Meiotically Stable Natural Epialleles of *Sadhu*, a Novel Arabidopsis Retroposon

Sanjida H. Rangwala¹, Rangasamy Elumalai^{2¤a}, Cheryl Vanier³, Hakan Ozkan^{2¤b}, David W. Galbraith², Eric J. Richards^{1*}

1 Department of Biology, Washington University in St. Louis, St. Louis, Missouri, United States of America, 2 Department of Plant Sciences, University of Arizona, Tucson, Arizona, United States of America, 3 Department of Biological Sciences, University of Nevada, Las Vegas, Nevada, United States of America

Epigenetic variation is a potential source of genomic and phenotypic variation among different individuals in a population, and among different varieties within a species. We used a two-tiered approach to identify naturally occurring epigenetic alleles in the flowering plant Arabidopsis: a primary screen for transcript level polymorphisms among three strains (Col, Cvi, Ler), followed by a secondary screen for epigenetic alleles. Here, we describe the identification of stable, meiotically transmissible epigenetic alleles that correspond to one member of a previously uncharacterized non-LTR retroposon family, which we have designated Sadhu. The pericentromeric At2g10410 element is highly expressed in strain Col, but silenced in Ler and 18 other strains surveyed. Transcription of this locus is inversely correlated with cytosine methylation and both the expression and DNA methylation states map in a Mendelian manner to stable cis-acting variation. The silent Ler allele can be converted by the epigenetic modifier mutation ddm1 to a meiotically stable expressing allele with an identical primary nucleotide sequence, demonstrating that the variation responsible for transcript level polymorphism among Arabidopsis strains is epigenetic. We extended our characterization of the Sadhu family members and show that different elements are subject to both genetic and epigenetic variation in natural populations. These findings support the view that an important component of natural variation in retroelements is epigenetic.

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Introduction

Epigenetic information in the form of differential DNA methylation, histone modification, and chromatin packaging is important for the management of large, complex eukaryotic genomes [1,2]. The stability of both animal and plant genomes depends heavily on epigenetic modification of repetitive DNA, including transposable elements and long tandem arrays of short repeats. For example, loss of genomic DNA methylation leads to meiotic defects [3], chromosome decondensation [4-7], transcription of previously quiescent transposons [8-11], and increased mutagenesis via DNA rearrangements [12]. Another component of genome stability is the integrity of epigenetic states that cement transcription rates of individual genes. These epigenetic states can be remarkably stable and transmitted faithfully through mitosis [13,14]. Alterations in these states form epigenetic alleles, or "epialleles," that lead to aberrant gene expression. The accumulation of epialleles in somatic tissues is now recognized as an important component of human carcinogenesis (e.g., tumor suppressor gene silencing) [15-17] and degenerative diseases associated with aging, such as atherosclerosis [18,19]. Transmission of epialleles is not restricted to mitotic divisions, but can also occur between generations of organisms [20-24], thereby mimicking traditional genetic mutations. This situation may be commonplace in plants where DNA methylation can be inherited through meiosis with high fidelity [25–28]. These findings raise the possibility that a significant portion of inherited information may be epigenetic and partially independent of the genetic sequence.

We are interested in determining the contribution of meiotically stable epigenetic alleles in the generation of genomic and phenotypic diversity. We exploited the availability of different accessions of the flowering plant *Arabidopsis thaliana* to evaluate the significance of the epigenetic component of inheritance in natural populations. We previously reported variation among Arabidopsis accessions in 5-methylcytosine (5mC) levels in the long tandem arrays of the major ribosomal RNA gene repeats [27]. Further, we showed that cytosine methylation patterns in inter-strain crosses are controlled by a combination of epigenetic inheritance of parental methylation patterns and the action of *trans*-acting loci [27,29]. Here, we describe a screen for natural epigenetic variation in cytosine methylation that is associated with transcript level polymorphisms among strains of Arabidopsis. This screen has led to the discovery of a new class of non-autonomous retroposons that are subject to epigenetic variation among natural accessions.

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Abbreviations: MPSS, massively parallel signature sequencing; RI, recombinant inbred; tpm, transcripts per million

* To whom correspondence should be addressed. E-mail: richards@biology.wustl.

 ${\tt \tiny ZA}$ Current address: Operon Biotechnologies, Huntsville, Alabama, United States of America

 ${\tt mb}$ Current address: Department of Field Crops, Faculty of Agriculture, University of Cukurova, Adana, Turkey



Synopsis

Differences among biological strains or individuals in a population can arise either from changes in DNA sequence (genetic) or in the packaging of DNA within the nucleus independent of DNA sequence (epigenetic). Both types of changes can alter gene activity, although epigenetic variation is often thought to be transient and unable to affect inherited differences among organisms. The authors compared the amount of RNA transcripts—a measure of gene activity—from a comprehensive set of genes among different strains of the flowering plant Arabidopsis. This approach led to the discovery of a novel family of DNA sequences, termed Sadhu, which show both genetic and epigenetic variation in gene activity. Alternative epigenetic states of one Sadhu element were created using mutants defective in epigenetic regulation. Both natural and induced epigenetic states were inherited. These results demonstrate that inherited differences among natural populations can be caused by epigenetic as well as genetic differences. Sadhu elements are a type of transposon, a class of DNA sequences that can move from one position in the genome to another. Epigenetic variation in gene activity of transposons modulates their movements within the genome and can influence genome diversification and evolution.

Results

Screen for Candidate Natural Epigenetic Variants

We set out to discover naturally occurring epialleles by identifying transcripts that were, first, differentially expressed in Arabidopsis accessions derived from wild populations and, second, whose transcription activity correlated with epigenetic state. An Arabidopsis long-oligo array containing approximately 26,000 predicted gene targets was hybridized with cDNA synthesized from whole-seedling RNA from the accessions Col, Ler, and Cvi. 279 loci were found to be differentially expressed (ANOVA p-values <0.1) with changes greater than 2-fold in pair-wise comparisons of these accessions. Here, we describe our characterization of one locus, At2g10410, identified in this screen for natural epialleles.

At2q10410 Is Subject to Natural Variation That Maps in cis

The microarray data indicated lower expression of At2g10410 in Ler and Cvi compared with Col. Robust transcription of At2g10410 in Col was confirmed by RT-PCR (Figure 1A, Table 1) and RNA gel blot analysis (Fig 1B, unpublished data); we did not detect expression of this locus in the Cvi and Ler accessions. Expression of the Col At2g10410 allele was corroborated by the massively parallel signature sequencing (MPSS) cDNA project (http://mpss.udel. edu/at) [30], which indicated a transcript level on the order of 180 transcripts per million (tpm). In addition, whole-genome transcriptome analysis in Col using a high-density oligonucleotide array [31] revealed that this locus is the most highly expressed feature within a 600-kb window of the transposonrich pericentromeric region of Chromosome 2. The lack of detectable At2g10410 expression in Cvi and Ler is not caused by the absence of this locus, which could be amplified from Ler and Cvi genomic DNA templates (Table 1). We also examined 22 additional Arabidopsis accessions and detected the presence of the At2g10410 locus in the majority of these natural strains (Table 1). However, of the accessions containing the locus, we detected expression by RT-PCR in only three (Col, N13, and Pu2-7). These data demonstrate the existence of natural variation in At2g10410 expression.

