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Type II Protein Arginine Methyltransferase 5 (PRMT5) Is Required for Circadian Pperiod Determination in Arabidopsis Thaliana

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multiple clock genes and clock-controlled output genes. Thus, protein methylation represents a posttranslational modification that modulates circadian clock function.

Results

Forward Genetic Screen Based on *pCAT3:LUC* Expression. We screened an EMS-mutagenized M2 population of seedlings carrying *pCAT3:LUC*, which exhibits dusk-peaking expression (25), using seedlings entrained to LD cycles and released into DD, and we identified a long-period mutant, *prmt5-54* (H54). In DD, Columbia 3 (Col-3) seedlings had a period (\pm SEM) of 24.7 ± 0.1 h ($n = 11$), whereas H54 seedlings had a period of 26.9 ± 0.1 h ($n = 12$; significant by two-tailed *t* test at $P < 0.0001$) (Fig. 1 A and B). The long period of *pCAT3:LUC* expression also was seen in seedlings entrained in LD cycles and released into LL. In experiments where wild-type Col-3 seedlings had a period of 24.8 ± 0.2 h ($n = 15$), an F3 line derived from a backcross of H54 to Col-3 had a period of 27.31 ± 0.1 h ($n = 12$; $P < 0.0001$) (Fig. 1 C and D).

The Gene Mutated in H54 Is Required for Multiple Circadian Rhythms.

To determine whether the gene mutated in H54 is required for multiple clock outputs, we examined the rhythm in cotyledon movement in seedlings. Consistent with the period lengthening seen in *pCAT3:LUC* expression, the period in cotyledon movement was lengthened. Col-3 seedlings had a period of 24.8 ± 0.2 h ($n = 14$), whereas H54 seedlings had a period of 26.9 ± 0.1 h ($n = 12$; $P < 0.0001$) (Fig. S1 A and B).

H54 Identifies a Role for *PRMT5* in Circadian Clock Function. Low-resolution mapping by bulked segregant analysis (30) places H54 on the lower arm of chromosome 4, in a 42-cM region between *ciw7* and *nga1107* (Fig. S24). In organisms with smaller genomes, resequencing has proven a fast and economical alternative to map-based cloning for mutation identification (31–33). This approach is particularly attractive when the phenotypic characterization necessary for conventional positional cloning is made difficult by high individual variation among genotypically identical individuals, requiring confirmation in F3 lines, as is the case for

