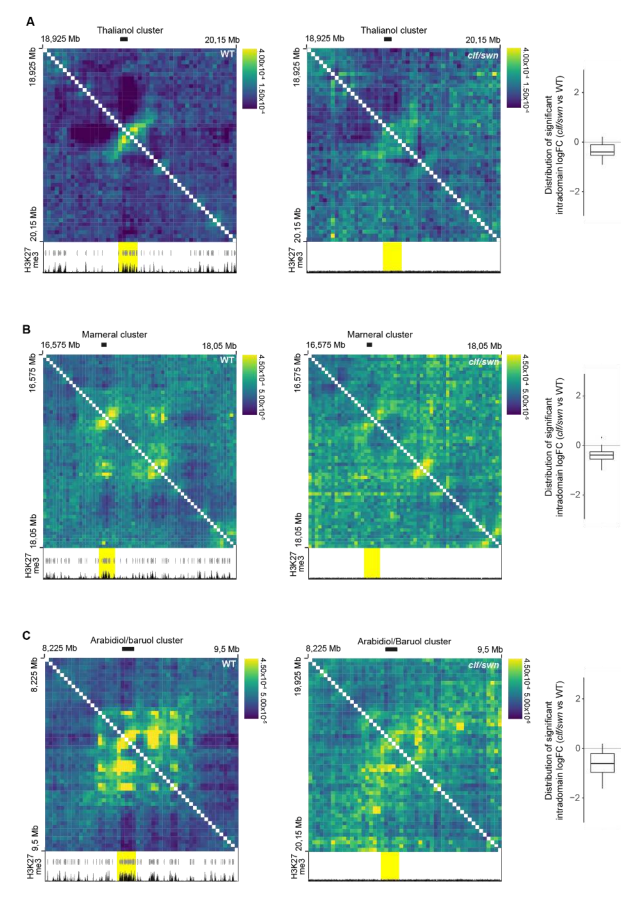


Fig. 4. Loss of cluster-associated interactive domains in a H3K27me3 mutant.

(A, B, C) Two-dimensional Hi-C interaction maps of the 3D conformation surrounding the thalianol cluster (A), marneral (B) and arabidiol/baruol (C) clusters in seedlings. Left, wild type; right *clf/swn* mutant. Chromosomes are labelled from left to right and top to bottom. Yellow to blue colouring indicates strong to weak interaction tendency. Genomic bin size: 25 kb. The boxplots on the right show logarithmic Fold Change values for the wild type vs the *clf/swn* mutant intradomain interaction counts. The tracks at the bottom of each panel show significant peaks and enrichment tracks of H3K27me3 markings in wild-type (left panels) and *clf/swn* double mutants extracted from Shu et al (2019) (55). In yellow, area of cluster associated interactive domain.

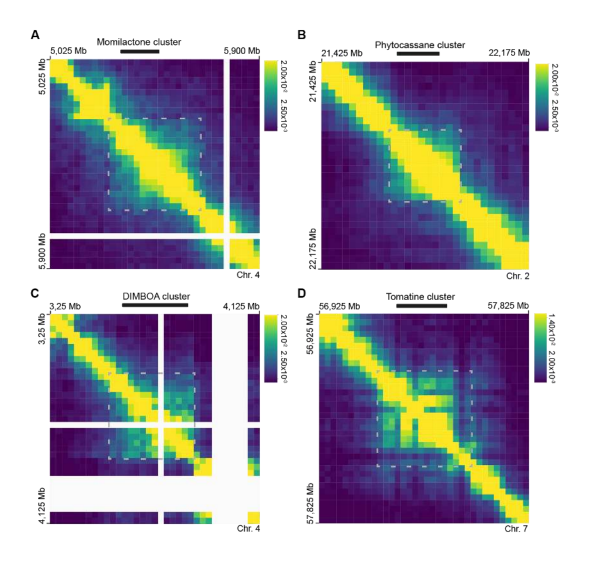


Fig. 5. Metabolic gene clusters from other plant species are located in single TADs.

(A, B, C, D) Two-dimensional Hi-C interaction maps of the 3D conformations of the momilactone (A) and phytocassane (B) cluster-associated genomic regions in rice; the DIMBOA cluster-associated genomic region in maize (C); and the tomatine cluster-associated genomic region in tomato (D). Chromosomes are labelled from left to right and top to bottom. The biosynthetic gene clusters are indicated with black line and associated TADs are shown in the grey boxes. Yellow to blue colouring indicates strong to weak interaction tendency. Genomic bin size: 25 kb