We examined a Col/Ler recombinant inbred (RI) population [32] to determine whether At2g10410 expression states mapped in cis or to a trans-acting transcriptional modifier. We selected fifteen RI lines that were Col homozygous and fifteen lines that were Ler homozygous at markers flanking the At2g10410 locus. RNA gel blot analysis indicated that all the lines containing the Col allele expressed At2g10410, while all the lines containing the Ler allele were silenced at this locus (Figure 1B). These results argue against an unlinked Col or Ler specific factor that influences expression at At2g10410. Instead, expression of this locus maps in cis and is stable through the eight generations of self-fertilization used to generate the recombinant inbred lines.

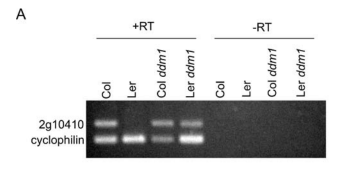
At2q10410 Is a Unique, Non-Coding Sequence That Arose by Retroposition

At2g10410 does not contain a long open reading frame and is not significantly similar to any known protein-coding sequence in any of the plant, animal, bacterial, or viral sequence collections. There is likewise no obvious sequence or structural similarity to any known non-coding RNA gene. At2g10410 is composed of 901 bp of unique sequence inserted within a hAT family DNA transposase pseudogene (Figure 2). EST data suggest a polyadenylated full-length transcript of 1,054 bp, ending 130 bp within the flanking transposase sequence. There is also a poly(A) stretch of ten nucleotides at the boundary of the hAT transposon and unique sequence of this element. In addition, the entire unique sequence is flanked by a direct duplication of 12 nucleotides of hAT sequence. The corresponding region in the Ra-0 accession (Table 1) contains a continuous hAT transposase pseudogene, lacking the At2g10410 unique sequence, the poly(A) tract, and the 12 nucleotide duplication (Figure 2). These genomic structure comparisons support a model in which the At2g10410 gene in the Col accession derived from a polyadenylated RNA precursor that has retrotransposed into the ancestral genomic sequence, represented by the Ra-0 allele.

Genetic and Epigenetic Variation of At2q10410

We explored the possibility that genetic differences between the expressed Col At2g10410 allele and the unexpressed Ler allele are responsible for transcription level differences. We determined that the nucleotide sequence of approximately 1.7 kb of Ler genomic DNA encompassing the At2g10410 locus was 98.6% identical to the Col reference genome sequence (Figure S1). The polymorphism frequency observed in the transcribed region was similar to that in the 5' and 3' non-transcribed flanking regions. No large indels or rearrangements were observed in our comparison of this region between the Col and Ler accessions and only two SNPs exist within +/- 150 bp of the transcription start site (Figure S1).

We next investigated cytosine methylation at At2g10410 using a PCR-based assay that monitors digestion of genomic templates by the methylation-requiring restriction enzyme McrBC [33]. In every natural accession we examined, transcriptionally silent alleles at this locus were also methylated (Table 1). By contrast, both the expressed N13 and Col alleles of At2g10410 are hypomethylated within the transcribed region. We observed one exception to the trend that methylated sequences are silent: the Pu2-7 At2g10410 allele



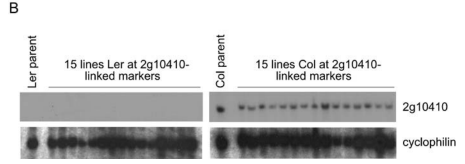


Figure 1. At2g10410 Is Differentially Expressed in the Arabidopsis Accessions Col and Ler
(A) RT-PCR of At2g10410 expression in Col, Ler, Col ddm1-1, and Ler ddm1-2. Cyclophilin is shown as an internal control for amplification.
(B) RNA gel blots of Col/Ler recombinant inbred lines. Lines homozygous for Col alleles at At2g10410-linked markers express the locus, while lines homozygous for Ler alleles at linked markers do not express At2g10410. All lanes were included on the same filter. The filter was rehybridized with a cyclophilin probe as a loading control.
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was both methylated and expressed (Table 1). We also examined cytosine methylation at At2g10410 in the Col/Ler RI lines described above. The At2g10410 locus was hypomethylated in all RI lines containing an expressed Col allele, while the locus was methylated in all lines carrying the silent Ler allele (unpublished data). Therefore, cytosine methylation was strictly correlated to the expression state of the Col and Ler allele. Moreover, these data suggest that the parental cytosine methylation state of the two alleles is stably inherited through the multiple generations required to construct the independent RI lines.

We also examined cytosine methylation flanking the At2g10410 transcribed region in Ler and Col accessions to determine the boundaries of the differential methylation states of these alleles. Alleles from both of these accessions had comparable methylation levels in the regions 1 kb upstream or 400 bp downstream of the transcript, even though they were differentially methylated within the gene (Figure 3A and 3B). These data indicate that the differential methylation that we observed between silenced and expressed accessions is limited to the region of transcription.

A higher resolution map of cytosine methylation of the *At2g10410* locus was constructed using bisulfite-mediated genomic sequencing [34] of a 380-bp region encompassing the start of transcription. In the Col allele a boundary of cytosine methylation coinciding with the transcription start site was observed; the region downstream of transcription was almost entirely free of methylation. On the other hand, the Ler allele was methylated both downstream and upstream

of transcription (Table 2; Figure S2). In the Ler allele, the methylation occupancy at CpG sites was high (\sim 90%), and less methylation was observed at cytosines at CpHpG (\sim 30%) or asymmetrical CpHpH sites (\sim 14%). These data corroborate the McrBC-PCR results and confirm that cytosine methylation is absent from the transcribed region in the expressed Col At2g10410 allele.

DNA Hypomethylation of the Ler *At2g10410* Locus Induces Ectopically Expressing and Meiotically Stable Epialleles

Having established that cytosine methylation correlates with the expression state of At2g10410 in different strains, we asked whether the silent Ler allele could be reactivated by manipulating DNA methylation of the locus. We first treated Ler seedlings with the cytosine-DNA-methyltransferase inhibitor 5-aza-deoxycytidine [35,36] and observed ectopic transcription of the At2g10410 locus (unpublished data). Next, we monitored expression of this locus in Ler strains carrying DNA hypomethylation mutations: met1-1 (disrupting the major Dnmt1-class CpG "maintenance" methyltransferase [37,38]) or ddm1-2 (disrupting a SNF2-class ATP-dependent nucleosome remodeling protein gene [28,39]). As shown in Figure 1A, loss of DDM1 function leads to ectopic expression of the Ler At2g10410 allele; similar results were observed for the met1-1 mutant (unpublished data). McrBC-PCR suggested that the expressing At2g10410 allele in the Ler ddm1-2 background is hypomethylated relative to the silenced allele in Ler wild-type (Figure 3C). Bisulfite-mediated