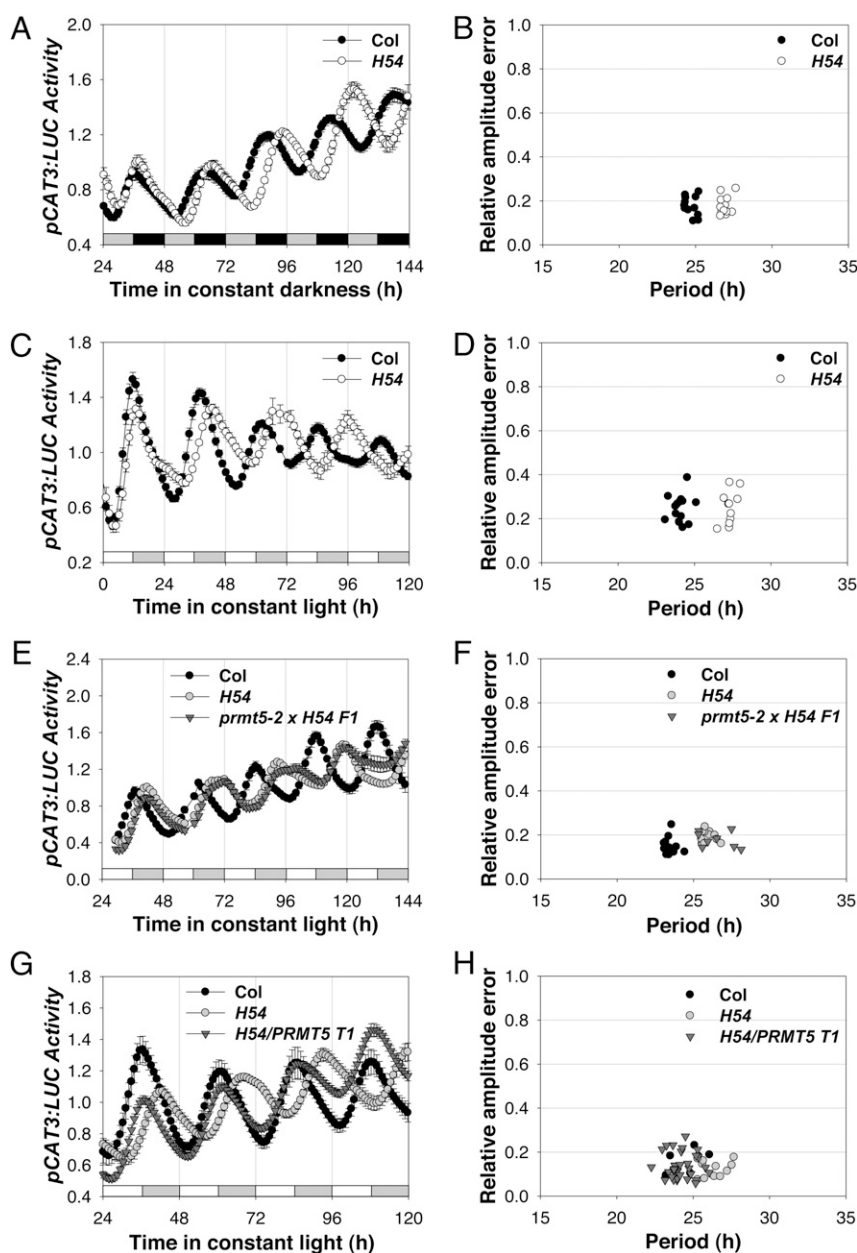


Fig. 1. *prmt5* mutations lengthen the period of *pCAT3:LUC* in DD and LL. Seedlings of the indicated genotypes containing were entrained to photoperiods (LD: 12/12 h) for 7 d before release into DD (A and B) or LL (C–H). Average traces of *pCAT3:LUC* expression (mean \pm SEM; $n = 11$ –13 in A–D; $n = 8$ –36 in E–H) and period versus relative amplitude error plots show the long period of H54 (A–D), the allelism of H54 and *prmt5-2* (E and F), and the rescue of the long period of H54 by *PRMT5* (G and H). The gray and black bars in A and the white and gray bars in C, E, and G indicate subjective day and night, respectively.

circadian phenotypes. In *Arabidopsis*, high-throughput short-read sequencing has identified spontaneous and chemically induced mutations in the reference (34, 35) and nonreference genomes (36). Accordingly, we first resequenced the Col-3 parent used for the mutagenesis to an average base coverage of 25-fold (122 million reads of 35 bp were mapped) and identified ~6,000 heterozygous and homozygous SNPs relative to the Col-0 sequence (Fig. S2B and C and Table S1). We then resequenced a pool of F2 progeny from a backcross of H54 to Col-3 that were long period and, hence, homozygous for the H54 mutation. Thirty-two homozygous SNPs were clustered in the region of chromosome 4 to which H54 mapped; 15 of these SNPs were intergenic, and 17 mapped to annotated genes, although one was in an intron. Two intragenic and one intergenic SNPs also were seen in Col-3 and therefore were unlikely to be responsible for the H54 mutant long-period phenotype. Eight of the intragenic SNPs were predicted to alter the coding sequence of eight distinct genes (Table S1). One of these, a G642A transition that is consistent with EMS mutagenesis, is predicted to confer the nonsense mutation W214* and to result in a truncated form of *PRMT5* (At4g31120).

