Biased transcription and selective degradation of small RNAs shape the pattern of DNA elimination in *Tetrahymena*

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The ciliated protozoan *Tetrahymena* undergoes extensive programmed DNA elimination when the germline micronucleus produces the new macronucleus during sexual reproduction. DNA elimination is epigenetically controlled by DNA sequences of the parental macronuclear genome, and this epigenetic regulation is mediated by small RNAs (scan RNAs [scnRNAs]) of ~28–30 nucleotides that are produced and function by an RNAi-related mechanism. Here, we examine scnRNA production and turnover by deep sequencing. scnRNAs are produced exclusively from the micronucleus and nonhomogeneously from a variety of chromosomal locations. scnRNAs are preferentially derived from the eliminated sequences, and this preference is mainly determined at the level of transcription. Despite this bias, a significant fraction of scnRNAs is also derived from the macronuclear-destined sequences, and these scnRNAs are degraded during the course of sexual reproduction. These results indicate that the pattern of DNA elimination in the new macronucleus is shaped by the biased transcription in the micronucleus and the selective degradation of scnRNAs in the parental macronucleus.

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The phenotypes of a parent can be epigenetically transmitted to offspring through the germline in many different eukaryotes. Epigenetic information can be inherited as chromatin modifications that are incompletely reprogrammed in gametes and thereby transmitted to descendants. For example, flower shape in plants and coat color in mice can be transmitted transgenerationally by variations in DNA methylation patterns (Cubas et al. 1999; Morgan et al. 1999). Alternatively, some diffusible factors can transmit epigenetic information from parents to progeny through the germline, such as the inheritance of transposon silencing states by maternally supplied Piwi-associated small RNAs in fruit flies (Brennecke et al. 2008). Transgenerational epigenetic inheritance potentially allows organisms to transmit acquired characteristics to the next generation and thus could increase the fitness of progeny in rapidly changing environments. However, the detailed molecular mechanism of transgenerational epigenetic inheritance is largely unclear. Several lines of evidence suggest that the patterns of programmed DNA elimination in ciliated protozoans are epigenetically and transgenerationally inherited (for review, see Mochizuki 2010). Therefore, DNA elimination in ciliated protozoans serves as a model to understand how transgenerational epigenetic inheritance can be achieved at a molecular level.

Like most ciliated protozoans, *Tetrahymena thermophila* exhibits nuclear dimorphism. Each cell contains a germ-line micronucleus [Mic] and a somatic macronucleus [Mac] (Fig. 1). Most, if not all, gene expression occurs in the Mac. In the sexual process of conjugation (Fig. 1A–G), the Mic undergoes meiosis (Fig. 1A), and one of the four meiotic products is selected and mitotically divides to form two haploid pronuclei (Fig. 1B). One of the pronuclei is reciprocally exchanged between the two conjugating cells (Fig. 1C). The migratory and stationary pronuclei then fuse to create a zygotic diploid nucleus (Fig. 1D) that divides twice mitotically (Fig. 1E) to produce the next generation of new Macs and Mics (Fig. 1F). The parental Mac is then destroyed (Fig. 1G).

Programmed DNA elimination occurs during the development of the new Mac. Previous studies have estimated that ~6000 internal eliminated sequences (IESs) are removed from the new Mac (for review, see Yao et al.
The Mic sequences that remain in the future Mac are called Mac-destined sequences (MDSs). IESs in Tetrahymena vary in size (0.5–20 kb) and sequence, and recent Mic genome sequencing indicates that they account for about one-third of the Mic genome (Tetrahymena Comparative Sequencing Project, Broad Institute of Harvard and Massachusetts Institute of Technology [http://www.broadinstitute.org]). Some IESs are moderately repeated in the Mic genome [Yao and Gorovsky 1974] and are related to transposable elements [Wuitschick et al. 2002, Fillingham et al. 2004]. IESs are primarily located in nongenic regions, although some are located in genic regions [Fass et al. 2011]. The excision of IESs can occur reproducibly at a specific site or with a limited number of alternative boundaries [Austerberry et al. 1989; Fass et al. 2011]. However, despite this precision, the sites of DNA elimination do not seem to be specified by the primary DNA sequences of the Mic genome because no obvious consensus sequence has been identified in or around IESs, and the in vitro DNA specificity of the enzyme responsible for the DNA excision process is ambiguous [Cheng et al. 2010]. Instead, DNA elimination in Tetrahymena and some other ciliates is epigenetically regulated by the DNA sequence of the parental Mac such that the elimination of a new Mac DNA is inhibited by the presence of the same sequence in the parental Mac [Koizumi and Kobayashi 1989; You et al. 1991; Duharcourt et al. 1995; Chalker and Yao 1996]. Because the chromosomes of the parental and the new Macs do not directly interact, some sequence-specific diffusible signal must be transmitted from the parental to the new Macs through the cytoplasm to mediate the epigenetic effect [Chalker et al. 2005].

