# The mop1 (mediator of paramutation1) Mutant Progressively Reactivates One of the Two Genes Encoded by the MuDR Transposon in Maize

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### ABSTRACT

Transposons make up a sizable portion of most genomes, and most organisms have evolved mechanisms to silence them. In maize, silencing of the *Mutator* family of transposons is associated with methylation of the terminal inverted repeats (TIRs) surrounding the autonomous element and loss of *mudrA* expression (the transposase) as well as *mudrB* (a gene involved in insertional activity). We have previously reported that a mutation that suppresses paramutation in maize, *mop1*, also hypomethylates *Mu1* elements and restores somatic activity to silenced *MuDR* elements. Here, we describe the progressive reactivation of silenced *mudrA* after several generations in a *mop1* background. In *mop1* mutants, the TIRA becomes hypomethylated immediately, but *mudrA* expression and significant somatic reactivation is not observed until silenced *MuDR* has been exposed to *mop1* for several generations. In subsequent generations, individuals that are heterozygous or wild type for the *Mop1* allele continue to exhibit hypomethylation at *Mu1* and *mudrA* TIRs as well as somatic activity and high levels of *mudrA* expression. Thus, *mudrA* silencing can be progressively and heritably reversed. Conversely, *mudrB* expression is never restored, its TIR remains methylated, and new insertions of *Mu* elements are not observed. These data suggest that *mudrA* and *mudrB* silencing may be maintained via distinct mechanisms.

RANSPOSABLE elements make up at least half of the maize genome (SANMIGUEL *et al.* 1996). Given the mutagenic potential of transposons and their ubiquity in plant and animal genomes, it is not surprising that most transposable elements remain quiescent most of the time. Transposon activity is held in check by a well-conserved set of mechanisms that include both post-transcriptional and transcriptional components (ZILBERMAN and HENIKOFF 2004). Mutants have been identified that reactivate quiescent transposons in a variety of species (LIPPMAN et al. 2003) (reviewed in Окамото and HIROCHIKA 2001). In plants, loss of DNA methylation is often associated with transposon reactivation (KATO et al. 2003). Arabidopsis MULE (Mutatorlike element) class II DNA transposons are reactivated in several recessive homozygous mutants, including DDM1 (DECREASE IN DNA METHYLATION1) (SINGER et al. 2001) and MET1 (METHYLTRANSFERASE1) (KATO et al. 2003). Reactivation of Arabidopsis MULEs in a ddm1 mutant background is associated with hypomethvlation of terminal inverted repeats (TIRs), expression of the putative transposase, and new insertions (SINGER et al. 2001). Hypomethylation is heritable in that it persists even in ddm1/+ progeny outcrossed from a ddm1/ddm1parent (LIPPMAN et al. 2003). ddm1 mutants also cause

reactivation and mobilization of CACTA transposable elements, and, like MULE reactivation, this reactivation is heritable (KATO *et al.* 2004).

MET1 is a DNA methyltransferase that is also involved in maintaining global CG methylation (BARTEE *et al.* 2001; KANKEL *et al.* 2003). In *met1* mutants, MULEs become hypomethylated, and the transposase is expressed (LIPPMAN *et al.* 2003). CACTA elements in Arabidopsis are strongly reactivated in lines double mutant for *MET1* and the DNA methyltransferase gene *CMT3* (*CHROMOMETHYLASE3*), which is involved in maintaining non-CG methylation.

In addition to chromatin-remodeling factors, genes involved in RNAi have been implicated in transposon silencing, both in animals (ARAVIN *et al.* 2001; VASTENHOUW *et al.* 2003) and in plants (LIPPMAN and MARTIENSSEN 2004). Indeed, the initiation and maintenance of transposon silencing almost certainly involves a complex interaction between RNAi and chromatin modification.

Arabidopsis is an excellent system in which to study the biochemistry and molecular biology of transposon reactivation in plants. The Arabidopsis genome is fully sequenced, and there are a number of mutants that affect transposon activity. However, the transposon systems in Arabidopsis are poorly characterized; the autonomous elements have not been generally identified, and the means by which the transposons were silenced in the first place is not known. In this respect, maize is unique

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in that it contains several DNA transposon systems that are highly active and that have been extensively characterized.

The Mutator family of transposons is the founding member of the MULE class of elements that are present in plants, fungi, and bacteria (reviewed by (LISCH 2002). All Mutator transposons are characterized by flanking 220-bp TIRs. The Mutator family consists of the autonomous element MuDR and at least nine classes of nonautonomous elements (BENNETZEN 1996). MuDR carries two genes, *mudrA*, which encodes the transposase, and mudrB, a helper gene that is required for transposon integration (LISCH et al. 1995; RAIZADA and WALBOT 2000). The mudrA gene is found in all autonomous MULEs in all species that carry MULEs; the *mudrB* gene is found only in the genus Zea (LISCH 2002). The MuDR TIRs contain the promoters for both mudrA and mudrB (RAIZADA et al. 2001). Methylation of sequences within the TIRs of all elements is correlated with Mutator silencing (CHOMET et al. 1987, 1991; MARTIENSSEN and BARON 1994). A typical maize Mutator active line carries several active MuDR elements as well as up to several hundred nonautonomous elements, and duplication frequencies of these elements can average 100% (Alleman and FREELING 1986).

The large number of active autonomous and nonautonomous elements in the typical Mutator line makes it difficult to perform genetic analysis on the Mutator system. Therefore, we have developed a minimal Mutator line containing a single functional autonomous element, MuDR(p1), at position one (p1) on chromosome 2L, and a single nonautonomous element, Mu1, in the A1 color reporter gene (a1-mum2) (CHOMET et al. 1991) (LISCH et al. 1995). This minimal line provides a simple system in which to easily track the excision and duplication events of a single Mutator element. Excision of Mul from a1-mum2 results in pigmented spots in the kernel aleurone, which is a direct reflection of transposase activity, as is hypomethylation of Mu1 termini. Also, unlike typical Mu active lines, the minimal line does not exhibit spontaneous silencing, which would interfere with genetic analysis of factors influencing epigenetic regulation of the Mutator system.

We have identified and cloned a single dominant gene, *Mu killer* (*Muk*), which heritably and reliably silences the *Mutator* system via a double-stranded RNA mechanism that targets the 5' region of *mudrA* (SLOTKIN *et al.* 2005). *Muk* activity results in silencing of *mudrA* in the first generation and silencing of *mudrB* by the next generation. *Muk* is a variant of *MuDR* that carries an inverted and duplicated portion of the transposon that includes TIRA and a portion of the *mudrA* gene, but that lacks *mudrB* or TIRB. In the presence of *Muk*, all *Mu* TIRs become methylated, and there is loss of somatic and germinal activity (SLOTKIN *et al.* 2003). Once silenced by *Muk*, *Mutator* elements do not become spontaneously reactivated. Thus, *Muk* can be used as a tool to reliably and heritably silence *MuDR* elements to study the dynamics of epigenetic *Mutator* regulation. This, along with the convenience of the minimal *Mutator* line, makes the maize *Mutator* system particularly useful for genetic analysis of transposon silencing and reactivation.

