

Seasonal stability and dynamics of DNA methylation in plants in a natural environment

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20 **Abstract**

21 Organisms survive in naturally fluctuating environments by responding to

22 long-term signals, such as seasonality, by filtering out short-term noise. DNA

23 methylation has been considered a stable epigenetic mark but has also been

24 reported to change in response to experimental manipulations of biotic and

25 abiotic factors. However, it is unclear how they behave in natural environments.

26 Here, we analyzed seasonal patterns of genome-wide DNA methylation at a

27 single-base resolution using a single clone from a natural population of the

28 perennial *Arabidopsis halleri*. The genome-wide pattern of DNA methylation was

29 primarily stable, and most of the repetitive regions were methylated across the

30 year. Although the proportion was small, we detected seasonally methylated

31 cytosines (SeMCs) in the genome. SeMCs in the different contexts showed

32 distinct seasonal patterns of methylation. SeMCs in CHH context were detected

33 predominantly at repetitive sequences in intergenic regions. Additionally, we

34 found that CHH methylation within *Ahglc* locus showed a seasonal pattern

35 that was negatively associated with changes in gene expression. Gene-body CG

36 methylation (gbM) itself was generally stable across seasons, but the levels of
 37 gbM were positively associated with seasonal stability of RNA expression of the
 38 genes. These results suggest the existence of two distinct aspects of DNA
 39 methylation in natural environments: sources of epigenetic variation and
 40 epigenetic marks for stable gene expression.

41 **Keywords:** *Arabidopsis halleri*, DNA methylation, Natural environment,
 42 seasonal changes, seasonally methylated cytosines

43 **Introduction**

44 DNA methylation at cytosine residues is an epigenetic mark that can be
 45 maintained through cell divisions in a wide range of eukaryotic genomes (1). The
 46 previous analyses in diverse organisms have revealed genomic distributions of
 47 DNA methylation vary among organisms (2-7). In plants, DNA-methylation varies
 48 both between and within species (8), and sometimes is associated to phenotypic
 49 variation (9-11). Although the level and patterns of DNA methylation are
 50 heritable to a certain extent, the mechanisms that produce and maintain
 51 epigenetic variation across generations are largely unknown.

52 DNA methylation can vary between individuals also by non-genetic causes.
 53 It has been shown that both biotic and abiotic treatment can modify DNA
 54 methylation (12-14). Because of its semi-stable and semi-labile nature,
 55 non-genetic changes in DNA methylation are not explained by simple
 56 environmental effects. For example, even in a genetically homogeneous
 57 background under stable laboratory conditions, epigenetic variation in DNA
 58 methylation can occur during repeated self-pollination in a transgenerational

manner (15, 16). Therefore, it is difficult to predict whether DNA methylation is stable or dynamic under natural conditions. Recently the need for '*in natura*' studies has been highlighted in order to understand how organisms respond to environmental signals by filtering out innumerable fluctuations and noise (17, 18). In the temperate regions, seasonality is the most prominent cause of environment fluctuations. However, we still do not understand how DNA methylation behaves across seasons.

In order to reveal the seasonal dynamics of DNA methylation, here, we conducted an "*in natura*" study on genome-wide DNA methylation using a single clone growing in a population of *Arabidopsis halleri* (L.) O'Kane & Al-Shehbaz subsp. *gemmaifera* (Matsum.) O'Kane & Al-Shehbaz (hereafter referred to as *A. halleri*), a close relative of *Arabidopsis thaliana*. Its clonal propagation and perennial life-cycle allowed us to sample leaves all year round from the single clonal individual (19). We studied seasonal dynamics of key flowering-time genes and whole transcriptome previously in the site (20, 21). In this study, to understand the dynamics of DNA methylation, we performed whole-genome

bisulfite sequencing (WGBS) at 1.5-month intervals, over a year, under natural conditions. We adopted the strategy of monitoring the seasonal pattern in a single clonal individual with a uniform genetic background.

DNA methylation occurs in three contexts, according to the flanking sequence, i.e. CG, CHG and CHH (H = A, C, or T). The former two form symmetrically and the latter one asymmetrically, in terms of sequences on the complementary strands. Since these contexts of methylation are distinctly regulated and associated with DNA replication, histone modification, and non-coding RNA production (22-24), we examined seasonal patterns of DNA methylation by conducting single-base resolution analyses.

The results suggested the existence of seasonally methylated cytosines (SeMCs) in the genome, at least for the examined clone. Interestingly, DNA methylation changes occurred in a context-dependent manner. There were distinct patterns of seasonal changes among CG, CHG, and CHH methylation. Moreover, our analysis revealed that genic CG methylation, i.e. gene-body methylation (gbM), was seasonally stable by itself, and was associated with

seasonal stability of RNA expression. This study suggested not only that there is a dynamic nature in DNA methylation in plants in their natural habitats, but also highlights the implications of DNA methylation in robust maintenance of stable RNA expression.

Results

Seasonal stability in large-scale distribution of DNA methylation. To investigate the dynamics of DNA methylation in a natural environment, we analyzed genome-wide DNA methylation in a natural population of *A. halleri* (Fig. 1 A and B). In the study site, the hourly air temperature ranged from -4.3 °C to 36.3 °C during the one-year study period, from Nov 2014 to Sep 2015 (Fig. 1 C). We collected leaves from a single clonal patch of *A. halleri* at 8 sampling times, at 1.5-month intervals across a year, and performed WGBS (Fig. 1 C). From this, we obtained a series of genome-wide DNA methylation data (Table S1).