The injection of dsRNAs complementary to a MDS locus into an early conjugating cell induces ectopic DNA elimination at the MDS locus [Yao et al. 2003], indicating that an RNAi-related mechanism plays a role in DNA elimination. Indeed, the Dicer homolog Dcl1p and the Argonaute protein Twi1p are essential for DNA elimination [Mochizuki et al. 2002, Malone et al. 2005]. Bidirectional Mic noncoding transcripts [Chalker and Yao 2001], which are probably transcribed by RNA polymerase II [Mochizuki and Gorovsky 2004b], are processed by Dcl1p [Malone et al. 2005; Mochizuki and Gorovsky 2005] to small (~28- to 30-nucleotide [nt]) RNAs, named scan RNAs (scnRNAs). scnRNAs are then complexed with Twi1p [Mochizuki and Gorovsky 2004a]. The fact that
the Twi1p–scnRNA complex first localizes to the parental Mac and then to the new Mac (Mochizuki et al. 2002; Noto et al. 2010) makes scnRNA a strong candidate to be the mediator of the epigenetic signal.

Consistent with the production of scnRNAs from the Mic, scnRNAs complementary to MDSs and those complementary to IESs have been detected in the early stages of conjugation. However, scnRNAs complementary to IESs become enriched as conjugation proceeds (Mochizuki and Gorovsky 2004a; Aronica et al. 2008). Moreover, most of the scnRNAs cloned from the late stage of conjugation are complementary to IESs (Lee and Collins 2006). Thus, scnRNAs are selected for IES specificity after they are diced. Based on these and other observations, a model has been proposed to explain how scnRNAs mediate the epigenetic regulation of DNA elimination (Fig. 1H–M). In the model, scnRNAs are produced from the entire Mic genome, including both IESs and MDSs, in the early stages of conjugation (Fig. 1H,J), and scnRNAs that share homology with any Mac genome sequences (MDSs) are degraded in the parental Mac in the middle stages of conjugation (Fig. 1I,K). This process is called scnRNA selection. It has been further proposed that these IES-specific scnRNAs move to the developing Mac in the late stages of conjugation (Fig. 1L), where they identify homologous sequences as IESs and target them for heterochromatin formation (Coyne et al. 1999; Taverna et al. 2002; Liu et al. 2007), followed by DNA elimination (Fig. 1M).

Three issues with this proposed model have not been resolved: (1) Are scnRNAs produced exclusively from the Mic? (2) Are scnRNAs produced homogeneously from both IESs and MDSs at a local level and from the entire Mic genome at a global level? (3) Does scnRNA selection occur at a genome-wide level? To address these points, we analyzed the production and turnover of scnRNAs by deep sequencing.

**Results**

*scnRNAs are predominantly produced from the Mic*

It has been proposed that scnRNAs are produced exclusively from the Mic because some fraction of scnRNAs is complementary to the Mic-limited sequences (Mochizuki et al. 2002; Mochizuki and Gorovsky 2004a; Aronica et al. 2008), and the Dicer homolog Dcl1p, which is responsible for the production of scnRNAs, is detected only in the Mic (Malone et al. 2005; Mochizuki and Gorovsky 2005). However, another fraction of scnRNAs is complementary to MDSs (Mochizuki et al. 2002; Mochizuki and Gorovsky 2004a; Aronica et al. 2008), which are shared in the Mic and the Mac; therefore, the production of scnRNAs from the Mac has not been completely excluded. To determine whether the Mac produces scnRNAs, we compared scnRNAs from wild-type cells with those from chromosome 4 [Chr4]–nullisomic (Nulli4) strains. In Nulli4 strains, Chr4 is completely missing in the Mic, while the Mac chromosomes derived from Chr4 are maintained (Bruns et al. 1983). Therefore, if any Chr4-derived scnRNA is detected in Nulli4 strains, it must be transcribed from the Mac genome.