Table 1. Natural Variation of *At2g10410* Structure, Cytosine Methylation, and Expression

Accession	Source ^a	DNA ^b	5mC ^c	RNA ^d		
Br-0	CS22628	No				
Bur-0	CS22656	Yes	Yes	-		
C24	CS22620	Yes	Yes	-		
Col	Lehle WT-2	Yes	No	+++		
Cvi	Lehle WT-18	Yes	Yes	-		
Cvi-0	CS22614	Yes	Yes	-		
Ct-1	CS22639	Yes	Yes	-		
Fei-0	CS22645	Yes	Yes	-		
Hi-0	CS6736	No				
Kn-0	CS6762	Yes	Yes	-		
Kondara	CS22651	Yes	Yes	_		
Kz-1	CS22606	Yes	Yes	-		
Ler	Lehle WT-4	Yes	Yes	-		
N13	CS22491	Yes ^e	No	+++		
No-0	CS6805	Yes	Yes	-		
Po-0	CS6839	Yes	Yes	_		
Pro-0	CS22649	Yes	Yes	-		
Pu2-7	CS22592	Yes	Yes	++		
Ra-0	CS22632	No				
Tamm-27	CS22605	Yes	Yes	-		
Ts-1	CS22647	Yes	Yes	-		
Tsu-1	CS22641	Yes	Yes	-		
Van-0	CS22627	Yes	Yes	-		
Wei-0	CS22622	Yes	Yes	_		
Ws-2	CS22659	Yes	Yes	-		

^aLehle, Lehle Seeds; CS[number], Arabidopsis Biological Resource Center (ABRC) stock number.

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genomic sequencing of the At2g10410 locus in Ler ddm1-2 individuals confirmed complete loss of cytosine methylation in all sequence contexts (Table 2). RNA gel blots indicated that the ectopic At2g10410 transcript in Ler ddm1-2 plants is approximately the same size as the transcript in Col (unpublished data). We determined the DNA sequence from 245 bp upstream to 650 bp downstream of transcription in the Ler ddm1-2 mutant and found no differences from the Ler wild-type sequence (unpublished data). These findings indicate that the ddm1-2 mutation did not alter the genetic information at the Ler At2g10410 locus, but did change the DNA methylation and transcription states of the locus.

To see whether ddm1-induced expression of the Ler At2g10410 allele was stable in the presence of a functional DDM1 allele, we outcrossed a Ler ddm1-2 individual to wild-type Col. F1 hybrids generated from reciprocal crosses maintain expression of both the Col and Ler alleles (Figure 4). By contrast, there was no expression of the Ler allele in F1 hybrids of a control cross between wild-type Col and Ler individuals. Therefore, parental expression states at At2g10410 are faithfully maintained in an inter-strain cross, and there is no evidence for a Col or Ler specific trans-acting modifier. When F1 $At2g10410^{\text{CollLer}}$ hybrids were backcrossed to the Col parent strain (\rightarrow BC1), individuals heterozygous for



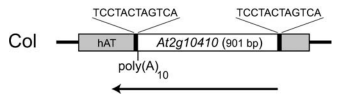


Figure 2. Structure of At2g10410 in Two Accessions

At2g10410 in Col is inserted within a sequence that is 93% identical to a hAT10 (1536 to 2399 coordinates) DNA transposase gene (grey box, hAT). The poly(A) tract and target site duplication in Col are shown. The At2g10410 transcript is indicated by the arrow. The Ra-0 accession lacks the At2g10410 insertion, poly(A) tract, and target site duplication. DOI: 10.1371/journal.pgen.0020036.g002

At2g10410 alleles from both Col and Ler parents continued to maintain the expression states inherited from the original parents. Ten Col \times [Col \times Ler ddm1-2] BC1 individuals examined showed bi-allelic expression, while five Col×[Col× Ler wild-type] BC1 individuals examined showed expression of the Col allele only (Figure 4). We note that $Col \times [Col \times Ler]$ ddm1-2] BC1 individuals must be either heterozygous or homozygous wild-type at the DDM1 locus. We conclude that ectopic expression of the Ler allele can be maintained in the absence of a homozygous ddm1-2 mutation. Moreover, we examined two F2 individuals from the Col \times Ler ddm1-2 cross that were homozygous wild-type at the DDM1 locus and homozygous Ler at the At2g10410 locus. Both of these F2 individuals, which no longer carried the ddm1-2 mutation or the Col At2g10410 allele, persisted in their expression of the Ler At2g10410 allele (Figure 4). These data indicate that expression states at At2g10410 can be modified in a ddm1 mutant background and that once established, ectopic expression states can be inherited as meiotically stable epialleles. Taken together with the RI mapping results detailed above, we conclude that silenced or active expression states at At2g10410 behave as stable epialleles.

At2g10410 Is a Member of a Previously Uncharacterized Non-Autonomous Retroposon Family

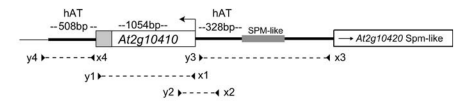
After establishing that At2g10410 was a novel retroposon subject to natural epigenetic variation, we searched the available Arabidopsis genomic sequence from strain Col for related sequences. Fourteen other sequences in the Col genome share 55%–75% identity over the \sim 850–900 bp of At2g10410 unique sequence (Table 3; Figure 5A). Consistent with being generated through retroposition, 13 of the 14 homologs contain 3' poly(A) tracts, while eight feature recognizable target site duplications of at least ten nucleotides (Table 3). The 3' target site duplication always occurs adjacent to the poly(A) tract; however, the other target site duplication occurs anywhere between 8 bp and 75 bp 5' of the conserved sequence. This observation suggests that the 5' boundary of retroposition can vary in both length and sequence. Apart from conservation of genomic structure,

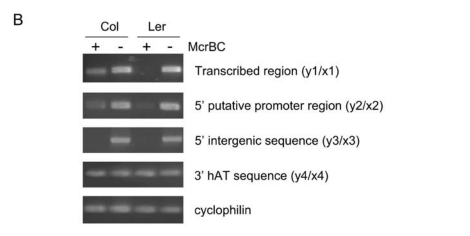
^bGenomic DNA amplification using specific primers (X1 and R1, Table S2) within the locus. ^cCytosine methylation was determined by the McrBC PCR assay shown in Figure 3.

^dSteady-state At2g10410 transcript levels were assayed by RT-PCR. —, none detected; +, low; ++, moderate; +++, high.

^eWe could amplify a full-length copy of the element from this accession using primers located inside the element, but could not amplify the full-length element using primers located in the flanking region. A full-length element is present in the N13 accession, but is located in a different genomic region than in the Col accession.

Α





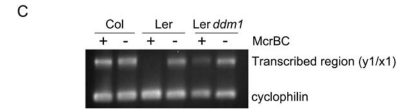


Figure 3. Analysis of Cytosine Methylation at At2g10410 by McrBC Digestion followed by PCR

(A) Diagram of the At2g10410 locus in Col including nearby repetitive elements and intergenic regions. Arrow indicates start and direction of transcription. The positions of primers used for PCR analysis are indicated as triangles.