Previous work has demonstrated that a mutant that affects paramutation in maize, mediator of paramutation1 (mop1) (DORWEILER et al. 2000), also reverses Mutator methylation and silencing (LISCH et al. 2002). Paramutation is the phenomenon whereby a silenced allele of a particular gene silences an otherwise active allele in trans (BRINK et al. 1968). The mop1 mutation prevents establishment of paramutation of the paramutable alleles of b1, r1, and pl1 and increases RNA expression of the paramutagenic b1 and pl1 alleles (CHANDLER and STAM 2004). The mop1 mutation also reverses Mu1 TIR methylation in Mutator lines silenced by Muk as well as restoring low levels of somatic activity that was lost due to Muk silencing of MuDR (LISCH et al. 2002). While mop1 affects both paramutation and Mutator TIR methylation, unlike DDM1 or MET1 in Arabidopsis, it neither affects global methylation in maize (DORWEILER et al. 2000) nor affects methylation of some other transposable elements, such as those just upstream of the paramutagenic allele of the b1 locus (LISCH et al. 2002).

In a previous study (LISCH et al. 2002) we reported that *mop1* hypomethylates *Mu1* TIRs and facilitates somatic activity of silenced MuDR elements. Here, we describe in detail the reactivation of silenced MuDR elements after they have been exposed to the mop1 mutation for multiple generations. We find that the TIR adjacent to mudrA, like Mu1 TIRs, is subject to demethylation in a *mop1* background. We find that there is a parent-of-origin effect on the maintenance of the reactivated state and that levels of somatic activity become progressively higher in each generation in a mop1 background, with excision frequencies approaching that observed in lines carrying active MuDR elements. We also find that after several generations in a mop1 background, MuDR elements become heritably active even in the absence of the *mop1* mutation. Surprisingly, despite high levels of somatic activity, we do not find evidence for new insertions, and we also find that, while mudrA expression is fully restored, mudrB remains silenced, consistent with a previously defined role for mudrB in insertional activity (LISCH et al. 1999). Finally, while TIRA becomes hypomethylated in a mop1 background, TIRB remains methylated. These data suggest that although both *mudrA* and *mudrB* are silenced by Muk, there are differences in the means by which that silencing is maintained. The progressive reactivation of silenced *mudrA* (but not *mudrB*) in a *mop1* background affirms that changes in the epigenetic state can be gradual and cumulative and supports the idea that, over time, the memory of the heterochromatic state can be lost in this mutant background.

# MATERIALS AND METHODS

Generation of lines: The a1-mum2 minimal Mutator line and the a1-mum2 minimal line tester: The W22-derived minimal Mutator line was previously generated and described (LISCH et al. 1995). It contains one full-length functional MuDR element and one nonautonomous Mutator element, the Mul element in the allele a1-mum2 (O'REILLY et al. 1985; CHOMET et al. 1991). When an active MuDR element is present, the Mu1 element excises out of a1-mum2 late in somatic development, creating characteristic *Mutator* spotting in the kernel. When MuDR activity is absent the kernel is uniformly pale. A hemizygous MuDR element on the long arm of chromosome 2 named MuDR(p1) (CHOMET et al. 1991) is the single active element in the minimal Mutator line. This element does not become spontaneously epigenetically silenced in the minimal *Mutator* line. When MuDR(p1) is absent, the minimal line is referred to as the *a1-mum2* tester.

Although other *MuDR*-homologous sequences (*hMuDR* elements) are present in this (and all) maize background (CHOMET *et al.* 1991; RUDENKO and WALBOT 2001), these sequences do not appear to contribute to *Mutator* activity in the minimal line either positively (CHOMET *et al.* 1991; LISCH *et al.* 1995) or negatively (SLOTKIN *et al.* 2003, 2005).

mop1 in the minimal Mutator line: Silenced MuDR(p1) elements were generated by crossing MuDR(p1) to Muk. The resulting progeny were crossed to an *a1-mum2* tester and then self-fertilized. Resulting plants were self-fertilized and testcrossed to MuDR(p1) to screen for plants that lacked Muk. Plants that were homozygous for silenced MuDR(p1) [referred to as  $MuDR(p1)^*$  and that lacked Muk were then crossed to *mop1* homozygotes. The resulting *mop1*-heterozygous progeny were crossed to a mopl-homozygous tester that carried the a1-mum2 reporter. None of the resulting  $F_1$  mutant kernels exhibited somatic mutability. A MuDR(p1)\* mop1 mutant progeny of that cross was then self-fertilized. Again, none of the F<sub>2</sub> progeny kernels were spotted. An F2 mop1-homozygous progeny plant carrying MuDR(p1)\* was then testcrossed to a mop1heterozygous tester and the resulting ear had 3% spotted kernels. In the next, F<sub>3</sub>, generation, one exceptional mutant individual carrying  $MuDR(p1)^*$  gave rise to 23% spotted kernels when testcrossed to a mop1-heterozygous tester. Spotted kernels from this cross were again testcrossed to mop1heterozygous testers and again, one exceptional F<sub>4</sub> mutant individual gave rise to 21% spotted kernels. In the next generation  $(F_5)$ , all *mop1*-homozygous individuals that carried  $MuDR(p1)^*$  gave rise to significant numbers of spotted kernels. In this and all subsequent generations the *mop1* mutant phenotype correlated well with the presence of large numbers of spotted progeny kernels. A schematic of all generations and crosses beginning with F<sub>5</sub> can be found in Figure 1.

The  $mop1/mop1;MuDR(p6)^*$  line: The  $mop1/mop1;MuDR(p6)^*$ line was generated in a manner similar to that used to generate the  $mop1/mop1;MuDR(p1)^*$  line, using in this case a MuDRelement at position 6 (p6) that had been previously silenced by Muk and then introgressed into the mop1 mutant background for three generations. MuDR(p6) resulted from the transposition of MuDR(p1) in the minimal line and is therefore a duplicate of that original element in an identical genetic background (the minimal line).

**DNA extraction and Southern blotting:** DNA preparation and genomic Southern blotting were performed as previously described (DORWEILER *et al.* 2000). Maize genomic DNA (10  $\mu$ g) was digested for >4 hr with an excess of 20 units of restriction enzyme. *Mutator* restriction sites and probes referred to in this report are shown in Figure 2.

**RNA extraction and RT–PCR analysis of** *mudrA* and *mudrB*: RT–PCR of *mudrA* and *mudrB* was performed as in (SLOTKIN



FIGURE 1.—Diagram of the crosses and generations discussed in this study.  $F_5$  is the first generation in which a significant percentage of kernel spotting was seen. Each generation indicates the percentage of spotted-kernel progeny produced from the previous generation as well as the genotypes selected for the crosses produced in the next generation. Arrows indicate the testcross.