Bulk DNA methylation levels were relatively constant across the eight time points, and ca 45%, 20%, and 6% of cytosines in the genome were kept

107 methylated in CG, CHG, and CHH contexts, respectively (Fig. S1A). The
108 large-scale distribution of DNA methylation was determined by the positions in
109 the genome, as represented by radial patterns in the circos plot for the longest
110 30 scaffolds (Fig. 2A). The distribution of methylated sites remained constant
111 across the 8 sampling times (represented by 8 concentric circles for each
112 methylation context in Fig 2A). We observed conspicuous aggregation of DNA
113 methylation on repetitive sequences (Fig 2A). For example, scaffolds 3 and 18 is
114 characterized by low and high density of repetitive sequences that corresponded
115 to relatively low and high methylation levels, respectively (Fig 2A). In the
116 comparative analysis between genic and repetitive regions for the whole
117 genome, the level of DNA methylation in repetitive sequences was higher than
118 that in genes (Fig. S1B). A similar pattern was confirmed by an analysis using
119 100 kbp windows for the whole genome (Fig. 2B). The level of DNA methylation
120 was correlated with density of repeats in each window for all three contexts in all
121 samples (e.g. Fig. 2B for Nov. 2014, Pearson's correlation coefficients were 0.64,
122 0.68, and 0.70 for CG, CHG, and CHH context, respectively; Table S2 for the

other sampling times). Although aggregation of DNA methylation at repetitive sequences was similar to the patterns previously reported in related Brassicaceae (2, 3, 25), seasonal stability in large-scale distribution of DNA methylation was reported for the first time here.

SeMCs: Seasonal dynamics of DNA methylation. Next, we searched for SeMCs by detecting differently methylated cytosines in the genome across the eight time points (Fisher's exact test; $P < 0.001$; FDR < 0.2). The majority of cytosines did not show statistical differences in methylation across the year, and the proportion of SeMCs was less than 0.5% of total cytosines (Fig. S2 A and B). Still, we detected 62,716, 47,140, and 179,385 SeMCs in CG, CHG, and CHH contexts, respectively (Fig. 3A). They showed diverse seasonal patterns in their level of methylation across the year (Fig. 3A). Interestingly, each context of DNA methylation showed distinct patterns of seasonal change: the number of SeMCs that peaked in a particular month was the highest in July, March, and September in CG, CHG, and CHH contexts, respectively (Fig. 3B). Differences among

contexts of SeMCs would reflect the responsiveness of the regulatory mechanisms of DNA methylation to environmental cues.

CHH SeMCs in repetitive sequences. The distribution of SeMCs in the genome differed between the contexts of DNA methylation. The majority of SeMCs in CHH context (CHH-SeMCs) were found in intergenic regions (those annotated as neither exons nor introns) while CG-SeMCs and CHG-SeMCs were in both genic and intergenic regions (Fig. 4A). Given that large fractions of the eukaryotic genome are intergenic regions and consist of repetitive sequences such as transposable elements (26, 27), we compared the level of CHH methylation for all repetitive sequences in the genome of *A. halleri*. The median level of methylation was the highest in autumn, i.e., September and November (Fig. 4B), and the pattern was similar to that of whole-genome CHH-SeMCs. An example of seasonal changes in CHH methylation level in a LINE/L1 showed low and high levels in February-March and September-November, respectively (Fig. 4C). The highest methylation levels in

autumn were detected in other selected families of repetitive sequences, such as LTR/Copia, LTR/Gypsy, LINE/L1, DNA/MULE-MuDR, DNA/hAT, and RC/Helitron (Fig. S3).

In addition, we found some repetitive sequences that showed a unique seasonal pattern of CHH methylation. For example, in the intronic repeat at the locus of *FLOWERING LOCUS C* homolog (*AhgFLC*). *AhgFLC* expression is regulated seasonally and suppresses flowering when it is upregulated (20). We found that a SINE-like repetitive sequence in the first intron of *AhgFLC* showed a seasonal change of CHH methylation. The level was the highest in March and the lowest in June (Fig. 4D). Interestingly, this seasonal pattern was opposite to that of RNA expression of *AhgFLC* (Fig. 4E). The presence of interactions between transcription and CHH methylation in the intragenic repeat at *AhgFLC* is expected.

Gene-body CG methylation (gbM) and constant RNA expression. Previous studies have reported that gbM is localized to active genes in a wide range of

eukaryotes (5, 6). In *A. thaliana*, DNA methylation at the gene body is primarily observed in CG context (2, 3, 25). We examined seasonal patterns of gbM in *A. halleri* and found that the level of gbM was constant across seasons, both in medians and quantiles of all genes (Fig. S4A), and at the individual gene level (Fig. S4B).