Total RNA was extracted from wild-type and Nulli4 strains at 3 h post-mixing, when the production of scnRNAs is expected to be almost complete but scnRNA selection is expected to have just initiated (see below). The RNA was gel-fractionated, and a small RNA population, including scnRNAs [26–32 nt], was used to construct the cDNA library for deep sequencing. To determine from which Mic chromosome each scnRNA was derived, we chose two ~100-kb Mic genome loci for which chromosomal origins have been identified. One, named the Cal locus, contains the calmodulin gene and 12 IESs, including the well-studied Cal-IES element (Katoh et al. 1993); the other, named the LMR locus, contains 13 IESs, including the well-studied L-IES, M-IES, and R-IES elements (Austerberry et al. 1984). The Cal and LMR loci are on Chr2 and Chr4, respectively (E. Hamilton and E. Orias, pers. comm.).

The numbers of scnRNAs that uniquely mapped to the loci (unique mappers) were calculated. The first 25-nt sequences of scnRNAs [26–32 nt] that did not match to tRNAs, trRNAs, snRNAs, and the mitochondrial genome (filtered reads) (Supplemental Table S1) were used for mapping. The number of scnRNAs from wild-type and Nulli4 cells that mapped to the Cal locus were similar [Fig. 2A,B]. In contrast, the number of scnRNAs that mapped to the LMR locus was significantly lower in Nulli4 strains [Fig. 2D] than in wild-type strains [Fig. 2C]. We believe that the small number of scnRNAs from Nulli4 strains that mapped to the LMR locus were transcribed from non-Chr4 Mic chromosomes because some repeated sequences likely failed to be assembled into the current draft Mic genome, and a small proportion of repeated sequences that were present on both Chr4 and a different chromosome could have been falsely recognized as unique to Chr4 in this study. Consistent with this view, most of the scnRNAs from Nulli4 strains that mapped to the LMR locus were complementary to IESs [Fig. 2D, IESs are marked in red] that did not exist in Nulli4 strains. Therefore, we believe that Nulli4 strains do not produce scnRNAs from the LMR locus and conclude that scnRNAs are produced predominantly, if not exclusively, from the Mic.

*scnRNAs are preferentially produced from IESs*

The current model of scnRNA-directed DNA elimination (Mochizuki and Gorovsky 2004a; Aronica et al. 2008) suggests that scnRNAs are produced homogeneously from both IESs and MDSs in the Mic genome. To determine whether this occurs at the genome-wide level, we first annotated the draft Mic genome. All possible 25-nt sequences from the draft Mic genome were sorted to three classes: sequences found only in MDSs, sequences found only in IESs, or sequences found in both MDSs and IESs (see the Materials and Methods for details), which correspond to 98.8 Mb, 33.1 Mb, and 3.0 Mb of genomic sequences, respectively [Fig. 3A, left]. We used the sum of these classified sequences as the annotated Mic genome.
in this study. To analyze the original scnRNA population produced by Dcl1p in the Mic, we used TWI1 knockout cells because the scnRNA selection process (Fig. 1K; see also below) that occurs after binding of scnRNAs to Twi1p should not occur in the absence of Twi1p.

The first 25-nt sequences of scnRNAs (26–32 nt) from TWI1 knockout cells at 3 h post-mixing were mapped to the annotated Mic genome. Among the scnRNAs from TWI1 knockout cells, 67.0% mapped to the annotated Mic genome without any mismatch (Supplemental Table S1). Of these matched scnRNAs, only 15.3% were exclusively complementary to MDSs, while 80.1% mapped exclusively to IESs (Fig. 3A, right; Supplemental Table S1). This distribution differed dramatically from the profile of the annotated genome sequence, which included 73.2% MDSs and 24.6% IESs (Fig. 3A, left). Therefore, in contrast to expectations, the production of scnRNAs is heavily biased toward IESs.