*et al.* 2003). The same oligo (dT)-primed cDNA used in the RT– PCR analysis of the *mudrA* and *mudrB* transcripts was also amplified with primers specific for the alanine aminotransferase (*aat*) transcript to ensure equal starting amounts of RNA. Amplification was performed for 29 cycles using the primers *aat*F, 5' ATGGGGTATGGCGAGGAT and *aat*R, 5' TTGCAC GACGAGCTAAAGACT. Amplification of *aat* cDNA generates a band of 281 bp, while amplification of the DNA produces a band of 454 bp.

Assay to determine the presence of MuDR(p1): The presence of a full-length MuDR element was assayed by Southern blots using DNA digested with SacI and probing with any internal region of MuDR (see below for generation of probes). If a full-length MuDR element is present, a fragment of 4684 bp is observed. To determine if the MuDR element was at the p1 position, probing with the methyl TIRA probe that flanks p1 and the TIRA with a HinfI-digested blot (see below) assured the presence of MuDR at p1.

Mutator TIR methylation assay: Mutator activity is associated with the methylation status of the HinfI restriction site present in all Mu element TIRs (LISCH et al. 1995). Mutator-active individuals have hypomethylated Mu1 TIRs and will produce a 1.3-kb band when digested with HinfI and probed with the internal region of Mu1. Individuals without MuDR or with silenced MuDR elements have hypermethylated Mu1 TIRs that are not digested by the methyl-sensitive HinfI restriction enzyme, producing Mu1 restriction fragments >1.3 kb (Liscн et al. 1995). The exact size of the inactive Mu1 restriction fragment is dependent on the position of the hypermethylated Mul element. In the allele a1-mum2, the size of this fragment is 2.1 kb. Additional fragments can also be observed that are the result of hybridization of the Mu1 probe to MRS-A, a maize gene that is homologous to Mu1 (CHANDLER et al. 1986). The HinfI sites in this gene, which lack Mu TIRs, are not affected by the presence or absence of MuDR.

The methylation and activity status of MuDR(p1) TIR at mudrA (TIRA) can also be assayed by restriction digestion

using methyl-sensitive restriction enzyme *Hin*fI. Digestion of TIRA with *Hin*fI produces a 607-bp fragment when the *Hin*fI site in TIRA is hypomethylated, and a larger fragment of 1052 bp when it is methylated. Additional fragments visible on Southern blots represent sequences homologous to *MuDR* (*hMuDRs*) that do not contribute either positively or negatively to *Mu* activity.

The methylation status of TIRA and TIRB can also be assayed using the restriction enzyme AfeI, which is blocked by CG methylation. A double digest of AfeI with BamHI produces a 2552- bp MuDR(p1) fragment when the AfeI site is hypomethylated at TIRA and a 2875-bp fragment when AfeI is methylated using the methyl TIRA probe. An AfeI / BamHI double digest produces a 4-kb fragment when the TIRB is hypomethylated at the AfeI site when using the p1 flanking probe; when the AfeIsite is methylated the fragment is  $\sim$ 7 kb and is difficult to resolve on a blot.

**Generation of probes:** *Mu1 probe:* The plasmid that carries the probe for the internal region of *Mu1* has been previously described (TALBERT and CHANDLER 1988). The *Mu1* internal probe is generated by gel isolating an internal *AvaI/Bst*EII fragment.

*Internal MuDR probe:* The internal *MuDR* probe is generated by gel isolating an internal *Eco*RI/*Bam*HI fragment from the pBMP1.3 plasmid (Chomet *et al.* 1991).

*Methyl TIRA probe:* The methyl TIRA probe was generated by amplification using PCR primers TIRA methyl F, 5' CGCGCA CGAGGAAGGCGTTCT, and TIRA methyl R, 5' AGCACCCGT CGCTCCACTTCC. The PCR program was as follows: 94° for 2 min, 94° for 30 sec, 63° for 45 sec, and 72° for 40 min, and then repeated for 35 cycles at 72°, with a last step of 10 min. The PCR product was a 52-bp DNA band.

p1 flanking probe: The p1 flanking probe was generated by the *PstI* digestion of the pBMP1.3 plasmid. The 800-bp p1specific fragment hybridizes to a single-copy sequence in the maize genome (SLOTKIN *et al.* 2003).

All DNA probes in this report were gel isolated and prepared by the random-priming method using a Prime-It II kit (Stratagene, La Jolla, CA) and <sup>32</sup>P-radiolabeled dCTP (Perkin-Elmer, Norwalk, CT). All blots were exposed to a Molecular Dynamics phosphorimaging screen and processed using Adobe Photoshop.

**Genotyping for** *mop1: mop1* is on chromosome 2 at locus 2.04. The primers umc1259 forward and reverse were used to amplify the SSR linked to *mop1* (umc1259 forward: 5' CTCTTT GGTGGCAGAAGCAGAAT; umc1259 reverse: 5' TAGCTAAA CTTGAGTGCTCTGCCC). umc1259 is tightly linked to the *mop1* locus in our minimal lines. When comparing the *mop1* mutant phenotype and a linked umc1259-size polymorphism, we have seen crossovers in <1% of individuals analyzed. Additional SSR markers used for mapping included umc1541 forward and reverse (for primer sequences and amplification conditions refer to http://www.maizegdb.org/).

#### RESULTS

Plants homozygous for *mop1* are hypomethylated at *mudrA* as well as *Mu1*: In the absence of transposase, nonautonomous elements such as *Mu1* become methylated at the TIRs (CHOMET *et al.* 1991). This methylation is fully reversible when functional *MuDR* elements are introduced genetically, and it can be reintroduced *de novo* during development if the transposase is lost due to internal deletions within *MuDR* (LISCH *et al.* 1995). Thus, *Mu1* TIR methylation appears to be the default

state that occurs in the absence of transposase. When otherwise functional MuDR elements are silenced by Muk, their TIRs become methylated as well (SLOTKIN *et al.* 2003). Previous work has shown that Mu1 TIRs are hypomethylated in a *mop1*-homozygous background whether or not a functional MuDR is present, consistent with a role for MOP1 in the default methylation of MuTIRs that occurs in the absence of the transposase (LISCH *et al.* 2002).

To determine whether the TIRs at MuDR(p1) elements silenced by Muk (designated  $MuDR(p1)^*$ ) also become demethylated in a mop1 background, we examined several families that were segregating for mop1 heterozygotes and homozygotes and for  $MuDR(p1)^*$ . Muk was no longer present in any of these families, which had carried  $MuDR(p1)^*$  in a mop1 mutant background for five or six generations. Methylation was assayed by examining Southern blots of DNA digested with HinfI, which is blocked at certain overlapping CG sites. The blots were sequentially probed with an internal fragment of Mul and a fragment flanking MuDR(p1) that included sequences adjacent to the TIR flanking *mudrA* (TIRA) as well as a portion of the TIR (Figure 2). All 43 individuals that genotyped as *mop1* homozygous and that contained  $MuDR(p1)^*$  exhibited hypomethylation at both Mu1 and TIRA. Conversely, all 53 siblings that were mop1 heterozygous and that contained  $MuDR(p1)^*$  remained methylated at both Mul and TIRA (Figure 3A). These data demonstrate that the *mop1* mutation reverses the methylation of TIRA as well as *Mu1* TIRs.