In order to examine the potential role of gbM in gene regulation, we tested whether the level of gbM associated to seasonal patterns of RNA expression using previously published data of a two-year seasonal RNA-seq of *A. halleri* at the same study site (21). We found that genes with high gbM often have constant levels of RNA expression across the year. One such example was *AhgPP2AA3* (Fig 5 A and B), the homolog of *PROTEIN PHOSPHATASE 2A SUBUNIT A3*, a gene that is known as one that is constantly-expressed under various conditions in *A. thaliana* (28). This is in contrast to the situation of *AhgFLC* locus, a representative of seasonally expressed genes, which lacks CG methylation from the entire locus, except for the repetitive sequence in the first intron (Fig. 5 C and D). Based on these observations, we hypothesized that gbM

would be associated to constant RNA expression across seasons. To test this hypothesis, we calculated the seasonal average and range of RNA expression for each gene, then compared them between genes with different levels of gbM (Fig. 5 *E* and *F*). Genes that were highly methylated in CG context showed relatively high average levels of RNA expression (Fig. 5*E*). The magnitude of seasonal changes in RNA expression of these genes decreased with increasing levels of gbM (Fig 5*F*). These results are consistent with the hypothesis mentioned above. The genes with the highest methylation level (in ‘group 5’ in Fig 5 *E* and *F*) were weakly enriched with four GOs of basic functions (Table S3).

Discussion

In this study, we examined the seasonal patterns of DNA methylation at CG, CHG, and CHH contexts in *Arabidopsis halleri*, under natural conditions. We identified the genomic sites that showed seasonal changes in DNA methylation. Our observations suggest that seasonal factors in the natural habitat could affect DNA methylation differently according to its context and location in the genome.

We would like to note here that, since our data set came from a single clonal individual for a single year, the genetic and non-genetic effects between different clones should be explored in future studies by applying WGBS to a population-level study with multiple repeat samples.

CHH methylation showed seasonal changes in diverse repetitive elements, and the level of DNA methylation was high in autumn, and low in winter. In *A. thaliana*, it has been reported that the level of CHH methylation in transposable elements was higher in 16°C environments relative to 10°C (13). Additionally, it has been suggested that RNA-directed DNA methylation (RdDM) pathway, which is required for maintenance and *de novo* methylation and mainly targets transposable elements, are involved in heat tolerance (29). This suggests that our observation on CHH SeMCs could reflect the temperature-dependence of the regulation of CHH methylation.

Seasonal changes detected in CHH methylation at a repetitive element in *AhgFLC* was an interesting example showing how genic heterochromatin behaves in genes. We observed that the seasonal pattern of CHH methylation in

this particular repetitive element was opposite to that of the expression of *AhgFLC* gene. *FLC* and its homologs in Brassicaceae species contain conserved structures in its second intron called VRE (vernalization response element), which is involved in responsiveness to cold treatment (30, 31). Expression of *FLC* has been reported to be repressed by long-term cold treatment (vernalization), accompanied by enrichment of tri-methylation of histone H3 lysine 27 (H3K27me3) at the TSS site in the beginning of cold treatment, and then by H3K27me3 accumulation across gene body region after the plants returned to warmer temperatures (32). On the contrary, the expression of genes is associated with the removal of H3K27me3 from their bodies (33). These reports and our results suggest that epigenetic regulation of *AhgFLC* gene might be responsible for seasonal removal of CHH methylation in its intronic repeat. The disruption of transcription in genes has been reported to induce ectopic CHH methylation in *A. thaliana* (34).

Large-scale patterns of DNA methylation were constant throughout the year for all three contexts, although there were seasonal changes in DNA

methylation at some cytosine sites. Constant levels of CG methylation were observed in gene body, and, furthermore, we found that genes with high levels of gbM showed seasonal stability in their gene expression. Although the function of gbM is still unclear (35), our results support previous reports, in *A. thaliana*, that gbM associates with genes modestly expressed among different tissues and experimental conditions (6, 36-38), and imply the importance of gbM under seasonal environment in a natural habitat.

Currently, we cannot entirely explain the patterns of DNA methylation in SeMCs, particularly in CG or CHG contexts. It is very likely that there are unidentified processes associated with environmental responses of DNA methylation that have not been studied under laboratory conditions. As mentioned above, the question remains of how the epigenetic variations in plants have been established in natural fluctuating environment (39, 40). Here, we focused on representative environmental effects on DNA methylation in a single clonal individual. In future studies of DNA methylation, we need to evaluate not only genotype effects, but also, so called 'lineage effects' that

represent past genetic and environmental effects. Furthermore, we should test to what extent seasonal changes in DNA methylation contribute to the variation in phenotypes in natural environments.

Materials and Methods

Plant materials. This study was conducted in a natural population of *Arabidopsis halleri* subsp. *gemmifera* located in central Japan (Omoide-gawa site, Naka-ku, Taka-cho, Hyogo Pref., 35°06' N, 134°55' E, altitude 190–230 m). Details of the study site have been described previously (19, 20). Leaf samples were collected at noon on the following dates: November 11 and December 22, 2014, and February 9, March 24, May 7, June 23, July 28, and September 8, 2015. In the study site, *A. halleri* forms patches of rosettes that consist of clonally propagated plants and sometimes genetically-related seed-originated plants. Originally six small patches of plants were chosen for leaf sampling, three of them were used for further analyses because the others were heavily damaged by deer herbivory between November 11 and December 22 in 2014. At each

sampling date, we harvested multiple mature and intact leaves from each plant. Each leaf was ca. 3-cm long, and weighed ca. 0.1 g. To detect DNA methylation and RNA expression in the same set of leaves, a small piece was collected from each leaf for RNA extraction. For DNA extraction, leaves were frozen in an ethanol bath with dry ice, then stored at -80 °C. For RNA extraction, the small pieces of leaves were stored in RNA*later* solution (Invitrogen, Carlsbad, CA, USA), then stored at -20 °C according to the manufacturer's instructions. The data for one patch were analyzed and shown in the main text, because the other two patches were found to be genetically mixed (Fig. S5). The samples analyzed here were confirmed to share whole-genome SNPs at levels that were safely judged to be a single clone (Fig. S5).