A similar conclusion was obtained by mapping scnRNAs to the Cal and LMR loci. The numbers of scnRNAs mapping to each position of these loci were calculated, and the repetitiveness of the sequences in the Mic genome was normalized by either dividing the numbers of mapped scnRNAs by their sequence frequencies in the draft Mic genome [Fig. 3B,D, “weighted”] or excluding scnRNAs that were complementary to more than one position in the draft Mic genome [Fig 3C,E, “uniquely mapped”]. Both of the analyses demonstrated that the majority of scnRNAs were complementary to IESs and that the other scnRNAs were complementary to MDSs [Fig. 3B–E, IESs are marked in red]. These results further indicate that scnRNAs are preferentially produced from IESs rather than MDSs.

**scnRNAs are produced from a variety of Mic genome loci at various levels**

It has been proposed that scnRNAs are produced not from specific chromosomal regions, but rather from the entire Mic genome (Mochizuki et al. 2002; Mochizuki and Gorovsky 2004c). As shown above, this is not the case at a local level because the scnRNA production is biased to IESs [Fig. 3A–E]. Therefore, we next analyzed the origin of scnRNAs at a global level. We mapped scnRNA sequences to MDSs to analyze scnRNA production from different chromosomal locations without the effect of varied IES densities at different loci (Fass et al. 2011). All Mic genome scaffolds longer than 100 kb, which account for ~60% of the sequenced Mic genome in total, were divided into 100- to 200-kb fragments and used as representative MDS fragments.

We first analyzed scnRNAs from TWI1 knockout cells at 3 h post-mixing. The density [reads per kilobase per million scnRNAs, RPKM] of scnRNAs that uniquely mapped to each of the MDS fragments was analyzed. Because the total length of the mappable sequences in the MDS fragments was 87.3 Mb, and 2.16 million of the 18.4 million scnRNA reads uniquely mapped to these fragments, the density was expected to be 1.34 RPKM if the scnRNA sequences mapped homogeneously to the genome. However, the densities of the mapped scnRNAs differed dramatically between the MDS fragments [Fig. 3F, column a], with a median density of 0.506 RPKM.
We hypothesized that the observed variability in the scnRNAs from TWI1 knockout cells that mapped to the MDS fragments could account for the timing but not the levels of scnRNA production from different chromosomal locations. This is because scnRNAs are unstable in the absence of Twi1p (Mochizuki and Gorovsky 2004a), and thus only scnRNAs produced immediately before harvesting the cells were likely sequenced from TWI1 knockout cells. In wild-type cells, scnRNAs are stabilized in a complex with Twi1p (Mochizuki and Gorovsky 2004a). On the other hand, in wild-type cells, Twi1p transports scnRNAs to the parental Mac, where scnRNAs complementary to the Mac genome are degraded (Fig. 1K; Aronica et al. 2008). The Twi1p-associated RNA helicase Ema1p is required for this degradation in the parental Mac but not for the formation of Twi1p–scnRNA complexes, which stabilize scnRNAs (Supplemental Fig. S3; Aronica et al. 2008).

Therefore, to circumvent scnRNA degradation and determine the level of scnRNA production from each locus more precisely, scnRNA sequences obtained from EMA1 knockout cells at early to late stages of conjugation were mapped to the MDS fragments (Fig. 3F, columns b–e). Although many of the MDS fragments were sparsely (<1 RPKM) mapped by scnRNA sequences from TWI1 knockout cells (Fig. 3F, column a), this fraction was greatly