Progressive restoration of high levels of somatic activity of silenced  $MuDR(p1)^*$  elements in a mop1 mutant background: Plants silenced by Muk lose somatic activity of Mutator elements, including Mu1 excision from the a1-mum2 reporter allele (SLOTKIN et al. 2003), resulting in the loss of kernel spotting in the aleurone. In previous work, we observed that mop1-homozygous mutants restored somatic activity to previously silenced MuDR(p1)\* elements in a small percentage of kernels (3%) (LISCH et al. 2002). We wanted to know if somatic activity would increase if  $MuDR(p1)^*$  were carried in a mop1 mutant background for additional generations. After four generations of exposure to the mop1 mutant, a line was derived in which mop1 mutants carrying MuDR(p1)\* consistently produced a high frequency of spotted kernels (see MATERIALS AND METHODS and Figure 1 for derivation of this line). In one family in the F<sub>6</sub> generation, 4 mop1 mutants gave a total of 43% spotted kernels. In contrast, 12 siblings that carried  $MuDR(p1)^*$  but that were heterozygous for mop1 gave only 1% spotted kernels. None of the progeny of 4 plants that were *mop1* mutant but that lacked  $MuDR(p1)^*$  gave rise to spotted kernels (0/291) (data not shown), confirming that the somatic activity we observed is due to MuDR(p1). A subsequent round of crossing (the F7 generation) gave rise to a similar frequency of spotted kernels (Table 1A);



FIGURE 2.—Representation of MuDR at p1 and Mu1 at a1-mum2. Shaded boxes represent Mutator TIRs; open boxes are Mutator internal sequences that differ between MuDR and Mu1. The mudrA and mudrB exons are solid boxes just below the MuDR element; the lines that angle down represent introns. Arrows indicate the direction of transcription. Restriction sites used in this study are indicated. Probes used in this study are indicated below the transcripts. Primers used in this study are shown as arrowheads.

10 *mop1*-homozygous mutants gave a total of 36% spotted kernels (360/989). Again, only individuals that were *mop1* homozygous and carried  $MuDR(p1)^*$  gave rise to significant numbers of spotted kernels (Figure 3B); the 17 *mop1*-heterozygous plants gave rise to only 7% spotted kernels (192/2700) (Table 1B and Figure 3B).

Although the frequency of spotted kernels was consistently much higher in the  $F_6$  and  $F_7$  generations than in earlier generations, there was significant varia-

tion between individual ears, suggesting a stochastic component to the degree of activity achieved by *MuDR* in this mutant background. Similarly, there was a range of spotting intensity among individual kernels on a given ear. Despite this variation, the proportion of heavily and medium spotted kernels was clearly much higher in progeny of *mop1* mutant plants than in their heterozygous siblings (21% vs. 3%). The frequency of excision in these kernels is indistinguishable from that



segregating for mop1/mop1; MuDR\*

FIGURE 3.—(A) Southern blots of a family from the F<sub>7</sub> generation segregating for the mop1 mutation as well as the presence of  $MuDR(p1)^*$ . The same blot was probed for Mu1 and then stripped and reprobed with TIRA. The genotype of each individual is indicated. Individuals that genotype as mop1 homozygous ("\*") are hypomethylated at both Mu1 and TIRA. Individuals that are mop1 heterozygous are methylated at both Mu1 and TIRA. Individuals that lack  $MuDR(p1)^*$  are missing fragments corresponding to both methylated and hypomethylated TIRA. Active MuDR(p1) and  $MuDR(p1)^*$  silenced by Muk are presented as controls. (B) Ear progeny of a cross between indicated individuals represented by the Southern blot by a mop1/+ tester. Individuals that are  $mop1/mop1; \hat{MuDR}(p1)^*$  gave rise to ~36% spotted kernels when crossed to mop1/+. Conversely, individuals that are mop1/+;  $MuDR(p1)^*$  gave rise to very few spotted kernels when crossed to mop1/+. Individuals lacking  $MuDR(p1)^*$ , regardless of *mop1* phenotype, gave rise to no spotted kernels.

Cross <sup>a</sup>	Heavy/medium spotted	Weakly spotted	Pale	Total	Spotted (%)	Heavy/medium spotted (%)
		A. mop1/mop1	; MuDR(p)	l)* siblings		
1	44	33	90	167	46	26
2	8	0	33	41	20	20
3	15	0	33	48	31	31
4	17	3	47	67	30	25
5	27	10	28	68	57	42
6	12	21	105	138	24	9
7	38	30	83	151	45	25
8	12	20	106	139	24	9
9	38	29	84	150	45	25
10	1	0	20	21	5	5
Total	212	148	629	989	36	21
		B. $mop1/+;$	MuDR(p1)	* siblings		
1	14	9	164	187	12	7
2	10	8	104	122	15	8
3	0	2	104	106	2	0
4	2	10	178	190	6	1
5	2	1	146	149	2	1
6	13	4	35	52	33	25
7	4	15	140	159	12	3
8	10	22	170	202	16	5
9	0	5	251	256	2	0
10	8	16	77	101	24	8
11	0	1	16	17	6	0
12	1	4	216	221	2	0
13	3	4	204	211	3	1
14	3	13	173	189	8	2
15	1	0	130	131	1	1
16	2	5	200	207	3	1
17	0	0	200	200	0	0
Total	73	119	2508	2700	7	3

TABLE 1

Somatic excision activity is restored in an mop1 background

<sup>*a*</sup> In the  $F_7$  generation  $mop1/mop1;MuDR(p1)^*$  (A) and mop1/+;MuDR(p1) siblings (B) were crossed to mop1/+ testers.

observed in kernels carrying a fully active MuDR(p1) element.

These results indicate that extended exposure of  $MuDR(p1)^*$  to the mop1/mop1 background can restore high levels of  $MuDR(p1)^*$  activity with respect to somatic excisions. However, in these generations mop1 heterozygotes grown from heavily spotted kernels derived from mop1 mutant parents had methylated Mu1 and MuDRTIRs (Figure 3A) and did not transmit a significant number of spotted kernels (Table 1 and Figure 3B), demonstrating that in these generations  $MuDR(p1)^*$  was not heritably reactivated by mop1.

Somatic excision activity is largely dependent upon the female parental genotype: We observed that in a  $mop1/mop1;MuDR(p1)^* \times mop1/+$  cross, each ear had more than the expected 25% spotted kernels, presuming kernel spotting is dependent upon the kernel genotype being mop1 homozygous and carrying  $MuDR(p1)^*$ . This suggested that at least a subset of the spotted kernels were not *mop1* homozygous. This turned out to be the case. Of 135 individuals genotyped, only 41% of the spotted kernels derived from the above cross were *mop1* homozygous (data not shown). This suggested that kernel spotting is dependent upon the female parent being *mop1* homozygous, and that the genotype of the kernel itself is irrelevant to kernel- spotting intensity.