DNA extraction and WGBS library preparation. Genomic DNA was extracted from collected leaves (two leaves per plant, ca. 0.2 g) using the CTAB method (41). Libraries for WGBS were constructed as described previously (42). Sequencing was performed with the Illumina Hiseq 2500 system.

283

284 **Processing of WGBS data.** Sequenced reads were trimmed using
 285 Trimmomatic (43). Trimmed reads were mapped onto the genome sequence of
 286 *A. halleri* (44) using Bismark (45) and Bowtie2 (46). Repetitive sequences in the
 287 genome were detected using RepeatModeler (47), and repeats with at least 50
 288 bp length were used for further analyses. The level of DNA methylation was
 289 calculated for each context using the ratio of the number of methylated cytosines
 290 to the number of total sequenced cytosine included in any region of the genome.
 291 Efficient bisulfite treatment (> 99% in all samples) was confirmed using the level
 292 of DNA methylation in unmethylated lambda DNA (Table S4). SeMC was defined
 293 as any cytosine differently methylated between at least two time points, detected
 294 by Fisher's exact test, with genome-wide FDRs that were calculated using
 295 Storey's method (48). To draw the heatmaps of methylation of SeMCs, cluster
 296 3.0 (49) and Java Treeview (50) were used. To draw a circos plot for
 297 scaffold-wide DNA methylation, Circos software (51) was used. To make
 298 browser views of DNA methylation, Integrated Genome Viewer (52) was used.

To draw a dendrogram of collected samples, MethylExtract (53), VCF-kit (<https://vcf-kit.readthedocs.io>), and Dendroscope3 (54) were used.

Transcriptome analysis. RNA extraction and library preparation for RNA-seq were performed using the shotgun type method of BrAD-seq protocol (55). Sequencing was performed with the Illumina Hiseq 2500 system. Mapping of the reads and calculation of RPM were processed as described previously (21) except that the reference sequence was replaced with a newer annotation (44). To calculate seasonal average and range of expression of genes more precisely, weekly sampled RNA-seq data from a two-year period (21) were re-analyzed. To quantify the expression, Kallisto software was used (56). Calculation of seasonal average and range of RNA expression was based on previously described methods (21).

Data availability

Raw WGBS and RNA-Seq reads are available under the DDBJ BioProject

315 PRJDB7785.

316

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327

328 **Author contributions**

329 H.K. and T.K. conceived the study. T.I., H.N., H.K., and T.K. designed the
330 experiments. H.N. collected samples. Y.T., A.T., and A.F. performed

whole-genome bisulfite sequencing. N.E. and M.N.H. performed RNA sequencing. T.I. analyzed the data. T.I. and H.K. wrote the paper and incorporated comments from co-authors.

Conflict of interest

The authors declare that they have no conflicts of interests.

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Figure Legends

Fig. 1. Sampling site and dates for the seasonal DNA-methylation analysis in a natural population of *Arabidopsis halleri* subsp. *gemmifera*. (A) A photograph of the study site alongside a small stream through the *Cryptomeria* and *Chamaecyparis* plantation in Hyogo Prefecture, Japan. Red arrows indicate individuals of *A. halleri*. (B) An individual of *A. halleri* in the study site. The cage was used for protection against deer herbivory. White bar indicates 100 mm. (C) Sampling dates (8 time points) and temperature regimes during the study period. Sampling was performed every ca. 1.5 month for a year. The red line indicates hourly air temperature, and dotted lines represent timings of the sampling.

Fig. 2. Genomic pattern of DNA methylation across a year. (A) A circos plot showing seasonal patterns (8 time points) of DNA methylation in CG, CHG, and CHH contexts for the longest 30 scaffolds of *A. halleri*. On the outermost circle, scaffold positions are indicated by numbers and black bars with scales (one

scale = 0.4 Mbp). Each shaded/colored circle represents the scaffold-wide distribution of genomic attributes for each 100-Kb window. The second outermost circle represents density of repetitive sequences (including transposable elements). Each tile indicates the density: the lowest in white, the highest in black (0–0.5). The next 24 circles represent DNA methylation levels at 8 time points (starting from Nov 24, 2014 to Sep 8, 2015, towards the inner circles shown by the arrows) for CG, CHG, and CHH contexts, respectively. Colors in each tile indicates level of methylation: the lowest as blue, the highest as red (0–0.9 for CG context, 0–0.5 for the others). (B) Scattering plots comparing CG, CHG, and CHH DNA methylation against TE density for all 100 kbp windows across the 30 scaffolds.