Figure 3. Production of scnRNAs from the Mic genome. (A) The draft Mic genome was annotated as three sequence classes: sequences that appear only in MDSs [blue], sequences that appear in both MDSs and IESs [green], and sequences that appear only in IESs [red]. The percent occupancy of genomic sequences in each class is shown in the left column (Genome). The fraction of the sequenced scnRNAs from TWI1 knockout cells at 3 h post-mixing that is complementary to each of the classes of the annotated Mic genome is shown in the right column (scnRNA). (B–E) The Cal (B, C) and LMR (D, E) loci are mapped by the sequenced scnRNAs from TWI1 knockout cells at 3 h post-mixing. For B and D, the number [reads per million scnRNAs (RPM)] of scnRNAs mapping to each position of the loci was divided by its sequence frequency in the draft Mic genome ("weighted"). For C and E, only the number of scnRNAs that mapped to a single location of the draft Mic genome is shown ("uniquely mapped"). In both cases, the numbers of scnRNAs matching the sense and antisense strands are shown in the plus and minus directions, respectively, as histograms with 100-nt bins. (C, E, bottom) The fraction of unique sequences that can be mapped by scnRNAs by the latter method in each 100-nt bin is shown as "Mappability". The positions of IESs are marked in red. (F) The number of scnRNA sequences that uniquely matched each of the 100- to 200-kb MDS fragments was divided by the total length of the unique sequences of the MDS fragments [scnRNA density, RPKM (reads per kilobase of unique sequences per million) scnRNAs] and is shown as box plots. scnRNAs from TWI1 knockout cells at 3 h post-mixing and from EMA1 knockout cells at 3, 4.5, 6, or 8 h post-mixing were analyzed. The box indicates the interquartile range (IQR). The horizontal bar in the box indicates the median value. The bars on the top and bottom of the box indicate the minimum and maximum values, respectively, within 1.5 × IQR. Circles indicate outliers. P-values were calculated using the Wilcoxon rank-sum test.
reduced when the MDS fragments were mapped by scnRNA sequences from EMA1 knockout cells at 3 h post-mixing [Fig. 3F, column b]. This result indicates that some population of scnRNAs that was derived from specific chromosomal locations existed in EMA1 knockout cells but was missing in TWI1 knockout cells. The variability of the mapped scnRNA density on different MDS fragments was smaller at 4.5 h post-mixing [Fig. 3F, column c] than at 3 h post-mixing [Fig. 3F, column b] in EMA1 knockout cells but did not significantly change further at later stages [Fig. 3F, columns d–e]. These results indicate that scnRNA production from different chromosomal locations indeed occurs at distinct conjugation stages before 3 h post-mixing and between 3 and 4.5 h post-mixing.

This different timing of scnRNA production from distinct chromosomal locations partially explains the variable densities of TWI1 knockout scnRNAs mapped to different MDS fragments [Fig. 3F, column a]. However, different MDS fragments were variably mapped even by the scnRNAs from the EMA1 knockout cells at late stages of conjugation [Fig. 3F, columns d–e], when scnRNA production was expected to be completed, indicating that, although scnRNA production occurs from a variety of chromosomal locations in the Mic, it is not homogeneous at either the local (i.e., IES-biased) [Fig. 3A–E] or global [Fig. 3F] level.

Sequence repetitiveness in the genome does not fully explain IES-biased scnRNA production

The IES-biased production of scnRNAs may be caused by preferential production of scnRNAs from repeated sequences in the genome because some IESs are related to transposon-derived sequences and are repeated in the Mic genome (Yao and Gorovsky 1974; Wuitschick et al. 2002; Fillingham et al. 2004), and scnRNAs are processed from dsRNA precursors (Malone et al. 2005; Mochizuki and Gorovsky 2005; Noto et al. 2010), which can be more frequently formed by hybridization of transcripts from repeated sequences. To evaluate this possibility, we classified the annotated Mic genome sequences into three classes according to their repetitiveness. All possible 25-nt sequences from the draft Mic genome were sorted to three classes: unique ("Unique" in Fig. 4A), moderately repeated (two to 10 times in the genome, "Mid-Rep" in Fig. 4A), and highly repeated (>10 times, in the genome) ("High-Rep" in Fig. 4A), which comprise 86.9%, 10.0%, and 3.1% of the annotated Mic genome, respectively [Fig. 4A]. The first 25-nt sequences of scnRNAs were then mapped to the sequences in each class.

Among the scnRNAs from TWI1 knockout cells at 3 h post-mixing, 52.2%, 29.1%, and 18.6% were complementary to Unique, Mid-Rep, and High-Rep sequences, respectively [Fig. 4E], indicating that scnRNAs are indeed produced from repeated sequences more frequently [Fig. 4, cf. A and E]. However, because half of the scnRNAs were derived from unique sequences, repetitiveness in the genome seemed not necessary for scnRNA production. To determine whether the repeat bias explains the IES-biased production of scnRNAs, we separately analyzed the IES and MDS biases of scnRNA production from the three repetitiveness classes (Unique, Mid-Rep, and High-Rep) [Fig. 4B–H]. scnRNAs complementary to all three repetitiveness classes were strongly biased for IESs [Fig. 4F–H]. Importantly, even scnRNAs complementary to Unique sequences were biased for IESs [Fig. 4, cf. D and H]. Therefore, the IES-biased production of scnRNAs cannot be explained by the repeated nature of IESs.
IES-biased scnRNA production is attributed to biased transcription in the Mic