Two aspects of the H54 phenotype are consistent with published reports of *prmt5* mutants: H54 has dark green, curled leaves and flowers late (Fig. S3A) (27–29). However, clock defects had not yet been associated with loss of *PRMT5* function. We therefore obtained additional *prmt5* alleles and tested for defective clock function. Consistent with our observations with H54, *prmt5-2* showed long period in cotyledon movement: Col-3 seedlings had a period of 23.4 ± 0.2 h ($n = 10$), whereas *prmt5-2* seedlings had a period of 26.4 ± 0.6 h ($n = 5$; $P < 0.01$) (Fig. S1C and D).

To confirm that the H54 mutant phenotype results from the W214* nonsense mutation in *PRMT5*, we crossed H54 to *prmt5-2* and observed that the F1 progeny all showed a long period in *pCAT3:LUC* expression. In LL, Col-3 seedlings had a period of 23.5 ± 0.1 h ($n = 12$), whereas H54 seedlings had a period of 25.9 ± 0.1 h ($n = 12$; $P < 0.0001$), and F1 seedlings also exhibited a long period of 26.5 ± 0.4 h ($n = 8$; $P < 0.0001$) (Fig. 1E and F). In addition, the F1 progeny of a cross of H54 to *prmt5-2* were all late flowering (Fig. S3A), indicating allelism. Finally, we introduced a wild-type genomic copy of *PRMT5* driven from its endogenous promoter into H54 and observed full rescue of the wild-type period in *pCAT3:LUC* expression. In LL, Col-3 seedlings had a period of 24.3 ± 0.3 h ($n = 9$), and T1 transgenic lines also exhibited a wild-type period of 24.2 ± 0.1 h ($n = 36$; $P = 0.7$), whereas H54 seedlings had a period of 26.3 ± 0.1 h ($n = 11$; $P < 0.001$) (Fig. 1G and H). We tested the expression of *PRMT5* in the H54 background by quantitative real-time PCR (qPCR); expression was significantly decreased (Fig. S4A). Accordingly, we conclude that H54 is a loss-of-function allele of *PRMT5* that we name “*prmt5-54*.”

***prmt5* Mutants Are Impaired in Light Inhibition of Hypocotyl Elongation.** Disruption of circadian clock function has many manifestations. Many clock mutants show defects in light inhibition of hypocotyl elongation, and we observed that *prmt5-54* and *prmt5-2* show long hypocotyls relative to their respective wild-type parents in blue but not in red light (Fig. S3B and C).

***PRMT5* Is Broadly Expressed, and *PRMT5* Protein Is Found in both the Nucleus and Cytoplasm.** We used the *PRMT5* promoter to drive β -glucuronidase (GUS) expression and observed widespread expression throughout the shoot and root (Fig. S4B–G). To address the subcellular localization of *PRMT5*, we fused GFP in frame to the carboxyl terminus of *PRMT5* (driven by the cassava vein mosaic virus promoter) and observed *PRMT5*-GFP accumulation in both nucleus and cytoplasm of transfected protoplasts (Fig. S4H–J).

***PRMT5* Expression Is Regulated by the Circadian Clock and Responds to both Light and Temperature Cues.** To establish whether *PRMT5* mRNA abundance is circadian and/or diurnally regulated, we used the publicly available DIURNAL (<http://diurnal.cgrb.oregonstate.edu/>) database of *Arabidopsis* microarrays sampling a range of time