Fig. 3. Annual patterns of seasonally-methylated cytosines (SeMCs). (A) Heatmaps of seasonally methylated cytosines (SeMCs) in CG, CHG, and CHH contexts (from left to right; n = 62,716, 47,140, 179,385, respectively). Each row indicates a series of DNA methylation ratios across a year in each position in the

500 genome (0: unmethylated, 1; fully methylated). The dendrogram on the left of
501 each heatmap represents the result of hierarchical clustering of SeMCs.
502 Distance in the dendrogram was based on Pearson's correlation coefficient. (B)
503 Barplot showing the distribution of peak timings of methylation level for the
504 SeMCs in CG, CHG, and CHH contexts (from left to right, respectively).

505

506 **Fig. 4.** SeMCs in the CG, CHG, and CHH contexts, and seasonal patterns in
507 CHH DNA methylation levels. (A) Pie charts indicating locations (exon, intron,
508 and intergenic regions) of seasonally methylated cytosines (SeMCs) in CG, CHG,
509 and CHH contexts (from left to right, respectively). (B) Boxplots of CHH
510 methylation at 8 time points in repetitive elements. The boxes span from the first
511 to the third quartiles, the thick black bars inside the boxes are the medians,
512 whiskers above and below the boxes represent $1.5 \times$ interquartile ranges from
513 the quartiles. Dotted line indicates the median CHH methylation level in Nov.
514 2014. (C and D) Browser views for seasonal patterns of CHH methylation on
515 repeat sequences in one of LINE/L1 sequences (C) and *AhgFLC* locus (D).

516 Orange rectangles indicates exons. A gray rectangle in the *AhgFLC* locus
 517 indicates a repetitive sequence. (E) Comparison between RNA expression of
 518 *AhgFLC* and CHH methylation of the repetitive sequence in its intron.

519

520 **Fig. 5. CG DNA methylation and stability of gene expression.** (A-D) Browser
 521 views of seasonal patterns of CG methylation (A and C) and the two-year
 522 dynamics of RNA expression level (B and D; from September 2011 to August
 523 2013) for *AhgPP2AA3* (*g04731*) and *AhgFLC* (*g19190*), respectively. Orange
 524 rectangles indicate exons. A gray rectangle indicates a repetitive sequence in
 525 the *AhgFLC* locus. (E, F) Boxplots showing relationship between DNA
 526 methylation in CG context and the average (E) and range (F) of RNA expression
 527 of genes. Only expressed genes (average expression level ($\log_2(\text{RPM}) > 1$)) are
 528 used for these analyses. Genes were split into five bins according to quintiles of
 529 genic DNA methylation in CG context. The boxes span from the first to the third
 530 quartiles, the bands inside the boxes are the medians, whiskers above and
 531 below the boxes represent $1.5 \times$ interquartile ranges from the quartiles. Different

letters represent significant differences between groups in the Mann-Whitney test, $P < 0.01$ adjusted for multiple comparisons.

Fig. S1. Genome-wide bulk DNA methylation level at eight time points across a year. Genome-wide bulk DNA methylation levels are shown in CG, CHG, and CHH context for the whole genome (A), and gene and repetitive sequences (B). Eight sampling timings are represented by the different shades.

Fig. S2. DNA methylation was seasonally stable at a majority of CG, CHG, and CHH sites. Histograms of seasonal differences of DNA methylation (max. – min.) for all cytosine sites (A), and for SeMCs (B) in CG, CHG, and CHH contexts.

Fig. S3. Seasonal patterns in CHH DNA methylation in repetitive elements that belong to the six major families of transposable elements (TEs). The boxes span from the first to the third quartiles, the thick black bars inside the boxes are the medians, whiskers above and below the boxes represent $1.5 \times$ interquartile

548 ranges from the quartiles.

549

550 **Fig. S4.** Seasonal patterns in CG gene body methylation (gbM) across a year.

551 (A) Boxplot of DNA methylation in genes in CG context at eight time points

552 across a year. The boxes span from the first to the third quartiles, the thick black

553 bars inside the boxes are the medians, whiskers above and below the boxes

554 represent $1.5 \times$ interquartile ranges from the quartiles. (B) A histogram of

555 seasonal difference for DNA methylation in genes (max. – min.) in CG context.

556

557 **Fig. S5.** Some patches of *A. halleri* turned to be genetically mixed. A

558 dendrogram shows Kimura's genetic distance using genome-wide SNPs among

559 the samples in three patches for eight timepoints. The samples from replicate 1

560 were judged to be a single clone. *A. halleri* is an obligate outcrossing species

561 with a self-incompatible breeding system, and therefore a large portion of SNPs

562 are expected to be heterozygous. Because heterozygous SNPs can be

563 designated as homozygous to either of the alleles in a certain probability, a

564 terminal radial branching is expected to be observed even for genetically

565 identical plants that belong to a single clonal patch (Rep. 1).