To determine whether or not the genotype of the male parent had any effect on kernel spotting in the progeny, we performed a series of exact reciprocal crosses of  $F_7$  plants, whereby  $mop1/mop1;MuDR(p1)^*$  individuals were crossed to and by mop1/+ plants (Figure 4). When the female parent was  $mop1/mop1;MuDR(p1)^*$ , 36% of the kernels were spotted, 11% of the total being heavy and medium spotted (Table 2A). In contrast, when the male parent was  $mop1/mop1;MuDR(p1)^*$ , only 19% of the kernels were spotted, and only 1% of which

mop1/mop1; MuDR\* x mop1/+ mop1/+ x mop1/mop1; MuDR\*

FIGURE 4.—Ears resulting from an exact reciprocal cross between *mop1/mop1;MuDR(p1)*\* and *mop1/+*.

were heavy or medium spotted (Figure 4 and Table 2A). A subset of the spotted progeny kernels from several reciprocally crossed individuals were genotyped for mop1. As seen previously, 51% of all spotted kernels genotyped from the female cross were *mop1/+*; the rest were *mop1/* mop1 (Table 3). Interestingly, almost all (93%) of the spotted kernels from the male cross were mop1 homozygous. We also performed reciprocal crosses of *mop1*/  $mop1;MuDR(p1)^*$  to a1-mum2, which is wild type for Mop1. In these crosses the percentage of spotted kernels (31%) (290/921) (Table 2B) produced when *mop1*/ *mop1*;*MuDR(p1)*\* was female and crossed by an *a1-mum2* male was similar to that seen when the female mop1/mop1;MuDR(p1)\* was crossed to a mop1/+ male. In the reciprocal cross, there were fewer spotted kernels (12%) (136/1174) (Table 2B), and there were no heavily or medium spotted kernels.

Together, these data suggest that when the female parent is *mop1/mop1;MuDR(p1)\**, kernel- spotting intensity is independent of the kernel genotype; conversely, when the male parent is *mop1/mop1;MuDR(p1)\**, kernel-spotting intensity is largely dependent upon the kernel genotype.

*MuDR(p1)*\* becomes heritably reactivated after multiple generations in a mop1 background: To comprehensively examine the cumulative effects of *mop1* on  $MuDR(p1)^*$  in the same environment, three generations of kernels ( $F_6$ ,  $F_7$ , and  $F_8$ ) were planted simultaneously in the summer of 2003. In all generations, mop1 homozygotes and heterozygotes carrying reactivated  $MuDR(p1)^*$ were crossed as female to *a1-mum2* (wild type for *Mop1*) and/or *mop1/+*. We found that in the  $F_6$  and  $F_7$  generations, as noted above, all  $mop1/+;MuDR(p1)^*$  progeny plants were methylated at Mul or TIRA (Table 4). When testcrossed, these plants gave rise to few or no spotted kernels (Table 5 and Figure 5). Surprisingly, in the next (F<sub>8</sub>) generation, all  $mop1/+;MuDR(p1)^*$  plants grown from spotted kernels (50 of 50) were hypomethylated at TIRA (Table 4). Of these 50 mop1/+;MuDR(p1)\* plants that were hypomethylated at TIRA, roughly half

(29) were also hypomethylated at Mul; the remaining 21 were not (Table 4). When *Mul*-hypomethylated  $mop1/+;MuDR(p1)^*$  (F<sub>8</sub>) plants were crossed to either *a-1mum2* or *mop1/+* (resulting in the  $F_9$  generation), they gave rise to an average of 47% spotted kernels (Table 6), matching the percentages found in cases where the parent had been  $mop1/mop1;MuDR(p1)^*$ . In contrast, sibling plants that were methylated at Mul (but were hypomethylated at TIRA) gave rise to only 3% (80 of 2286) spotted kernels. These data suggest that in the F<sub>8</sub> generation, although there was a heritable effect on TIRA methylation in all *mop1*-heterozygous progeny that showed high levels of somatic activity in the aleurone, only a subset of those progeny were active enough to hypomethylate Mu1 in the embryo, and it was those individuals that went on to transmit continued somatic activity in a subsequent generation.

One family in the  $F_9$  generation that was segregating for both mop1/+ and homozygous wild- type individuals, was testcrossed by mop1/+ or to a1-mum2 (Table 7). MuDR activity continued to transmit in the resulting ears. Notably, one ear, derived from a cross between a homozygous wild-type plant and an a1-mum2 tester segregated 38% for spotted kernels, demonstrating that the somatic activity of  $MuDR(p1)^*$  persisted even in the absence of even one copy of the mop1 mutant allele.

We wanted to know whether the hypomethylation of Mu1 in  $mop1/+;MuDR(p1)^*$  individuals was the result of a generic effect on Mu1 due to its long-term exposure to mop1 (because mop1 has an effect on Mu1 independent of MuDR activity) or whether it was the result of reactivated  $MuDR(p1)^*$  transposase (which can hypomethylate Mu1 elements independent of mop1). To test this, we examined siblings in the F<sub>8</sub> generation that were mop1/+ but that lacked  $MuDR(p1)^*$ . Mu1 was methylated in all (20 of 20) of these individuals (data not shown), suggesting that MOP1 was present and competent to mediate default methylation of TIRs but that this process is prevented in plants that carry reactivated  $MuDR(p1)^*$  due to activity of the transposase in sibling plants that carried heritably reactivated MuDR elements.

mudrA but not mudrB expression is restored in mop1**reactivated** MuDR(p1)\*: When silenced by Muk,  $MuDR(p1)^*$  loses expression of polyadenylated and nonpolyadenylated *mudrA* and polyadenylated *mudrB* transcript. By the next generation, both mudrA and mudrB become transcriptionally silenced, even in the absence of Muk (SLOTKIN et al. 2003). We wanted to see whether or not the *mop1* mutation was able to restore expression of these genes. In the  $F_6$  and  $F_7$  generations, plants that were *mop1/mop1;MuDR(p1)*\* expressed polyadenylated *mudrA*, but  $mop1/+;MuDR(p1)^*$  sibling plants did not, nor did plants that were mop1 mutant but that lacked  $MuDR(p1)^*$  (Figure 6A). By the F<sub>8</sub> generation however, *mudrA* expression was maintained in *mop1/* +;*MuDR(p1)*\* individuals that are also hypomethylated at Mu1 (Figure 6B) and that transmitted significant

TABL	Æ	2
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Somatic activity exhibits a maternal effect in a mop1 background