If the repetitiveness of the sequences in the genome is not the main contributor to IES-biased scnRNA production, what mechanism is then responsible for the bias? One obvious possibility is that Mic transcription, which produces the precursors of scnRNAs, preferentially occurs at IESs. To investigate this possibility, we analyzed Mic transcription by global run-on sequencing (GRO-seq), which provides a genome-wide profile of the position and orientation of all transcription-engaged RNA polymerases [Core et al. 2008].

A Mic-enriched (>99.5%) nuclear fraction was prepared by a stepwise differential centrifugation method [Sweet and Allis 2006] from wild-type cells at 2 h post-mixing for use in the GRO-seq assay. We used the cells at 2 h post-mixing because the Mic extensively elongates at meiotic prophase (~3–4 h post-mixing) [Sugai and Hiwatashi 1974], making it technically difficult to prepare an intact Mic at these later stages. Although scnRNA accumulation peaks at ~4 h post-mixing, scnRNA production has already initiated at 2 h post-mixing [see Fig. 7, below; Mochizuki et al. 2002]. Therefore, Mic transcription should have started at this stage.

Figure 5. Nuclear run-on assay of the Mic-enriched nuclear fraction. (A, C) The sequences of nascent transcripts obtained by the global nuclear run-on assay using a Mic-enriched nuclear fraction of wild-type cells at 2 h post-mixing were mapped to the Cal (A) or LMR (C) locus. Numbers [reads per million RNAs [RPM]] of the sequences that uniquely mapped to the sense and antisense strands of the loci are shown in the plus and minus directions, respectively, as histograms with 100-nt bins. The IESs are marked in red. The regions of genes (including exons and introns) that are predicted to be transcribed from left to right and from right to left are labeled as purple and blue boxes, respectively. The reads that are predicted to be derived from unidirectional mRNA transcripts are marked with asterisks. (B, D) The profile of scnRNAs [TWI1 knockout cells, 3 h post-mixing] mapped to the Cal (B) or LMR (D) locus. The same results are also shown in Figure 3, C and E, respectively. (B, D, bottom) The fraction of unique sequences that can be mapped by scnRNAs or GRO-seq reads in each 100-nt bin is shown as “Mappability”.

Nuclear run-on transcripts were purified and sheared to ~30- to 250-nt fragments, a cDNA library was constructed, single-read sequences were produced, and the first 25-nt sequences were used for mapping [see the Materials and Methods for details]. Although we used a Mic-enriched nuclear fraction, ~9% of the reads matched rRNAs [Supplemental Table S2]. Because rDNA is amplified to ~10,000 copies only in the Mac and exists as a single copy in the Mic genome, most of the reads matching rRNAs were likely derived from Mac contamination of the Mic-concentrated nuclear fraction.

The filtered GRO-seq reads were mapped to the Cal locus [Fig. 5A]. Most of the transcripts occurred bidirectionally, and a large fraction was complementary to IESs. The pattern of these transcripts was highly similar to that of scnRNAs from TWI1 knockout cells at 3 h post-mixing [Fig. 5B]. Transcripts exhibiting mapping patterns that were highly similar to the scnRNA profiles were also detected from other loci [Supplemental Fig. S1]. Therefore, we conclude that the bias of scnRNAs toward IESs is mainly attributed to the IES-biased transcription of the precursor RNAs in the Mic.

When the GRO-seq reads were mapped to the LMR locus, most of the transcripts were complementary to IESs, and their pattern was highly similar to the scnRNA
profiles at the same locus [Fig. 5C]. However, overall levels of transcription from this locus were very low. The Mic LMR locus may not be highly transcribed at the stage at which the nuclei were prepared (at 2 h post-mixing) because different chromosomal locations probably produce scnRNAs at distinct times [Fig. 3F].