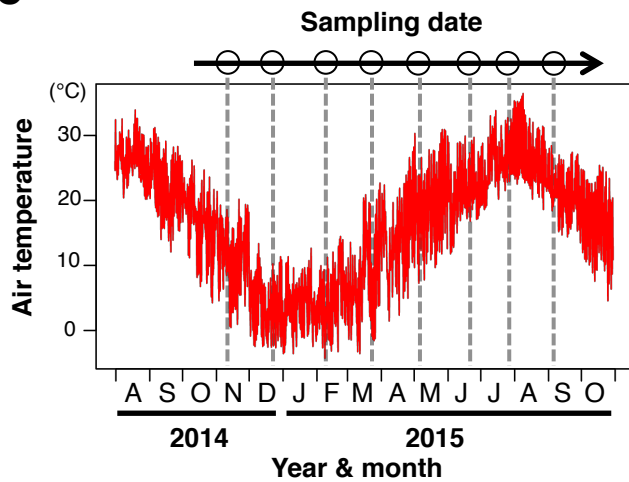
A**B****C**

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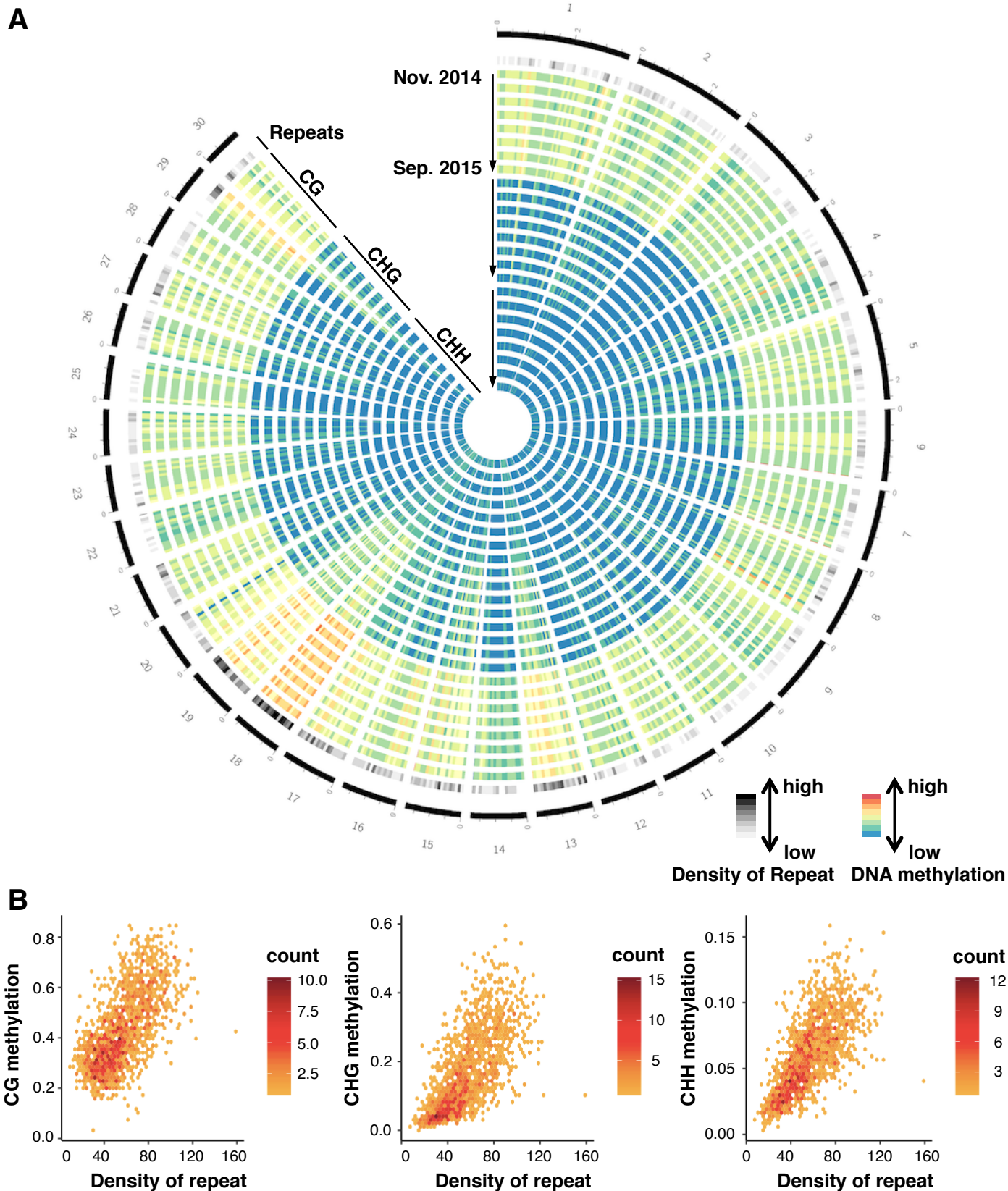