	A. mop1/mop1; $MuDR(p1)^* \times mop1/+$						$mop1/+ \times mop1/mop1; MuDR(p1)^*$				
Cross <sup>a</sup>	$h/m^b$	$\mathbf{W}^{c}$	$\mathbf{T}^d$	Spotted (%)	h/m (%)	$h/m^b$	$W^{c}$	$\mathbf{T}^d$	Spotted (%)	h/m (%)	
1	47	52	203	49	23	1	27	187	15	1	
2	26	49	141	53	18	14	93	355	30	4	
3	7	14	50	42	14	3	74	347	22	1	
4	8	45	231	23	3	0	6	206	3	0	
5	18	24	103	41	17	6	2	204	2	3	
6	1	10	21	52	5	0	26	226	12	0	
7	65	32	196	49	33	2	64	236	28	1	
8	3	38	137	30	2	6	68	260	28	2	
9	11	11	60	37	18	6	58	183	35	3	
10	12	51	155	41	8	3	61	225	28	1	
11	0	5	8	63	0	0	4	204	2	0	
12	2	28	139	22	1	0	1	231	0	0	
13	12	6	133	14	9	2	28	136	22	1	
14	0	42	126	33	0	0	42	199	21	0	
15	2	47	172	28	1	1	6	22	32	5	
Total	214	454	1875	36	11	44	556	3221	19	1	
	B. <i>m</i>	op1/mop1	1; MuDR(f	$(b1)^* \times a1$ -mum2			a1-mu	m2 $ imes$ mop	01/mop1; MuDR(p1	)*	
Cross <sup>a</sup>	$h/m^b$	$W^{c}$	$\mathrm{T}^d$	Spotted (%)	h/m (%)	$h/m^b$	$W^{c}$	$\mathrm{T}^d$	Spotted (%)	h/m (%)	
1	4	8	21	57	19	0	14	84	17	0	
2	6	25	70	44	9	0	10	35	29	0	
3	0	4	204	2	0	0	0	200	0	0	
4	7	41	96	50	7	0	6	56	11	0	
5	16	48	143	45	11	0	18	118	15	0	
6	4	11	32	47	13	0	1	81	1	0	
7	5	28	73	45	7	0	0	100	0	0	
8	4	6	20	50	20	0	4	20	20	0	
9	3	0	11	27	27	0	4	104	4	0	
10	9	1	34	29	26	0	26	118	22	0	
11	3	11	23	61	13	0	15	55	27	0	
12	0	0	100	0	0	0	0	25	0	0	
13	16	30	94	49	17	0	38	178	21	0	
Total	77	213	921	31	8	0	136	1174	12	0	

<sup>*a*</sup> In each pair of crosses (numbered), an individual plant was reciprocally crossed as a female (left) or as a male (right) to an mop1/+ (A) or an *a1-mum2* (B) tester.

<sup>*b*</sup> Heavy/medium spotted kernels;

Weakly spotted kernels; and

<sup>d</sup> Total number of kernels.

numbers of spotted progeny kernels when testcrossed (Table 6). However, polyadenylated *mudrB* expression was not restored in a *mop1* mutant background in any individuals, even in later generations in which  $MuDR(p1)^*$  activity had become independent of the *mop1* mutant.

To determine whether the differential reactivation of mudrA and mudrB expression in a mop1 background was due to a position effect of  $MuDR^*$  at p1, we examined mudrA and mudrB expression at another silenced MuDR locus, position 6 (p6).  $MuDR(p6)^*$  had been introgressed into a mop1 background for at least three generations and exhibited a small percentage of kernel spotting,

around 2% (data not shown). We found that  $mop1/mop1;MuDR(p6)^*$  individuals exhibited mudrA but not mudrB expression (Figure 7), just as was seen with  $MuDR(p1)^*$ . This suggests that the differential reactivation of mudrA and mudrB is a generic effect of the mop1 mutation and not due to a position effect on MuDR.

We wished to discover the methylation status of TIRB *vs.* TIRA to see if there was a methylation correlate with the differential *mudrA* and *mudrB* expression in a *mop1* background. Unfortunately our primary methylation assay, the *Hin*fI digest, did not produce conclusive data as the fragment sizes around *mudrB* were too small to

# TABLE 3

Frequency of mop1/mop1 mutants among spotted progeny of mop1/mop1;MuDR(p1)\* plants

mop1/mop1; MuDR(p1)* $\times$ mop1/+						$mop1/+ \times mop1/mop1; MuDR(p1)^*$						
Cross <sup>a</sup>	$m/m^{b}$	$m/+^{c}$	$\mathbf{T}^{d}$	<i>m/m</i> (%)	<i>m/</i> + (%)	Cross <sup>a</sup>	m/m	m/+	Т	<i>m/m</i> (%)	m/+ (%)	
1	15	13	28	54	46	1	12	0	12	100	0	
2	8	5	13	62	38	2	11	0	11	100	0	
3	13	13	26	50	50	3	17	1	18	94	6	
4	8	19	27	30	70	4	14	4	18	78	22	
Total	44	50	94	49	51	Total	54	5	59	93	7	

<sup>a</sup> Genotype of a subset of spotted kernels from reciprocal crosses.

<sup>b</sup> mop1/mop1.

<sup>c</sup> mop1/+;

<sup>d</sup> Total embryos from spotted kernels genotyped for *mop1*.

resolve on a Southern blot. Therefore, we looked to another restriction site, AfeI, which is found in both TIRA and TIRB and that is blocked by CG methylation. We digested DNA from a family segregating for *mop1* and *MuDR(p1)*\* with *Afe*I and *Bam*HI (which cuts once within *MuDR* and which is not methyl sensitive) and then probed the blot with the methyl TIRA probe (as mentioned previously) and subsequently with a *p1* flanking probe (Figure 8). We found that in individuals that were *mop1/mop1;MuDR(p1)*\* the TIRA was hypomethylated, but the TIRB remained methylated, indicating that at the *Afe*I site there is a difference in methylation between TIRA and TIRB.

New transpositions are not observed in a *mop1* background: One of the effects of *Mutator* silencing is the loss of new *Mu* element transpositions. In an active *Mutator* minimal line, new transpositions by *Mu1* and *MuDR(p1)* occur a rate of 10–20% transpositions per generation (LISCH *et al.* 1995). When new *MuDR(p1)* transpositions occur, the resulting ears exhibit a higher percentage of spotted kernels than that expected from an ear segregating for only MuDR(p1) (50% expected spotted kernels in a cross  $MuDR(p1)/- \times -/-$ ). We wanted to see if the *mop1* mutation restored transposition of either  $MuDR(p1)^*$  or Mu1 in silenced *Mutator* lines. To test this, we examined Southern blots for the appearance of unique fragments consistent with new transposition events, and we looked for ears that segregated for significantly more than 50% spotted kernels. HinfI digests of mop1-heterozygous progeny of mop1 mutant plants were probed with Mu1, and BamHI and EcoRI digests were probed with an internal fragment of MuDR. If full transpositional activity were restored, we would expect that between 10 and 20% of the individuals seen would exhibit new transpositions of either Mu1 or MuDR(p1)\*. No new insertions of Mul were observed in 437 progeny of *mop1/mop1* plants that carried reactivated  $MuDR(p1)^*$ . The same was true for the 119 progeny examined by Southern blot for evidence of new insertions of MuDR. In addition, of the 932 ears we examined that were derived from *mop1*-homozygous plants carrying reactivated  $MuDR(p1)^*$ , we have not detected any with a percentage of spotted kernels significantly greater than 50%. This suggests that while the *mop1* mutant can reactivate Mutator somatic activity, it fails to reactivate transpositional activity either in *cis* (*MuDR* insertions) or in trans (Mul insertions). Because mudrB regulates transposition, and because in a mop1 mutant mudrB remains silenced, the lack of transposition in a *mop1* mutant is further evidence for the relationship between transpositional activity and *mudrB* activity.