Some of the transcripts detected by GRO-seq occurred unidirectionally, and the number of these transcripts increased steeply at the 5' ends of predicted ORFs [signals marked with asterisks in Fig. 5A,C; Supplemental Fig. S1A]. These likely represent mRNAs derived from contaminating polyploid Mac because although the Mic-enriched fraction contained <0.5% Macs, the Mac contains ~20 times more DNA than the Mic, and thus some transcripts from Mac DNA were expected to be detected in this study. Consistently, the number of reads mapped to the coding regions was correlated to the RNA expression levels of the corresponding genes detected by microarray hybridization with total RNA extracted from whole cells at the same stage [Miao et al. 2009]. However, we cannot exclude the possibility that these mRNA-type transcripts were derived from the Mic. Nonetheless, they were transcribed unidirectionally and thus should not be processed to scnRNAs.

scnRNAs complementary to IESs are selectively enriched during conjugation

Despite the strong bias of scnRNA production for IESs, a significant fraction of scnRNAs at 3 h post-mixing was complementary to MDSs [Fig. 3A–E], including many protein-coding regions (see Fig. 5B,D; Supplemental Fig. S1B,D). Because scnRNAs are involved in the targeting of heterochromatin formation, which in turn induces DNA elimination [Liu et al. 2004, 2007], the presence of scnRNAs complementary to MDSs could cause the elimination of many essential genes. It has been proposed that this mistargeting of DNA elimination is prevented by the enrichment of scnRNAs complementary to IESs and that this enrichment is mediated by the interactions between scnRNAs and the Mac genome [Fig. 1K, Mochizuki et al. 2002; Mochizuki and Gorovsky 2004c; Aronica et al. 2008]. This proposed process is called scnRNA selection.

To determine whether scnRNA selection occurs at a genome-wide level, we compared scnRNAs from wild-type cells at 3, 4.5, 6, and 8 h post-mixing. In this analysis, scnRNAs were oxidized prior to cDNA library construction. The oxidation inhibits linker ligation at the first step of cDNA library construction, and thus any oxidation-sensitive RNAs were excluded from the sequencing. Because scnRNAs loaded onto Twi1p are 2'-O-methylated [Kurth and Mochizuki 2009], which prevents the peroxidation oxidation reaction, studying oxidation-resistant scnRNAs allowed us to specifically analyze scnRNAs that were loaded onto Twi1p [HM Kurth and K Mochizuki, in prep.] and committed to the DNA elimination pathway.

First, the sequenced scnRNAs were mapped to the annotated Mic genome sequence described above. Among the scnRNAs from wild-type cells at 3 h post-mixing, when the selective degradation process should have just begun, 13.0% and 84.0% of scnRNAs mapped to the MDSs and IESs of the annotated genome, respectively [Fig. 6A, Supplemental Table S1]. This ratio was similar to the one obtained from nonoxidized scnRNAs of TWI1 knockout cells [15.3% to MDSs and 80.1% to IESs] [Fig 3B, Supplemental Table S1], supporting our expectation that scnRNA selection had largely not begun at 3 h post-mixing.

The fraction of scnRNAs that was complementary to IESs was gradually enriched through conjugation [Fig. 6A; Supplemental Table S1]. At 8 h post-mixing, when the heterochromatin formation that preceded DNA elimination had begun and thus the scnRNA selection was expected to be completed, only 5.6% of the scnRNAs were complementary to MDSs, while 93.1% were complementary to IESs [Fig. 6A, “WT_8 h”, Supplemental Table S1]. These results are consistent with our previous Southern blot-based observations that total scnRNAs extracted from wild-type cells at 2 h post-mixing hybridized about three times more to Mic DNA than to Mac DNA, while scnRNAs at 8 h post-mixing hybridized ~22 times more to Mic DNA than to Mac DNA [Mochizuki and Gorovsky 2004a]. Therefore, we conclude that scnRNAs complementary to IESs are enriched during conjugation at a genome-wide level.

Enrichment of IES-complementary scnRNAs was also detected by analyzing scnRNAs from wild-type cells without oxidation [Supplemental Fig. S2]. This is not surprising because almost all scnRNAs in cells at the mid to late stages of conjugation [after ~6 h post-mixing] are loaded onto Twi1p [HM Kurth and K Mochizuki, in prep.]. Therefore, oxidation-resistant scnRNAs and total scnRNAs during the later stages of conjugation should mostly overlap. This result also indicates that the periodate oxidation reaction does not perturb the IES/MDS ratio of the cDNA libraries.