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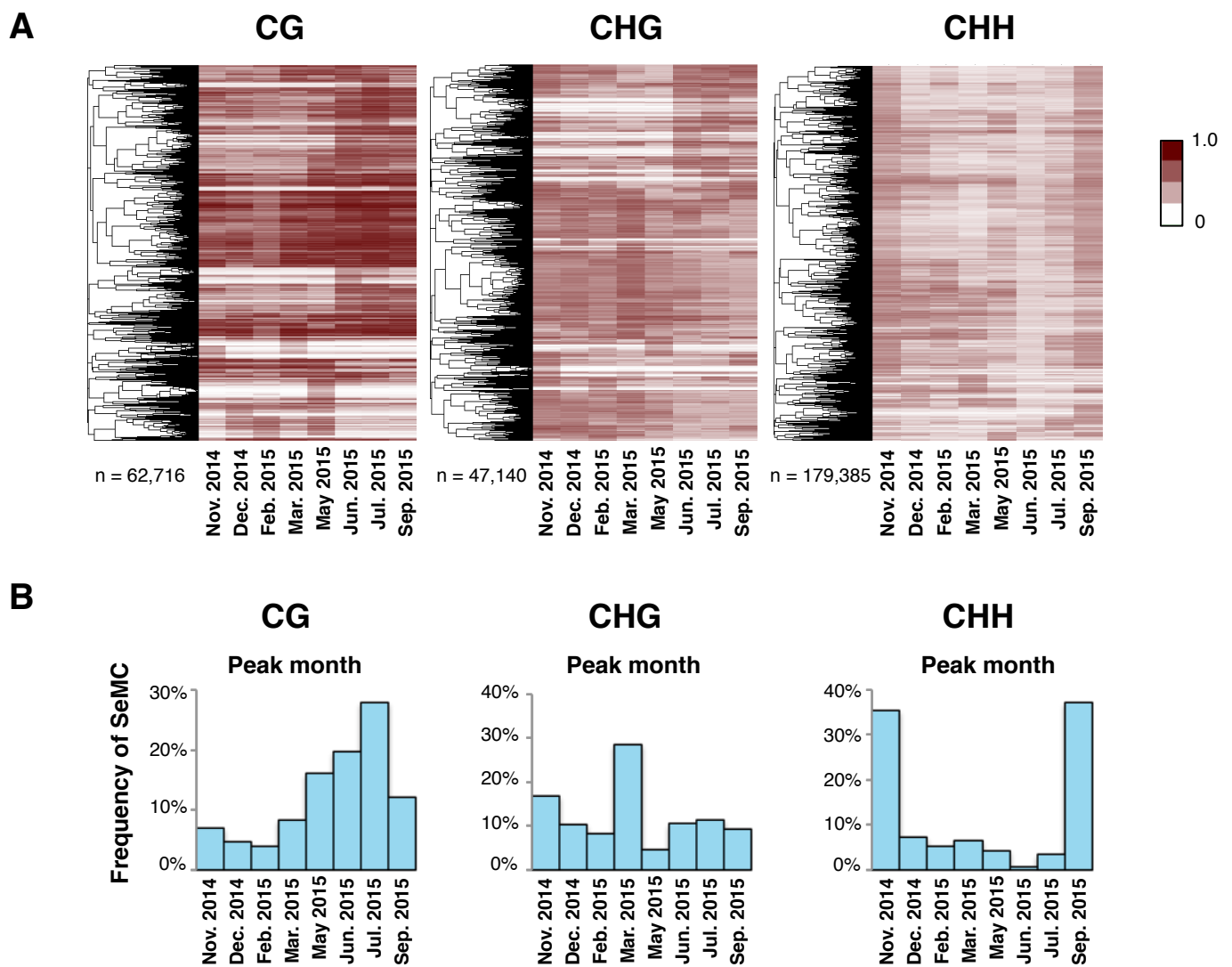


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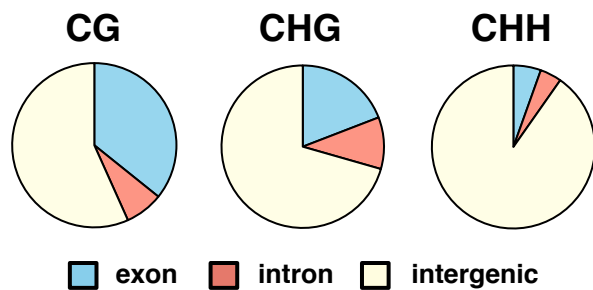
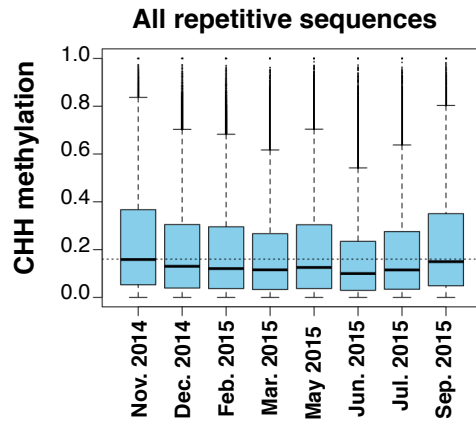
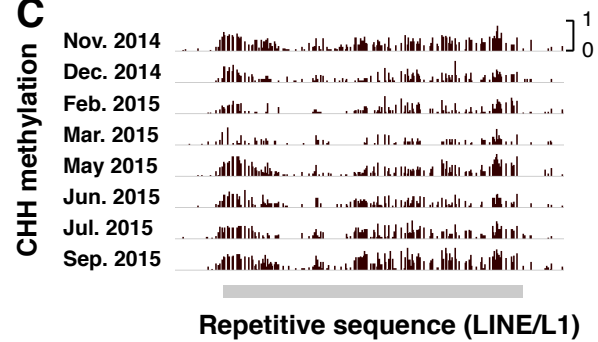
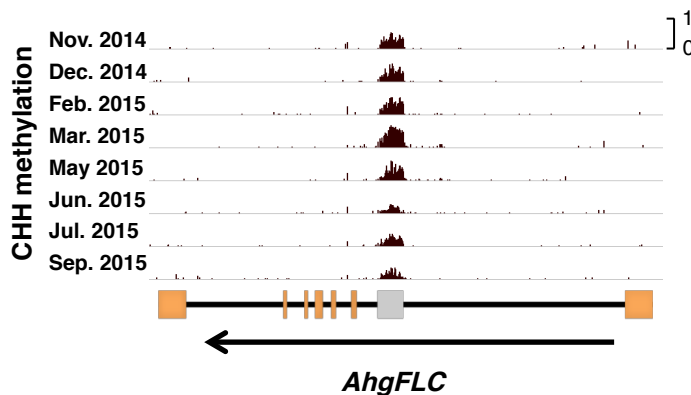
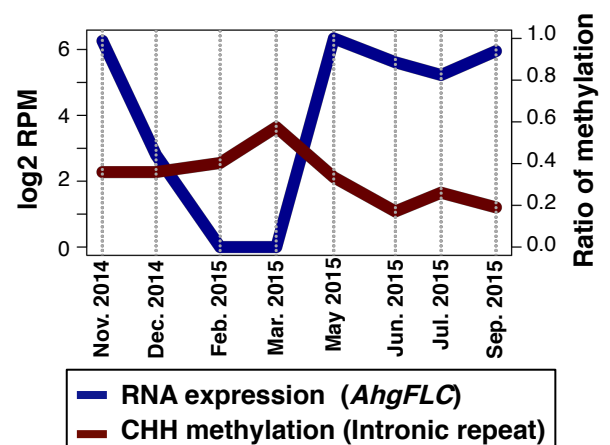
A**B****C****D****E**