(B) PCR amplification of genomic DNA +/- McrBC digestion. Cyclophilin is shown as an amplification control.

(C) McrBC PCR of the At2g10410-transcribed region in Col, Ler, and Ler ddm1-2 genomic DNA samples. Cyclophilin is shown as an internal amplification control.

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there are several short stretches of high DNA sequence identity among the 14 homologs separated by areas of little or no similarity (Figure 5B). Of note, there is a 13 nucleotide sequence (consensus 5'-GGACAATCGTTCC-3') near the

start of *At2g10410* transcription that is followed by a 10–20 nt CT-rich region (Figure 5B and 5C). DNA sequence conservation is restricted to the unique transcribed sequence of *At2g10410*; there is no similarity among family members in

Table 2. Number of Cytosines Methylated at the At2g10410 Locus in Col, Ler, and Ler ddm1-2 Plants

Strand	Number of Sites	CpG			СрНрС			СрНрН			
		Col	Ler	Ler ddm1	Col	Ler	Ler ddm1	Col	Ler	Ler ddm1	
Тор	Assayed Methylated (%)	192 0 (0)	238 226 (95.0)	168 0 (0)	176 4 (2.3)	170 56 (32.9)	120 0 (0)	912 38 (4.2)	918 198 (21.6)	648 0 (0)	
Bottom	Assayed Methylated (%)	121 0 (0)	132 113 (85.6)	144 0 (0)	77 12 (15.6)	66 18 (27.3)	72 1 (1.4)	583 41 (7.0)	594 16 (2.7)	648	
Combined	Assayed Methylated (%)	313 0 (0)	370 339 (91.6)	312 0 (0)	253 16 (6.3)	236 74 (31.4)	192 1 (0)	1495 79 (5.3)	1512 214 (14.2)	1296 2 (0)	

Determined by bisulfite-mediated genomic sequencing of top strand (-105 to +276) and bottom strand (-71 to +221) amplicons (see Materials and Methods). DOI: 10.1371/journal.pgen.0020036.t002



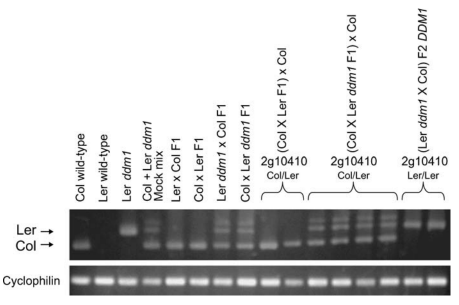


Figure 4. RT-PCR/CAPS (Cleaved Amplified Polymorphic Sequence) Detection of Allele-Specific Expression at At2g10410

The Col allele is cleaved with BstB1, generating a 480-bp fragment plus an undetected 70-bp fragment, while the Ler allele is uncleaved (550 bp). At2g10410 is not expressed in Ler wild-type, but is expressed in wild-type Col and Ler ddm1 mutant, as shown in Figure 1. A mixture of cDNA templates from Col and Ler ddm1 samples was used to illustrate the detection of bi-allelic expression; note an additional higher molecular weight heteroduplex band (see Materials and Methods) in samples showing bi-allelic expression. A total of five Col \times Ler heterozygous At2g10410 BC1 and ten Col \times Ler ddm1-2 heterozygous BC1 individuals were examined; all looked identical to the representative individuals shown. In addition, we examined two $DDM1^{+/+}$ F2 individuals homozygous for the Ler allele at At2g10410 that resulted from self-pollination of a Ler $ddm1 \times$ Col F1 individual. RT-PCR amplification of cyclophilin transcripts is shown as a control. DOI: 10.1371/journal.pgen.0020036.g004

the immediate upstream or downstream flanking genomic regions. These features suggest that each member has retroposed independently into its flanking genomic region without obvious selection for a particular region. Because none of these homologs contains an ORF with similarity to a transposase-related protein, these sequences fit the criteria of a family of previously uncharacterized non-autonomous

retroposons. We have named these sequences *Sadhu* elements, after the Sanskrit term for ascetic holy men who have renounced society.

In addition to the 14 family members described in Table 3, there are 25 sequences of $\sim 175-750$ bp in the Arabidopsis Col genome that have similarity to the 5', 3', or an internal section of the full-length *Sadhu* elements (Table S1). We noted

Table 3. Features of 14 Full-Length Members of the Sadhu Element Family in the Col Genome

Gene ID	Length (bp) ^a	Target Site Duplication (bp)	3'Poly(A) Tract	Transposable Elements within 2 kb	Incorporated into an Annotated Protein-Coding Gene	Largest ORF ^b (Amino Acids)	Expressed in Col (RT-PCR)
At1g03420	870	Yes, 13	Yes	No	Yes	171	Yes
At1q30835	902	Yes, 14	Yes	No	Yes	132 (AS)	Yes
At1g35112	876	No	Yes	Yes	Yes	66	Yes
At1q44935	846	No	No	Yes	Yes	54 (AS)	No
At1q50735	880	Yes, 13	Yes	Yes	No	87 (AS)	No
At2q10410	901	Yes, 12	Yes	Yes	No	39	Yes
At3q13438	886	Yes, 12	Yes	No	No	54 ^e	Yes
At3g02515	864	Yes, 12	Yes	No	No	55	No
At3g31442	870	Yes, 10	Yes	Yes	No	19	No
At3g42658	873	No	Yes	Yes	No	47	Yes
At3g44042	885	No	Yes	Yes	No	45	No
At4g01525	872	Yes, 11	Yes	Yes	Yes	56	Yes
At5g28626 ^c	932 ^d	No	Yes	Yes	No	59	Yes
At5g28913	908	No	Yes	Yes	No	58	Yes

^aExtent of similarity to At2g10410, not including the 3' poly(A) tract or 3' read-through of full-length transcript into the flanking genomic area.

^eEST BP867836 RAFL21 corresponds to a spliced transcript of the element that encodes a putative ORF of 92 amino acids. DOI: 10.1371/journal.pgen.0020036.t003



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^bEncoded by the element in isolation, not including incorporation into a larger predicted ORF. AS, antisense.

^cPreviously described as orphan transcript At_oRNA_590 and At_oRNA_478 [55].

dNot including the 760 bp ATLANTYS2-like LTR insertion.

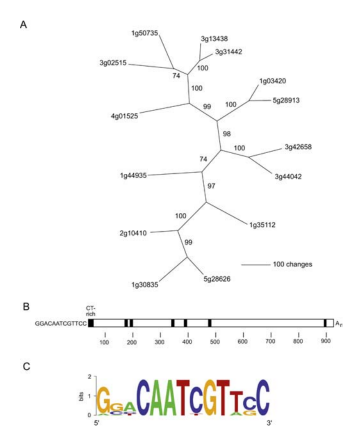


Figure 5. *Sadhu* Retroposon Family Members in the Arabidopsis Col Genome

(A) An unrooted phylogram of 14 full-length *Sadhu* elements built using a maximum parsimony algorithm. Bootstrap values are shown (100 bootstrap replicates).