courses (37–40). *PRMT5* mRNA abundance exhibits daily oscillations during growth in photocycles (LD), thermocycles (hot/cold, HC), and during growth in constant conditions following entrainment to either LD or HC cycles, indicating control by the circadian clock (Fig. 2A–C). In plants entrained to photocycles, *PRMT5* is phased to dusk (~8–12 h after dawn or subjective dawn) during driven conditions of long (LD: 16/8 h), intermediate (LD: 12/12 h), or short (LD: 8/16 h) days as well as during free-run conditions of constant light and temperature following entrainment to photocycles (Fig. 2A and C). Consistent with this observation, qPCR analysis using seedlings entrained to photocycles (LD: 12/12 h) before release into LL shows a dusk-specific peak of *PRMT5* mRNA accumulation (Fig. S4A). In contrast, the expression of *PRMT5* is phased to late evening/early dawn (~20 h after dawn or subjective dawn) during driven conditions of thermocycles or during free-run conditions following entrainment to thermocycles (Fig. 2B and C). This altered phase in response to thermocycles is seen both with and without concomitant exposure to photocycles (Fig. 2B) and indicates that *PRMT5* is sensitive to both light and temperature but is preferentially responsive to temperature entrainment.

To determine whether the circadian clock regulates transcription of *PRMT5*, we placed *LUC* under the control of the *PRMT5* promoter (*pPRMT5:LUC*) and observed robust circadian oscillations in *LUC* activity (Fig. 2D). *LUC* activity is minimal before dusk and then increases throughout the subjective night to peak late in the subjective night, before declining before subjective dawn. This expression pattern is seen in seedlings in constant conditions following entrainment to either photocycles or thermocycles and is consistent with the peak in mRNA abundance seen following entrainment to thermocycles (Fig. 2B and C) but contrasts with the peak in mRNA abundance seen at dusk following entrainment to photocycles (Fig. 2A and C). Although our data are inadequate to explain this apparent discrepancy, we speculate that changes in mRNA stability may contribute to the differences between transcription rates as measured with *pPRMT5:LUC* and transcript accumulation. The stability of a number of transcripts has been shown to be under circadian control (41), and the stability of others, including *CCA1* and *LHY*, is light regulated (11). The stabilization of *CAT3* mRNA in extended dark obscures an underlying rhythm in *CAT3* transcription (24, 42). Perhaps elevated temperature during the subjective night destabilizes *PRMT5* mRNA, preventing the continued accumulation of *PRMT5* mRNA at night in the absence of thermocycles. It also is possible that our *pPRMT5:LUC* construct lacks some sequence(s), presumably downstream of the transcription start site, necessary to recapitulate fully the transcriptional regulation of the endogenous *PRMT5* gene.

Using a large set of circadian and diel expression experiments (<http://diurnal.cgrb.oregonstate.edu/>), we established a list of 1,253 genes (Pearson correlation coefficient = 0.9) that share the same expression profile as *PRMT5* under cycling conditions (Table S2). These coexpressed genes are enriched for genes associated with ribosomal structural molecules, cytosol, cell organization and biogenesis, mitochondria, intracellular components, nucleus, cytoplasmic components, and DNA/RNA binding (Fig. S5). Using the PHASER tool (<http://phaser.cgrb.oregonstate.edu/>), we asked at what time of day expression of the top 500 genes peaks. Like *PRMT5*, this set of coreregulated genes peaks before dawn after entrainment to thermocycles or to concurrent thermocycles and photocycles, but in the absence of entrainment to thermocycles (entrainment to photocycles alone), these genes exhibit altered phase and peak between midday and dusk (Fig. S6). These results support the notion that *PRMT5* is part of an environmentally controlled transcription module.

Loss of *PRMT5* Function Affects both the Period and Strength of Expression of Multiple Clock Genes. To determine if the effect of loss of *PRMT5* function on period length results from the misexpression of specific clock genes, we measured the mRNA abundance of a number of clock and clock-regulated transcripts by