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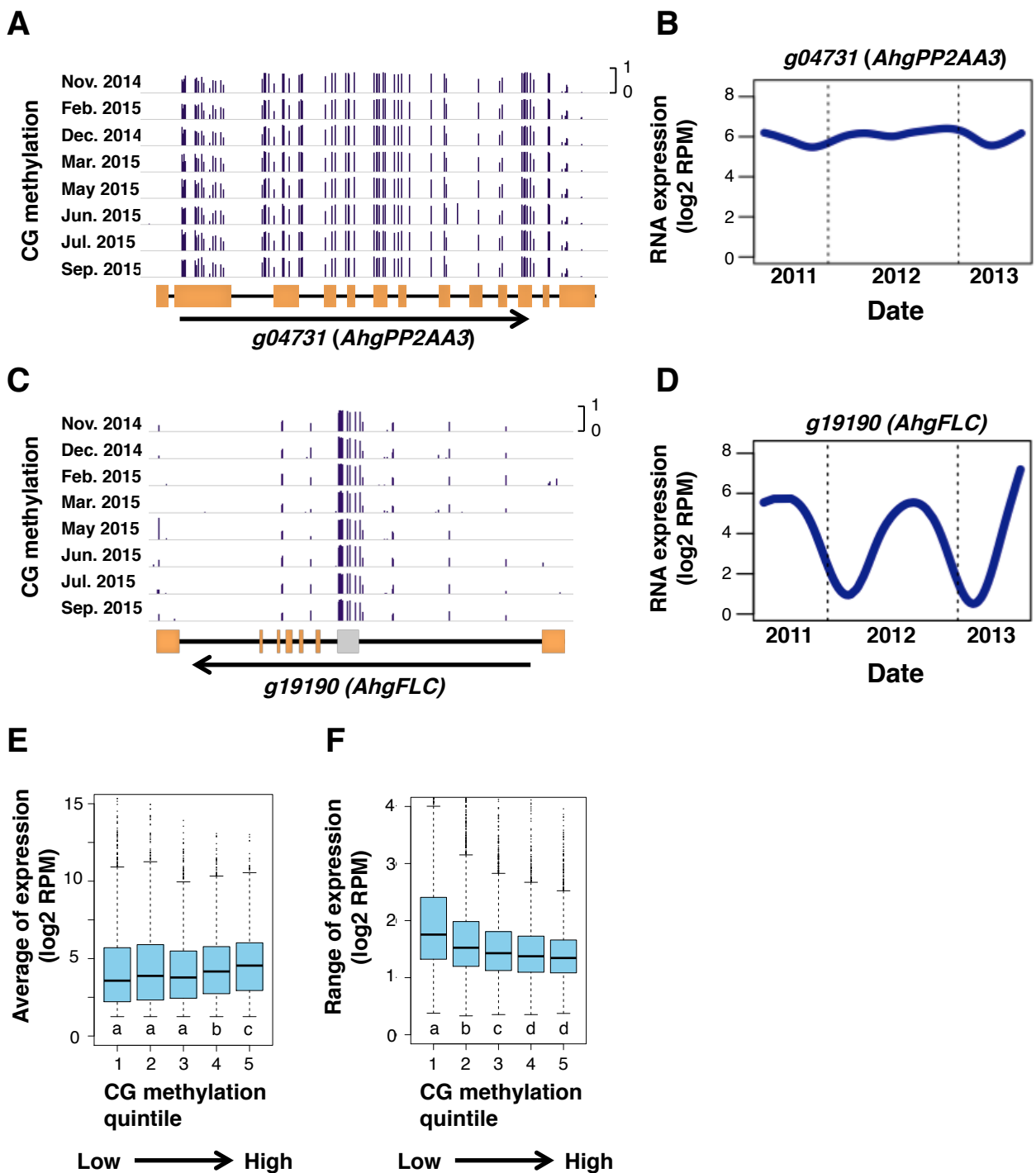


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