(B) Diagram of consensus sequence structure of a full-length *Sadhu* element based on ClustalX gapped alignment. Black boxes indicate short blocks (10–20 nucleotides) of high sequence conservation, including a CT-rich block toward the 5' end. The majority of elements end with a poly(A) tract (Table 3).

(C) A conserved 13-bp sequence at the predicted 5' boundary of the element is illustrated as a logo diagram based on alignment of the 14 homologs shown in (A) [72].

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the presence of truncated 3' elements, some of which contain poly(A) tracts and target site duplications. Such structures are predicted to arise from reverse transcription that did not proceed to the 5' end of the transcript prior to transposition. Some of these truncated sequences are 99% identical to their closest full-length *Sadhu* element, with no flanking sequence similarity shared among these elements. These observations indicate that the closely related *Sadhu* elements did not arise by recent segmental duplication but represent independent recent retroposition events.

Ten of the 14 full-length *Sadhu* elements and 12 of the 25 truncated elements are located within the vicinity of other repetitive elements, such as transposons (Table 3, Table S1; see Figure S3 for chromosome distribution). Many of these are integrated within transposons; *At5g28626* is disrupted by an *ATLANTYS2*-like retrotransposon LTR sequence, while *At2g10410*, *At3g44042*, and *At3g31442* are embedded within DNA transposons. Despite this preference for integration near repetitive environments, transcription of nine of the 14 full-length *Sadhu* elements is detected in the Col accession by

RT-PCR (Table 3). In addition, although the *Sadhu* elements are by themselves non-coding, five homologs have been annotated within putative protein coding genes (Table 3). EST data confirm that one of these, *At1g30835*, is in fact transcribed in antisense orientation to other *Sadhu* elements and encodes a 132 amino acid predicted protein. Two additional family members encode ORFs greater than 75 amino acids. The amino acid sequences encoded by these ORFs are independent of one another and do not resemble known protein coding sequences. In summary, most of the full-length *Sadhu* elements in Arabidopsis strain Col are expressed, and some members are candidates for generating potentially functional gene products.

Several Other *Sadhu* Family Members Are Subject to Naturally Occurring Epigenetic Variation

We were interested in whether other members of the Sadhu retroposon family are, as with At2g10410 characterized above, subject to natural variation in epigenetic transcriptional regulation. We focused on five full-length Sadhu elements-At1g30835, At5g28626, At1g35112, At3g42658, and At3g44042, which are closely related to At2g10410 (Figure 5A). First, we screened a set of 25 accessions for the presence of that particular family member, as indicated by the ability to amplify these loci from genomic DNA templates (Table 4). Second, we evaluated gene expression of the loci verified to be present in the various accessions. Third, we monitored cytosine methylation status of the loci using the McrBC-PCR assay. There was considerable variation among the accessions for all three criteria at these five loci. For example, At1g35112 was expressed at low levels in a few accessions, but transcriptionally silent or not amplified from genomic DNA in others. Some accessions contained methylated alleles at this locus, while others contained unmethylated alleles. The two accessions with the highest level of expression of this locus, Kz1 and N13, were both unmethylated. This result suggests that cytosine methylation may play a part in suppressing expression of some alleles of At1g35112. More notably, silencing was correlated with DNA methylation for most naturally occurring alleles of the loci At5g28626, At3g42658, and At3g44042. For these three Sadhu elements, the majority of the accessions containing hypomethylated alleles expressed these genes, while most accessions that contained methylated alleles were silent at these loci. It is likely that many of the alleles at these elements represent naturally occurring epialleles, as with the case for the Col and Ler alleles at At2g10410.

Discussion

We sought out transcript-level polymorphisms in natural populations that behaved as meiotically stable epialleles. Here we describe a locus, At2g10410, which is differentially expressed in different accessions of Arabidopsis. Silenced alleles are methylated predominantly at CpG sites over the transcribed region, while expressed alleles are correlated with an absence of cytosine methylation. Transcript level differences between the robustly expressed Col At2g10410 allele and the silenced Ler allele map in cis in recombinant inbred lines. In addition, ddm1-induced ectopically expressed Ler alleles are meiotically stable upon introduction of the wild-type DDM1 allele. Therefore, differentially expressing states

Table 4. Natural Variation in Five Sadhu Elements among a Set of 25 Arabidopsis Accessions

Accession	At3g44042			At1g30835			At1g35112			At5g28626			At3g42658		
	DNA	5mC ^b	RNAc												
Br-0	Yes	Yes	+	Yes		++	Yes	Yes	+	Yes	No	+++	Yes	No	+++
Bur-0	Yes	Yes	-	Yes	No	+	Yes	No	-	Yes	No	+++	Yes	Yes	-
C24	Yes	No	+++	Yes		++	No			Small	No	+	Yes	No	+++
Col	Yes	Yes	-	Yes	No	++	Yes	No	+	Yes	No	++	Yes	Yes	+
Ct-1	No			Yes		++	Yes	No	_	Yes	No	++	Yes	No	+++
Cvi	Rearr	Yes	++	Yes	No	++	No			Small	No	+++	Yes	No	+++
Cvi-0	No			Yes	No	++	No			Yes	Yes	+	Yes	No	+++
Fei-0	Yes	Yes	_	Yes		++	Yes	Yes	+	Yes	No	+++	Yes	Yes	_
Hi-0	Yes	No	+++	Yes		++	Yes	Yes	-	Small	No	+++	Yes	No	+++
Kn-0	No			Yes		++	Yes	Yes	+	Small	No	+++	Yes	No	_
Kondara	Yes	Yes	_	Yes	No	_	Yes	No	+	Yes	No	+++	Yes	No	+++
Kz-1	No			No			Yes	No	++	Yes	No	+++	Yes	No	+++
Ler	Rearr	Yes	++	Small	No	_	No			Small	No	+++	Yes	No	+++
N13	Yes	Yes	_	Yes	No	+	Yes	No	++	Small	No	++	Yes	No	+++
No-0	No			Yes		++	No			Small	No	+++	Yes	Yes	_
Po-0	Yes	Yes	++	Yes		++	Yes	Yes	_	Yes	No	+++	Yes	Yes	-
Pro-0	Yes	Yes	_	Yes		++	No			Small	No	+++	Yes	No	+++
Pu2-7	Yes	Yes	_	Yes		++	Yes	No	_	Small	No	+++	Yes	No	_
Ra-0	Yes	No	+++	Yes		++	Yes	Yes	-	Yes	No	+++	Yes	No	+++
Tamm-27	Rearr		_	Yes		++	No			Yes	No	+++	Yes	No	+++
Ts-1	Rearr		_	Yes		++	Yes	No	-	Small	No	+++	Yes	Yes	_
Tsu-1	Yes	Yes	_	Yes		++	Yes	No	-	Small	No	+++	No		
Van-0	Yes	Yes	-	Yes		++	Yes	No	+	Yes	No	+	Yes	Yes	_
Wei-0	Yes	Yes	_	Yes		++	No			Yes	No	++	Yes	Yes	++
Ws-2	Yes	Yes	-	Yes	No	++	Yes	No	+	Small	No	+++	Yes	Yes	-

^aGenomic DNA amplification using gene-specific primers (see Table S2); Rearr (Rearranged), refers to cases where we could amplify a full-length copy of the element from that accession using primers located inside the element, but could not amplify the full-length element using primers located in the flanking region. We believe the full-length element is present in these accessions, but located in a different genomic context than in the Col genome. Small, a lower molecular weight PCR product was generated compared to Col.

of At2g10410 in different Arabidopsis strains behave as stable epialleles.