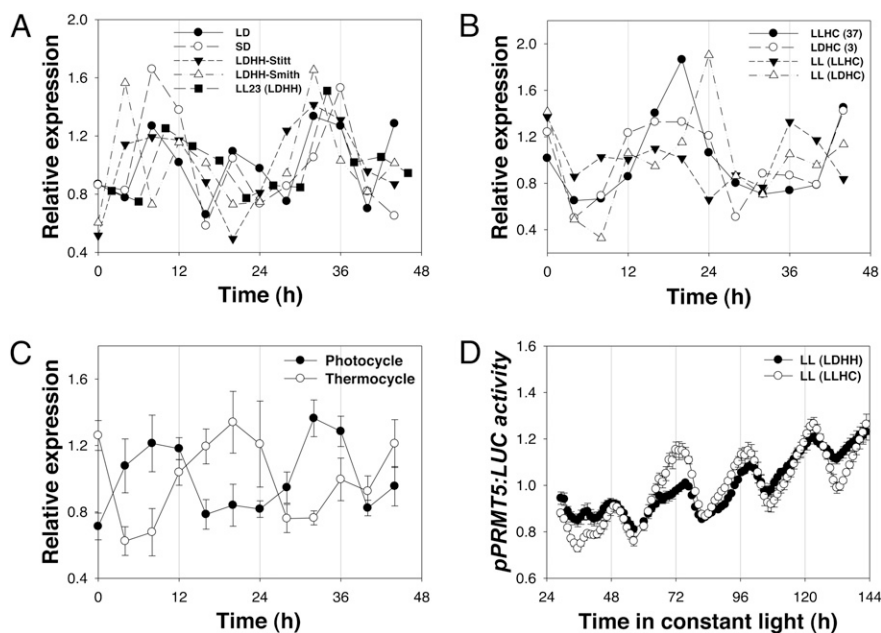


Fig. 2. *PRMT5* transcript abundance and transcription is clock regulated and sensitive to both light and temperature cues, but *PRMT5* phase is preferentially responsive to temperature entrainment. (A–C) *PRMT5* mRNA accumulates maximally in the late afternoon during photocycles and constant temperature or following entrainment to photocycles alone (A and C) but accumulates maximally in the subjective night during thermocycles or following entrainment to thermocycles or to both thermo- and photocycles (B and C). Data are from DIURNAL (<http://diurnal.cgrb.oregonstate.edu/>) and for each experiment are normalized to the average value to facilitate comparison of the temporal patterns despite differences in absolute signal strengths. LD, 16 h light/8 h dark; SD, 8 h light/6 h dark; LDHH-Stitt, 12 h light/12 h dark; LDHH-Smith, 12 h light/12 h dark; LLHC, constant light, 12 h at 22 °C/12 h at 12 °C; LDHC, 12 h in light at 22 °C/12 h in dark at 12 °C; LL (LLHC), constant conditions following entrainment to LLHC; LL (LDHC), constant conditions following entrainment to LDHC. (C) The data of A and B are averaged and plotted as mean \pm SEM. (D) *pPRMT5:LUC* expression (mean \pm SEM; $n = 25$) in seedlings entrained to photocycles (LL (LDHH)) or to thermocycles (LL (LLHC)) for 7 d before release into constant light and temperature.

qPCR and with *promoter:LUC* fusions. In each case the period was lengthened relative to wild type (Fig. 3 and Fig. S1 E–L). The amplitude and waveform of expression was relatively unaffected for several clock genes, including *CCA1*, *LHY*, and *TOC1*. However, the amplitude of mRNA abundance rhythms was increased dramatically for two *PRR* genes (*PRR7* and *PRR9*) and for an evening-expressed clock gene, *GIGANTEA* (*GI*) (Fig. 3), suggesting that these genes may be targets of *PRMT5* activity.