# DISCUSSION

Unlike the immediate and heritable reactivation of silenced transposons in a *ddm1* mutant background

Hypomethylation of mop1/+ progeny of mutant individuals over several generations

Cross	Hypo TIRA <sup>a</sup>	Нуро Ми1 <sup>ь</sup>	Meth Mu1 <sup>c</sup>	Total TIRA	Hypo TIRA (%)	Hypo <i>Mu1</i> (%)	Meth Mu1 (%)
$F_6$	0	0	0	75	0	0	0
$F_7$	0	0	0	27	0	0	0
F <sub>8</sub>	50	29	21	50	100	58	42

<sup>*a*</sup> Number of *mop1/+* progeny plants with hypomethylation at TIRA among heavily spotted mop1/+ progeny plants.

<sup>b</sup> Of plants that were hypomethylated at TIRA, number that were also hypomethylated at Mu1.

<sup>c</sup> Of plants that were hypomethylated at TIRA, number that were methylated at Mu1.</sup>

TABLE 5	TA	BL	Æ	5
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Cross <sup>a</sup>	Heavy/medium spotted	Weakly spotted	Pale	Total	Spotted (%)	Heavy/medium spotted <sup>b</sup> (%)	No. of crosses
		A. Crosse	es by a <i>ma</i>	pp1/+ test	ers		
$F_6 mop1/mop1$	241	64	411	716	43	34	4
$F_6 mop1/+$	6	13	1276	1295	1	0	12
$F_7 mop1/mop1$	212	148	627	987	36	21	10
$F_7 mop1/+$	73	119	2508	2700	7	3	17
$F_8 mop1/mop1$	124	31	248	403	38	31	5
$F_8 mop1/+$	644	272	1813	2729	34	24	17
$F_9 mop1/+$	504	179	799	1482	46	34	5
		B. Cross	es to al-n	num2 teste	ers		
$F_7 mop1/mop1$	734	177	1028	1939	47	38	15
$F_7 mop1/+$	108	105	1416	1629	13	7	11
$F_8 mop1/mop1$	126	71	191	388	51	32	3
$F_8 mop 1/+$	260	98	411	770	46	34	6
$F_9 mop1/+$	342	164	789	1285	39	26	7

MuDR(p1)\* activity becomes heritable after several generations in an mop1 background

<sup>*a*</sup> Generation ( $F_X$ ) and genotype of the female plants carrying  $MuDR(p1)^*$  crossed to the indicated male genotypes; all individuals from a particular generation are siblings.

<sup>b</sup> Percentage of heavy/medium spotted of total kernels.

(LIPPMAN *et al.* 2003; KATO *et al.* 2004), reactivation of *MuDR* elements in a *mop1* mutant background occurs only gradually and stochastically. Only after multiple generations of exposure to the *mop1* mutation do we observe evidence of heritable activity of silenced *MuDR* elements in the absence of the mutation. This process is

reminiscent of the gradual appearance of epimutations, such as *fwa*, after several generations in a *ddm1* mutant background (KAKUTANI *et al.* 1996; SOPPE *et al.* 2000). Interestingly, the promoter and upstream portion of the FWA transcript are composed of a transposable element (LIPPMAN *et al.* 2004). These data suggest that the effects



FIGURE 5.—(A) Mul and TIRA Southern blots of individuals from both F7 and F8 generations. All individuals are mop1/+. Note that in the  $F_7$ generation, all individuals are methylated at both Mu1 and TIRA. In the F<sub>8</sub> generation in this figure, all individuals that are  $mop1/+;MuDR(p1)^*$ are hypomethylated at Mu1 and TIRA, indicating that MuDR(P1)\* has been heritably reactivated in this generation. Individuals lacking MuDR(p1)\* are methylated at Mu1 and lack the methylated or the hypomethylated TIRA fragments. (B) Ear progeny of a cross between indicated individuals represented by the Southern blot to a mop1/+tester. mop1/+ individuals from the F<sub>7</sub> generation gave rise to few or no spotted-kernel progeny when outcrossed to a mop1/+ tester. However, Mu1-hypomethylated  $mop1/+;MuDR(p1)^*$  individuals from the  $F_8$  generation gave rise to  $\sim 46\%$ spotted kernels when outcrossed to mop1/+.

#### TABLE 6

		1	·	ų /	, 00	<i>.</i>	1 0 /		
$h^a$	Cross to	Spotted	Total	Spotted (%)	$m^a$	Cross to	Spotted	Total	Spotted (%)
1	mop1/+	80	184	43	1	mop1/+	3	123	2
2	mop1/+	48	99	48	2	mop1/+	1	112	1
3	mop1/+	108	207	52	3	a1-mum2	4	204	2
4	a1-mum2	58	123	47	4	mop1/+	3	158	2
5	mop1/+	178	334	53	5	a1-mum2	6	192	3
6	mop1/+	66	222	30	6	a1-mum $2$	7	207	3
7	mop1/+	127	258	49	7	mop1/+	1	121	1
8	mop1/+	96	172	56	8	mop1/+	2	202	1
9	a1-mum2	120	256	47	9	a1-mum2	4	203	2
10	a1-mum2	46	92	50	10	a1-mum2	14	214	7
11	mop1/+	96	263	37	11	mop1/+	27	227	12
12	a1-mum2	168	327	51	12	mop1/+	5	186	3
13	a1-mum $2$	144	280	51	13	mop1/+	3	137	4
Total		1335	2817	47	Total		80	2286	3

Persistence of somatic activity even in the absence of the *mop1* mutation: *Mu1* methylation of  $mop1/;+MuDR(p1)^*$  individuals, F<sub>8</sub> generation, and progeny

As indicated in Table 4, all plants were hypomethylated at TIRA, but only about half were also hypomethylated at Mu1 h, plant had hypomethylated Mu1 element; m, plant had methylated Mu1 element. Plants were crossed by either a1-mum2 or mop1/+ testers as indicated.

<sup>a</sup> Plants grown from the F<sub>8</sub> generation kernels that were spotted.

of missing one component of the silencing machinery (MOP1 in this case, DDM1 in Arabidopsis) can be cumulative over time, perhaps because the loss of one component results in destabilization of a complex, which in turn leads to the loss of additional components and subsequent additional destabilization. Since a number of factors involved in the construction of stabilized chromatin appear to be mutually reinforcing (i.e., DNA methylation and histone modification (JOHNSON et al. 2002; SOPPE et al. 2002), it is reasonable to assume that, although the loss of one component may not have an immediate effect of gene activity, the absence of reinforcement of the silenced state could eventually result in destabilization of the silencing chromatin. Several other mutations that affect both paramutable alleles and Mu element methylation have been identified (LISCH and HOLLICK, unpublished data). It will be

interesting to see how efficiently double mutants reactivate silenced *MuDR* elements.