At2g10410 is a previously undescribed, unique, non-coding retroposed element. It is a member of a small family of such elements in Arabidopsis—Sadhu elements. These elements are typically ~900 bp long, with a poly(A) tract at the 3' end and direct target site duplications. Because Sadhu elements do not share any sequence similarity to any known ORFs, they are unlikely to be processed pseudogenes, but are more reminiscent of SINE-class retroposons. However, Sadhu elements differ from canonical SINEs in that they are longer (>500 bp) and do not have recognizable RNA pol III promoter A or B boxes nor similarity to known SINE or SINE ancestral molecules such as tRNA, 5S rRNA or 7SL RNA (i.e., mammalian Alu) [40,41]. Therefore, Sadhu elements represent a family of novel retroelements.

Sadhu elements, like SINE retroelements, do not encode their own reverse transcriptase. SINE elements are thought to make use of LINE-encoded reverse transcriptase/endonuclease to create a DNA copy from RNA intermediates, which then inserts into the genomic DNA [40]. We do not find any reverse transcriptase- or transposase-encoding sequences related to the Sadhu elements in the available Arabidopsis Col genome sequence. We hypothesize that the Sadhu elements may be mobilized by LINE-encoded factors. It is unclear how exactly the LINE retrotransposition machinery recognizes its targets of transposition. SINE elements maintain significant conservation of motifs with their non-coding

RNA ancestor molecules [40,42]. While *Sadhu* elements do not resemble SINEs or LINEs at the primary nucleotide level, they do show conservation of short motifs (Figure 5B) that may be functional in promoting mobilization. Although most *Sadhu* elements in the Col genome share only 60%–70% sequence identity, the presence of partial elements with 99% identity to one another (Table S1) suggests that mobilization is ongoing or has occurred recently in this family.

Mobilization of Sadhu elements by retroposition is expected to require expression of these elements into RNA intermediates. Indeed, most of the full-length Sadhu elements can be detected by RT-PCR in at least the Col strain (Table 3). Seven out of the 14 full-length Sadhu elements in Col are represented in the MPSS database at greater than 20 tpm in at least one tissue examined; At2g10410 is expressed at greater than 100 tpm in most tissues. By contrast, we examined 170 annotated retroelements in the Col MPSS database [30], and found that less than ten were expressed at more than 20 tpm in any tissue examined (unpublished data). If the robustly expressed At2g10410 is mobile or has been recently mobile, we would expect multiple copies of near identity in the genome. Preliminary analysis by Southern blot detects only one copy of this locus in the Col genome (unpublished data). This result suggests that transcription of At2g10410 is not sufficient for transposition. Perhaps a reverse transcriptase source necessary to mobilize Sadhu elements is itself either nonfunctional or silenced in the Col strain. DNA transposons are mobile in ddm1 mutants

^bCytosine methylation was determined by a McrBC PCR assay using gene-specific primers. Blank spaces indicate that the sample was not tested.

^cSteady-state transcript levels were assayed by RT-PCR. –, none detected; +, low; ++, moderate; +++; high. Blank spaces indicate that the sample was not tested. DOI: 10.1371/journal.pgen.0020036.t004

[9,10,43], and we are interested in exploring the possibility that LINE retrotransposition factors may become re-expressed in ddm1 or other chromatin mutants, indirectly causing increased mobility of Sadhu elements.

The robust expression of full-length Sadhu elements, in contrast to the general non-activity of other retroelements in Arabidopsis, is puzzling for another reason-from what promoter are these elements transcribed? The 14 full-length Sadhu elements contain no similarity outside of the transcribed region, suggesting that the elements do not carry their own upstream conserved promoter or enhancer elements. Because the conserved motifs among the Sadhu elements downstream of transcription (Figure 5B and 5C) do not bear any resemblance to known RNA pol III promoters, it is possible that these sequences may instead represent novel, non-canonical RNA pol II or RNA pol III promoter elements. An alternative model is that transcriptionally active Sadhu elements have inserted near cryptic RNA pol II promoters. Pol II transcripts are polyadenylated, while pol III transcripts are typically not. In fact, there are oligo-d(T)-primed ESTs to At2g10410 that do not originate from the poly(A) tract in the DNA sequence, supporting polyadenylation of this transcript. This evidence suggests that At2g10410, if not other Sadhu elements, may be transcribed from a flanking cryptic pol II promoter. We hypothesize that conserved downstream motifs in the Sadhu elements may act as enhancers to promote robust transcription of elements in a flexible variety of genomic contexts.

Our study of At2g10410 and preliminary survey of other Sadhu family members suggest that there is considerable genetic, DNA cytosine methylation, and transcriptional variation in these elements among A. thaliana accessions (Table 1, Table 4). Across the Sadhu family members examined, there is a good correlation between cytosine methylation and lack of transcription. However, some exceptions exist. In the case of unmethylated alleles that are not expressed, genetic variation in promoter elements may be responsible for transcriptional inactivity. In instances where methylated alleles are transcribed, the expression level tends to be intermediate, consistent with partial silencing. Previous studies have highlighted the role of cytosine methylation in silencing of DNA transposons and retrotransposons [9-11,44–48]). However, this is the first report of strain-specific variation in both transcript abundance and cytosine methylation of a retroposon family.

The Sadhu family of retroelements represents a previously overlooked source of genetic and epigenetic variation in the genome. Barbara McClintock proposed over one half century ago that transposons ("controlling elements") existing in different states or distinct genomic locations can differentially affect gene expression [49]. Recent studies have lent support to McClintock's view: epigenetic states at transposons can indeed affect the spread of transcription or silencing into neighboring coding sequences [50,51]. In some cases, chimeric transcription units are formed and regulated by flanking transposon sequence [52-54]. We found that At2g10410 hypomethylation in the expressed Col allele is limited to the region of transcription (Figure 3). In addition, preliminary results suggest that flanking transposons are not expressed in genetic backgrounds expressing At2g10410. Because At2g10410 is present in a transposon-rich heterochromatic pericentromere, expression at this locus may not

be adequate to reverse the silenced chromatin state of the genomic region. It is possible that expression or cytosine methylation at other Sadhu family members present in more euchromatic, gene-rich environments may influence the expression of neighboring genes.