Discussion

Our results demonstrate a crucial role for *PRMT5* in the *Arabidopsis* circadian clock. *PRMT5* encodes a type II protein arginine methyltransferase that catalyzes the symmetric dimethylation of arginine residues (Rsm2). A number of posttranslational modifications have been established as important elements in the plant circadian clock. For example, many clock components are phosphorylated, as has been seen in other taxa from cyanobacteria through humans (43, 44). A second posttranslational modification, ubiquitination, also is encountered often and typically targets a protein for proteasomal degradation (14). Phosphorylation often serves as a prelude to ubiquitination and degradation. However, phosphorylation also can alter protein–protein and protein–DNA interactions as well as protein activity without concomitant effects on protein stability. Posttranslational decoration with *N*-acetylglucosamine also plays a significant role in *Arabidopsis* clock function, because mutational loss of or overexpression of the O-linked β -*N*-acetylglucosamine transferase, Spindly (SPY), lengthens or shortens, respectively, the circadian period (45). The period lengthening associated with mutational disruption of *prmt5* indicates that Rsm2 represents a decoration important for the functioning of the plant clock and possibly for clocks in general. In general terms, the methylation of arginine residues in proteins can be expected to alter binding interactions, because methylation alters both the shape and hydrogen-bonding potential of the modified arginine (46).

At this time we do not know the key targets for *PRMT5*-mediated methylation that are of functional significance to the circadian clock, although several can be proposed. *PRMT5* methylates components of the transcription complex, including the transcription elongation factor SPT5, altering its interaction with RNA polymerase II and potentially affecting global transcription rates (47). *PRMT5* also interacts with Switch/Sucrose non-

fermentable (SWI/SNF) chromatin remodeling complexes to act as a transcriptional coactivator (46). *PRMT5* is a transcriptional corepressor that methylates R8 of histone H3 and R3 of histone H4, modifications that repress gene expression (46, 47). In this role, *PRMT5* can be targeted to specific genes as well as exerting more ubiquitous effects. In *Arabidopsis*, *PRMT5* has been shown to methylate R3 of histone H4 (H4R3me2) at the Flowering locus C (*FLC*) promoter, which decreases expression of *FLC*, a flowering repressor, to promote flowering (27–29). Loss of *PRMT5* function elevates *FLC* expression and therefore would lengthen period, which lengthens in response to increased *FLC* expression (39, 48, 49). However, the period lengthening seen in *prmt5* mutants is greater than might be expected based on increased *FLC* expression, suggesting additional consequences of loss of *PRMT5* function. Thus, if the period lengthening associated with loss of *PRMT5* function is mediated entirely through histone modification and concomitant changes in gene expression, there are likely to be expression changes for genes in addition to *FLC*.

In addition to methylating histones, *PRMT5* methylates Sm proteins, constituents of small nuclear ribonucleoprotein (snRNP) components of the spliceosome, and inhibition of methylation disrupts premRNA splicing (47). It has been shown recently that histone modifications, including methylation of lysine residues in H3, regulate alternative splicing of a number of human genes (50). This observation may be of relevance to the period effects of *prmt5* mutations in *Arabidopsis*. *CCA1* premRNA undergoes alternative splicing which alters the balance between productive and unproductive mRNAs that cannot be translated to give full-length protein (51). There are alternatively spliced isoforms of other clock gene mRNAs, including at least two alternatively spliced isoforms of *PRR9* mRNA. Thus, the period-lengthening effects of the loss of *PRMT5* function could stem, at least in part, from either global or targeted changes in splicing.

Among the genes with altered expression in *prmt5* mutants, the morning loop components *PRR9* and *PRR7* and the evening loop component *GI* show increased amplitude as well as long period and are potential direct or indirect targets of *PRMT5* activity. In contrast, central loop components *CCA1*, *LHY*, and *TOC1* are relatively unaffected in terms of amplitude and exhibit only long period. Together with the sensitivity of *PRMT5* phase to entrainment by photocycles versus thermocycles, this finding suggests that *PRMT5* may be an important conduit of environmental