The absence of both new insertions and *mudrB* expression in our reactivated lines is consistent with earlier data suggesting a role for *mudrB* in *Mu* element transposition (LISCH *et al.* 1999) (RAIZADA and WALBOT 2000). Like our reactivated *MuDR* elements, deletion derivatives that lack *mudrB*, as well as transgenes that express only *mudrA*, can condition only excisions. Differential reactivation of two genes on the same transposon has not been observed previously. It suggests that the silenced state of these two genes differs in some way. Either *mudrB* is simply more deeply silenced than *mudrA* or the silenced state of the two genes is qualitatively different. There is some evidence that the latter may be the case, since TIRA becomes hypomethylated in a *mop1* background but TIRB does not. Further, although both

TABLE 7			
Crosses from one family of the F <sub>9</sub> generation segregating f	for v	vild	type

Cross <sup>a</sup>	$h/m^b$	$W^{c}$	Pale	$\mathrm{T}^{d}$	Spotted (%)	h/m (%)	Kernels (% wt)
$mop/+; MuDR(p1)^* \times a1-mum2$	2	6	93	101	8	2	75
$mop/+; MuDR(p1)^* \times a1-mum2$	62	48	145	255	43	24	75
$mop/+; MuDR(p1)^* \times a1-mum2$	7	12	140	159	12	4	75
$+/+; MuDR(p1)* \times mop1/+$	1	10	102	113	10	1	75
$+/+; MuDR(p1)^* \times mop1/+$	45	39	68	152	55	30	75
$+/+; MuDR(p1)^* \times a1-mum2$	38	14	84	136	38	28	100

<sup>a</sup> Plants in this generation were either heterozygous for mop1 or homozygous for Mop1.

<sup>*b*</sup> Heavy/medium spotted kernels.

<sup>e</sup> Weakly spotted kernels; and

<sup>d</sup> Total number of kernels.



FIGURE 6.—mudrA and mudrB expression. (A) RT–PCR of both mudrA and mudrB in the  $F_7$ generation. Individuals that are mop1/mop1; MuDR(p1)\* express mudrA, but mop1/+;MuDR(p1)\* individuals and those that lack MuDR(p1)\* do not express mudrA. However, mop1/mop1;MuDR(p1)\* individuals do not express mudrB. (B) RT–PCR of mudrA and mudrB in the  $F_8$  generation. mop1/+; MuDR(p1)\* as well as mop1/mop1;MuDR(p1)\* individuals now express mudrA, in conjunction with the heritable reactivation of MuDR(p1)\* in this later generation. Once again, mudrB is still not expressed in any individual, not even mop1 homozygotes. aat is the cDNA control.

genes are silenced after exposure to Muk, their mode of silencing appears to be different. Muk is an inverted duplicated version of *MuDR* that lacks the *mudrB* gene. It produces a long inverted repeat transcript that is homologous to the *mudrA* gene that produces small (23 and 26 nt) mudrA-homologous RNA molecules that are amplified when Muk silences MuDR elements. No such RNA molecules homologous to *mudrB* are observed (SLOTKIN et al. 2005). The mudrA gene is transcriptionally silenced by the immature ear of F<sub>1</sub> plants carrying both MuDR and Muk. In contrast, the mudrB gene remains transcriptionally active in these same F<sub>1</sub> immature ears, and it is only in the next generation that mudrB becomes transcriptionally silenced (SLOTKIN et al. 2003). This process requires that *mudrB* be in *cis* to *mudrA*; when a deletion derivative of *MuDR(p1)* that carries only *mudrB* is placed in *trans* to *MuDR(p1)*, it is not silenced in the presence of Muk (SLOTKIN et al. 2005). Thus, mudrB silencing is mediated via *mudrA* silencing, most likely via a distinct pathway that involves a signal that spreads in cis from mudrA to mudrB. If this is the case, then the chromatin at silenced *mudrB* may well be qualitatively different from that at *mudrA*.

The identity of *mop1* remains mysterious, so we can only speculate as to the mechanism of MuDR reactivation. The fact that this mutation affects both paramutation and MuDR activity but neither global methylation nor methylation of some transposon sequences upstream of the maize paramutagenic b1 allele implies a relatively restricted (or partially redundant) role in maize gene silencing. This is reinforced by our observation that only one of the two genes encoded by MuDR is reactivated in a mop1 background. The relative specificity of the mop1 mutation may be because the *Mop1* gene is qualitatively different from mutations in Arabidopsis that affect transposon activity, or because the more global functions of those genes have been partitioned among multiple genes in maize. This possibility is supported by the existence of multiple mutations that affect both Mutator methylation and paramutation (HOLLICK and CHANDLER 2001; LISCH and HOLLICK, unpublished data).



FIGURE 7.—mudrA and mudrB expression at another MuDR position  $(MuDR(p6)^*)$ . Individuals that are  $mop1/mop1;MuDR(p6)^*$  express mudrA but not mudrB, similar to what is observed in reactivated  $MuDR(p1)^*$ .

MuDR(p1)\* and AfeI methyl-sensitive restriction sites in the TIRs



FIGURE 8.—Methyation analysis of TIRA vs. TIRB. Shown is a Southern blot depicting individuals carrying  $MuDR(p1)^*$ and that are mop1/mop1 or mop1/+. Individuals were digested with AfeI (a methyl-sensitive enzyme that cuts once within each TIR) and BamHI (not methyl sensitive and that cuts once within the MuDR element as well as outside each TIR). The Southern blot was probed first with the methyl TIRA probe and then stripped and reprobed with the p1 flanking probe.  $mop1/mop1;MuDR(p1)^*$  individuals are hypomethylated at TIRA but are not hypomethylated at TIRB. Active MuDR is hypomethylated at both TIRA and TIRB. Silenced MuDRand  $mop1/+;MuDR(p1)^*$  individuals are hypomethylated at neither TIR.

An understanding of the differences between *mudrA* and *mudrB* silencing and resulting differences in chromatin at the *mudrA* and *mudrB* promoters at various stages of reactivation, should shed light on the relationship between the means by which genes are silenced and the nature of the silenced state once achieved. The evidence to date suggests that the initiation and maintenance of silencing is far more complex than simple "on" and "off" states. History, position, context, and timing may all play a role in these processes. The cumulative reactivation of *mudrA* in a *mop1* background suggests that changes in chromatin memory through several rounds of meiosis can be gradual, and that there are intermediate chromatin states between silenced and active chromatin that may in themselves have unique regulatory consequences and interesting implications for our understanding of heritable changes in the chromatin state.

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