Non-coding transcripts such as the Sadhu sequences have been discovered recently in a variety of organisms, from Drosophila to humans to Arabidopsis [55,56]. In some cases, these sequences are conserved in related species and may therefore be functional. There are no related sequences to the Sadhu sequences in any of the currently available plant genome sequence releases, suggesting that this family is rapidly evolving. In fact, there are variable numbers of given elements even among A. thaliana accessions (Table 1 and Table 4). We are currently searching for evidence of Sadhulike sequences in the genomes of other Brassicaceae species. Related sequences encoding autonomous retroelements, for instance, may provide clues to the origin of this nonautonomous retroposon family.

Finally, while Sadhu elements are by themselves nonprotein coding, at least one element, At1g30835, has been incorporated into a transcribed gene (expressed in antisense orientation to other family members) capable of encoding a 132-amino acid protein. Indeed, retroelement movement is thought to increase protein coding diversity, either through incorporation into new genes [57] or by shuffling of existing exonic sequences around the genome [58-60]. We believe that the Sadhu retroposons, in addition to being a reservoir of transcriptional variation, may serve as genetically important wells of novel genes and gene functions.

Materials and Methods

Plant materials. Col, Ler, and Cvi accessions were obtained by Lehle Seeds (Tucson, Arizona, United States); all other accessions were obtained from Arabidopsis Biological Resource Center (ABRC). Stock numbers are listed in Table 1. ddm1-1, ddm1-2, and met1-1 mutants were originally isolated in the Col strain and have been introgressed greater than five generations into Col or Ler genetic backgrounds [10,28,38]. Col/Ler recombinant inbred lines that had been self-fertilized eight generations [32] were obtained from ABRC. The fifteen lines used in this study that were homozygous Ler at markers linked to At2g10410 are CS1900, CS1911, CS1913, CS1929, CS1953, CS1954, CS1957, CS1959, CS1960, CS1968, CS1969, CS1970, CS1971, CS1974, and CS1988. The fifteen lines homozygous Col at markers linked to At2g10410 are CS1901, CS1903, CS1904, CS1909, CS1915, CS1916, CS1932, CS1945, CS1946, CS1948, CS1951, CS1963, CS1975, CS1978, and CS1984. The Col parent line is CS933, while the Ler parent line is CS20.

Plants were grown on soil or on 1× MS media with 1% sucrose. For 5-aza-dC treatment, seedlings were germinated on 1x MS media supplemented with 1% sucrose and 10 μg/ml 5-aza-dC. DNA hypomethylation by 5-aza-dC treatment was monitored by examination of cytosine methylation at the normally methylated 180 bp repeats and 25S rRNA repeats by DNA gel blot analysis as described previously [27]. RNA or DNA was extracted from 4-6-wk-old rosette leaves or from whole 3-wk-old seedlings.

For microarray analysis, seeds were surface-sterilized and plated on $1\times$ MS salts, 0.8% phytagar (Gibco BRL), $1\times$ Gamborg's B5 vitamin mix, and 3% sucrose. Petri plates were incubated vertically for 14 d within a Conviron growth chamber maintained at 21 °C under a 16 h light-8 h dark cycle with a light intensity of 150-175 μ mol \cdot m⁻² sec⁻¹. Whole plant tissue samples were collected and frozen in liquid nitrogen until the RNA was extracted.

RNA isolation and microarray hybridization. A detailed description of RNA isolation, labeling, and hybridization protocols can be found at http://www.ag.arizona.edu/microarray. Each biological replicate consisted of approximately 50 pooled seedlings. Total RNA was isolated using TRIZOL (Invitrogen, Carlsbad, California, United States), and poly(A⁺) mRNA was purified from 75 µg of total RNA using DynaBeads Oligo (dT)25 (Dynal AS, Oslo, Norway) according to manufacturer's instructions. Purified poly (A⁺) mRNA was labeled using either Cy3- or Cy5-dUTP (Amersham Pharmacia Biotech, Piscataway, New Jersey, United States) with Clontech Powerscript reverse transcriptase (Clontech, Mountain View, California, United States). The labeled products were purified using a Millipore Microcon YM30 column (Millipore, Billerica, Massachusetts, United States), washed five times with 100 μ l TE, and the final product was eluted in 40 μ l TE.

For the comparison of *A. thaliana* accessions, we employed long oligonucleotide microarrays provided by the Galbraith laboratory (University of Arizona, Tucson, Arizona, United States; http://www.ag.arizona.edu/microarray), which are produced from a set of 26,088 single stranded 5′ amino-modified oligonucleotides, each 70 bases in length (Qiagen-Operon, Valencia, California, United States, http://loligos.qiagen.com/arrays/omad.php). These oligos have been designed to contain less than 70% homology with any other gene, minimal secondary structure, and to have a single melting temperature of at least 70°C to permit stringent microarray hybridization and washing. Each microarray element was printed once so that all genes could be accommodated on a single slide.

Total RNA from four biological replicates from each of three accessions (Ler, Cvi, and Col) were split in half before being converted to targets, resulting in eight targets from each accession $(3 \times 8 = 24 \text{ total targets})$. Targets were hybridized pair-wise in a three-sided loop design (Ler-Col, Ler-Cvi, and Cvi-Col), giving a total of 12 slides hybridized.

Microarray data acquisition and analysis. Hybridized slides were scanned using a GSI Lumonics ScanArray 3000 (Packard BioChip Technologies, Billerica, Massachusetts, United States). Image processing, including spot finding and quantification of signal intensity, was done using ImaGene 5.0 (BioDiscovery, El Segundo, California, United States). The median fluorescence intensity values for each spot were log base 2 transformed and normalized using the quantile method [61]. No background correction was required. The normalization effectively reduced nonlinear and linear biases due to differential incorporation of dyes, differences between slides, and effects of scanning. Linearity of the data was checked for each pairwise comparison of accessions across all oligos using least-squares means from the final analysis in an RI plot [62]. The remaining genespecific effects and inference of differential expression among accessions was handled in a mixed-model analysis of variance (ANOVA) [63]. On an element-by-element basis, accession and dye were modeled as fixed effects, and slide modeled as a random effect (SAS Proc Mixed; SAS 8.2, SAS Institute). We tested both the raw data and residuals from the fit model for deviations from normality and homoscedasticity, on an element-by-element basis. Because there was no evidence for non-normality or unequal variance for almost all (e.g., $\sim 2\%$ non-normal; unadjusted $\alpha = 0.01$) of the oligos, significance was determined from F-ratios. Since the purpose of this study is to discuss the general relationships among accessions, providing the basis for future work, we employed a cutoff value of 0.001 from the raw p-values. This corresponds to a false discovery rate (FDR) [64] of 0.011 in this data set. After the ANOVA, post-hoc pairwise Tukey tests compared adjusted means (SAS Proc Mixed) for pairwise comparisons between accessions ($\alpha = 0.001$; FDR = 0.023).