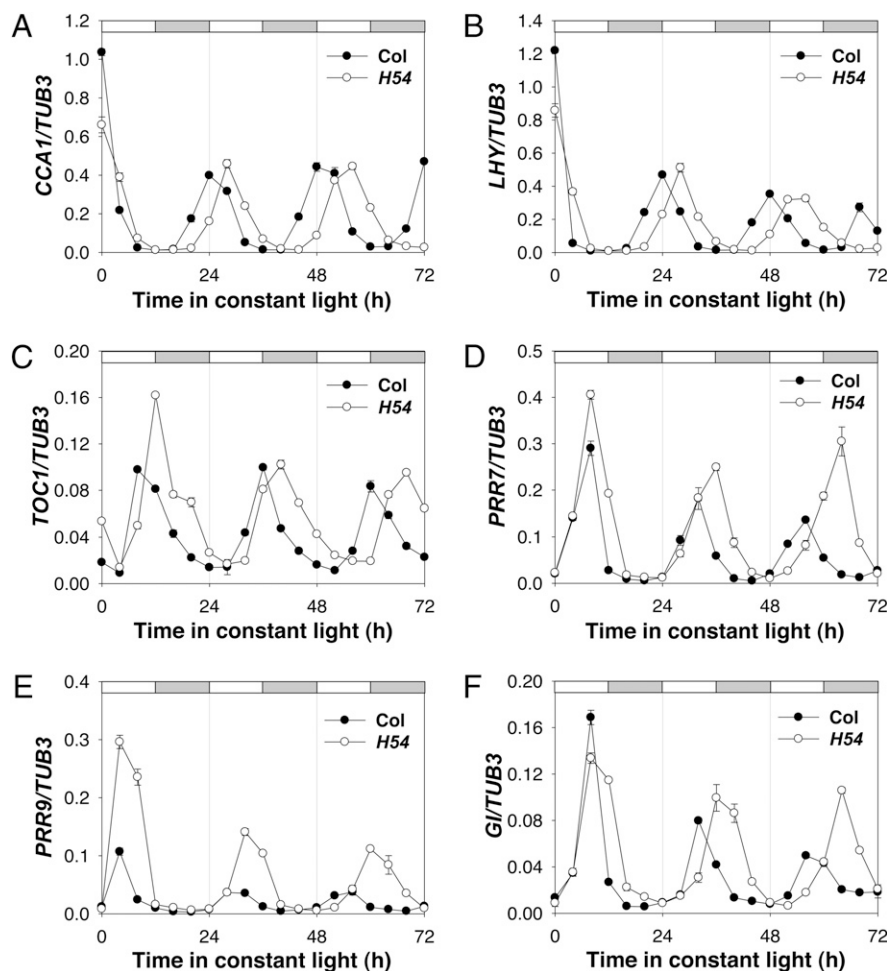


Fig. 3. *prmt5* mutation affects the period of all and the amplitude of expression of a subset of clock genes. Col and *prmt5-54* seedlings were entrained to photoperiods (LD: 12/12 h) for 10 d before release into LL. Transcript levels (mean \pm SEM from two independent experiments) of (A) CCA1, (B) LHY, (C) TOC1, (D) PRR7, (E) PRR9, and (F) GI were estimated by qPCR and normalized to tubulin (*TUB3*) expression. White and gray bars indicate subjective day and night, respectively.

input, via the morning and evening loops, to modulate clock entrainment and performance. An exciting possibility is that PRMT5 exerts this effect through environmentally induced epigenetic genome decorations.

Materials and Methods

Bioluminescence and Cotyledon Movement Assays. Rhythm assays were performed as described (52), except that LUC measurement in the dark was recorded with a Hamamatsu digital CCD camera (C4742-98 ERG; Hamamatsu Photonics) using MetaMorph software. Seedlings were entrained in photoperiods (LD: 12/12 h) for 6 (cotyledon movement) or 7 (LUC) d. Rhythms were analyzed by fast-Fourier transform nonlinear least-squares (53).

Generation of Constructs and Transgenic Plants. For complementation of H54, ~5.6 kb of the *PRMT5* gene including 1 kb of promoter was amplified by PCR from genomic DNA using gene-specific primers (Table S3), cloned into pCR8/GW/TOPO (Invitrogen), and recombined into pEarleyGate 301 (54). For *pPRMT5:GUS* or *pPRMT5:LUC* lines, ~1.1 kb of the *PRMT5* promoter was amplified from genomic DNA using gene-specific primers (Table S3), cloned into pCR8/GW/TOPO (Invitrogen), and recombined into pMDC 163 (55) or subcloned into pZP2LUC+ (56). The resulting binary vectors were introduced into *Agrobacterium tumefaciens* strain AGL1 by electroporation and transformed into Col (52).