Nucleic acid manipulation. DNA was isolated from rosette leaves or whole 3-wk-old seedlings as previously described [65]. RNA was isolated using TRIZOL reagent (Invitrogen) following the manufacturer's instructions, followed by DNAsel treatment (Invitrogen). First strand cDNA was primed with oligo-dT(15) primer using Superscript II reverse transcriptase (Invitrogen) following the manufacturer's protocol. PCR was done using standard conditions with *Taq* DNA polymerase (Qiagen) or KT1 polymerase (Clontech). Primers within the transcribed region of cyclophilin (*At4g38740*) (400 bp amplicon from genomic DNA) [66] were used as PCR amplification controls. All control amplifications utilized the same primer concentration, template amount, and number of cycles as test amplifications. PCR products destined for DNA sequencing were pretreated with exonuclease I/ Antarctic phosphatase (New England Biolabs, Ipswich, Massachusetts, United States) for 30 min at 37 °C.

For RNA gel blot analysis, RNA was size-fractionated by electrophoresis through 1% agarose formaldehyde gels and blotted to GeneScreen (NEN DuPont) nylon membranes using capillary action and $10\times$ SSC buffer. All hybridizations were done following the protocol of Church and Gilbert [67], and membranes were washed at $60\,^{\circ}\text{C}$ in $0.2\times$ SSC, 0.1% SDS. Hybridization probes were radiolabeled using the random priming protocol [68], and unincorporated radionucleotides were removed by size-filtration columns. Gel blots were hybridized with an amplicon from either At2g10410 (X1 + R1, Table S2) or cyclophilin (At4g38740 F + R). At2g10410 RT-PCR/CAPS

analysis involved amplification from cDNA template with primers X1 and R1, followed by a BstB1 (New England Biolabs) digest at 65 °C following the supplier's recommended conditions. The Col allele is cleaved with BstB1, generating a 480 bp fragment plus an undetected 70 bp fragment, while the Ler allele is uncleaved (550 bp) (Figure 4). The additional higher molecular weight band detected in individuals expressing both Ler and Col alleles results from unresolved heteroduplex formation during PCR [69,70]. McrBC (New England Biolabs) digests were carried out at 37 °C overnight using the supplier's recommended conditions.

Genomic DNA from Col and Ler was modified by sodium bisulfite using the CpGenome DNA Modification Kit (Chemicon, Temecula, California, United States) according to the manufacturer's protocols. PCR products were TA-cloned into pGEM-T Easy (Promega, Madison, Wisconsin, United States). Top strand products were amplified with primers Bt1 and Bt2, while bottom strand products were amplified with primers Bb1 and Bb2 (Table S2). For Col, 16 clones were sequenced from the top strand of At2g10410 from two independent amplifications, while 11 clones were sequenced from the bottom strand. For Ler, 17 clones were sequenced from the top strand of At2g10410 from two independent amplifications, and 11 clones from the bottom strand. For Ler ddm1-2, 12 clones were sequenced from the top strand of At2g10410 from two independent amplifications, and 12 clones from the bottom strand. As a control for efficient conversion, we sequenced four clones from each converted template in the promoter region of gene At1g01010, which we had previously determined to be unmethylated (H. Kuo and E. J. Richards, unpublished data); we found that nearly all cytosines (>98.5%) in these clones were converted. DNA sequencing was performed using Big-Dye Terminator Cycle Sequencing (Perkin-Elmer, Wellesley, Massachusetts, United States) protocols/reagents.

Bioinformatics. Sadhu family members and partial elements were identified based on sequence similarity to At2g01410 using iterative searches (BLOSUM62 matrix, gapped alignment, repeat filter off) on the Arabidopsis WU-BLAST server (http://www.arabidopsis.org/wublast/index2.jsp). The NCBI BLAST server (http://www.ncbi.nlm. nih.gov/BLAST) was used to confirm lack of signification sequence similarity of Sadhu elements to sequences outside of A. thaliana available in the public databases. Characterization of features in the vicinity of the Sadhu elements was aided by the repeat masker feature on the Censor server (http://www.girinst.org/censor) [71] and the genome browser at the Arabidopsis thaliana Small RNA Project (ASRP) website (http://asrp.cgrb.oregonstate.edu/cgi-bin/gbrowse/thaliana-v5). RT-PCR, microarray and RNA gel blot characterization of expression of At2g10410 and other Sadhu elements was supplemented by reference to the Arabidopsis Tiling Array Transcriptome Express Tool (http://signal.salk.edu/cgi-bin/atta) [31], the Arabidopsis MPSS database (http://mpss.udel.edu/at/?) [30], and BLASTn to the EST database on the NCBI server (http://www.ncbi.nlm.nih.gov/BLAST). Weblogo (http://weblogo.berkeley.edu/logo.cgi) [72] was used to generate the logo image in Figure 5C. The chromosome map tool at the TAIR website (http://arabidopsis.org/jsp/ChromosomeMap/tool. jsp) aided in generating Figure S3. The maximum parsimony phylogenetic tree in Figure 5A was generated using PAUP* 4.0 (http://paup.csit.fsu.edu/about.html) based on a ClustalX alignment (ftp://ftp-igbmc.u-strasbg.fr/pub/ClustalX). Updated annotations of the sequences of Sadhu family members have been submitted to The Arabidopsis Information Resource (TAIR) (http://arabidopsis.org). Sequence information for all elements listed in Table 3 and Table S1 are available upon request.

Supporting Information

Positions of the 5^\prime and 3^\prime ends of the transcript in Col is indicated. Polymorphisms are highlighted in red.

Found at DOI: 10.1371/journal.pgen.0020036.sg001 (1.5 MB EPS).

Figure S2. Diagram of Bisulfite-Mediated Genomic Sequencing at (A) Ler and (B) Col Alleles of At2g10410 (Positions -105 to +276)

The start of transcription is indicated. The number of circles above a given C indicates the number of clones sequenced; the filled circles denote the proportion of clones which were methylated. CpG sites are indicated in red, CpHpG sites in pink, and CpHpH in blue (where $H=A,\,C,\,$ or T).

Found at DOI: 10.1371/journal.pgen.0020036.sg002 (6.8 MB EPS).



Figure S3. Map Position of 14 Full-Length (Red) and 25 Partial (Blue) *Sadhu* Elements on the Five Chromosomes of the *A. thaliana* Col Genome

Partial elements are marked by their closest gene I.D. number. CEN = position of physical centromere.

Found at DOI: 10.1371/journal.pgen.0020036.sg003 (1.3 MB EPS).

Table S1. List of Partial *Sadhu* Elements in the Col Genome Found at DOI: 10.1371/journal.pgen.0020036.st001 (54 KB DOC).

Table S2. Primers Used in This Study

Found at DOI: 10.1371/journal.pgen.0020036.st002 (47 KB DOC).

Accession Numbers

The GenBank (http://www.ncbi.nlm.nih.gov/Genbank) accession numbers for the *At2g10410* genomic region sequences in Ler and Ra-0 strain backgrounds are DQ385059 and DQ385062, respectively.

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Author contributions. SHR and EJR conceived and designed the experiments. SHR conducted the experiments. SHR and EJR analyzed the data. CV analyzed data for the microarray experiment. RE and HO performed the microarray experiment. DWG conceived and organized the microarray experiment. SHR and EJR wrote the paper.

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