Expression Analysis by Quantitative RT-PCR. Seedlings were entrained for 10 d in photoperiods (LD: 12/12 h) and transferred to LL. Samples were collected every 4 h for the following 3 d. RNA was extracted using Qiagen RNeasy Plant Mini Kit (Qiagen). First-strand cDNA synthesis used 2 μ g of total RNA with the SuperScript III first-strand synthesis system (Invitrogen). The cDNA was diluted 10 \times with water, and 1 μ L was used for PCR amplification using a SYBR Premix Ex Taq II (Takara) with gene-specific primers (Table S3). mRNA abundances

were calculated using the comparative C_T method, with *TUB3* (At5g62700) as the normalization control.

Plant Growth and DNA Preparation for Sequencing. H54 and its parent (Columbia, Col-3) were grown under intermediate-length days (LD: 12/12 h) and constant temperature (22 $^{\circ}$ C) for 7–12 d on 1/2 strength Murashige and Skoog plates without sucrose. Whole plants were harvested, and nuclear DNA was extracted using a modified protocol (57). Briefly, groups of 20–50 seedlings were frozen in liquid nitrogen, homogenized in a Retsch mixer (MM301), and filtered through cheesecloth, and nuclei isolated using a sucrose gradient were used for DNA extraction. SOLiD fragment libraries were prepared according to manufacturer's protocol (Life Technologies).

SOLiD Sequence by Ligation Sequencing. Sequencing template beads were generated by emulsion PCR, affixed to a microscope slide, and sequenced by oligonucleotide ligation and detection according to the SOLiD manufacturer's instructions (Life Technologies). We generated 35-bp reads for Col-3 and 50-bp reads for H54. Primary images were processed, color bases were called, and colorspace quality values were generated on the SOLiD onboard cluster. Reads were rejected if any colorbase could not be called in a read; usually this problem resulted from poor quality toward the end of the read or beads that were out of focus during a specific ligation cycle.

SOLiD Colorspace Alignment, SNP and INDEL Detection. Colorspace SOLiD reads (.csfasta) were mapped to the TAIR7 annotation of the *Arabidopsis* genome (arabidopsis.org). Raw colorspace reads that included both polyclonal reads and low quality were mapped (in colorspace) to the TAIR7 *Arabidopsis* genome using the corona lite pipeline (<http://solidsoftwaretools.com/>) allowing three mismatches (mm, -e 3) for 35-bp reads and 5 mm (-e 5) for 50-bp reads, counting adjacent errors as one error (-a 1), and aligning a maximum of 10 hits across the genome (-z 10). Both uniquely and randomly (more than one time in the genome) matching reads were used to call SNPs, thus leading to an

inflated number of heterozygous SNP calls. In colorspace every base is queried twice by two separate ligation reactions (two base pair encoding) and provides two independent supports for SNP calling. Thus in colorspace mapping, an SNP results in two adjacent mismatches with the reference genome; two colorspace changes are required for each SNP. To maximize the number of reads that map to the genome, adjacent errors are considered as one error. This approach allows the detection of up to one SNP and one error, two SNPs or two errors per read. To handle reads that map to multiple places in the genome (random), usually the result of repetitive sequences, we limited the number of times that these reads can map to 10.

To identify the causative SNP in H54, we used several filtering criteria. First we removed all SNPs that were identified in both Col-3 and H54 as compared with the TAIR7 reference. Second, we only used homozygous SNPs, which were mapped back to annotated genes in the TAIR7 assembly to determine if they changed amino acids.

Additional materials and methods are included in *SI Materials and Methods*.

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