A similar scnRNA selection process was detected by mapping scnRNAs to the Cal and LMR loci [Fig. 6B,C]. Like the scnRNAs from TWI1 knockout cells [Fig. 3B–E], the majority of oxidation-resistant scnRNAs from wild-type cells at 3 h post-mixing were complementary to IESs, and only a smaller fraction of scnRNAs mapped to MDSs [Fig. 6B,C, “3h”]. While scnRNAs complementary to IESs persisted through conjugation, those complementary to MDSs were gradually diminished [Fig. 6B,C, peaks marked by pink arrows], and at 8 h post-mixing, most scnRNAs were complementary to IESs [Fig. 6B,C, “8h”]. Therefore, scnRNA selection occurs at both the genome-wide and local levels.

We previously demonstrated that Ema1p, the Twi1p-associated RNA helicase, is required for the degradation of scnRNAs complementary to some MDSs and for proper DNA elimination [Aronica et al. 2008]. Therefore, we next analyzed scnRNAs from EMA1 knockout cells. Although scnRNAs complementary to MDSs were reduced in wild-type cells during conjugation, as described above [Fig. 6A–C], scnRNAs complementary to MDSs remained at high levels throughout conjugation in EMA1 knockout cells [Supplemental Fig. S3A–C]. Therefore, we conclude that EMA1 is required for the genome-wide process of scnRNA selection.
Although scnRNAs complementary to MDSs were dramatically reduced during conjugation in wild-type cells, such scnRNAs still remained at 8 h post-mixing [Fig. 6A–C], when scnRNA-mediated heterochromatin formation had already begun. We believe that this was caused in part by asynchronicity of conjugation; although asynchronous mating initiation was limited by adding nutrient medium at ~3 h post-mixing, some cells at 8 h post-mixing might have mated for only 5 h. However, it is also possible that some scnRNAs complementary to MDSs remain in cells that are at late conjugation stages. There may be a threshold of scnRNAs required for efficient induction of DNA elimination, and MDS-complementary scnRNAs detected at 8 h post-mixing could be below this threshold.

**Turnover of scnRNAs complementary to MDSs is responsible for scnRNA selection**

Transcription in the Mic and the localization of Dcl1p, which are necessary for scnRNA biogenesis, both exclusively occur at the early stage of conjugation [Sugai and Hiwatashi 1974; Martindale et al. 1985; Mochizuki and Gorovsky 2004b, 2005; Malone et al. 2005]. Therefore, the enrichment of scnRNAs complementary to IESs must be attributed to either selective degradation of scnRNAs complementary to MDSs or Dcl1p-independent selective secondary amplification of scnRNAs complementary to IESs by an RNA-dependent RNA polymerase activity.

If the observed relative increase of scnRNAs complementary to IESs (82.4% at 3 h to 93.1% at 8 h) [Fig. 6A] was solely achieved by their selective secondary amplification of scnRNAs complementary to IESs, the total scnRNAs would have to be amplified ~2.6-fold [(100/C0)82.4]/[(100/C0)93.1]. However, the amount of scnRNA, which was analyzed by gel electrophoresis followed by fluorescent dye staining of RNAs [Fig. 7A], peaked at ~4 h post-mixing and decreased slightly at later conjugation stages. Moreover, we recently found that secondary scnRNA production is limited after scnRNA processing by Dcl1p because scnRNAs from all analyzed stages exhibited the signature of Dicer processing [HM Kurth and K Mochizuki, in prep.]. Therefore, selective amplification of IES-complementary scnRNAs, if any, cannot fully explain the enrichment of IES-complementary scnRNAs.

Our previous Northern hybridization studies have demonstrated that the levels of scnRNAs complementary to two different transposon-like IES elements remain constant during the course of conjugation, while the number of scnRNAs complementary to the MI repeat, a repetitive sequence that occurs in both IESs and MDSs, dramatically decreases at the mid-conjugation stage (Aronica et al. 2008). This is consistent with the observation in this study that the fraction of scnRNAs complementary to both MDSs and IESs of the annotated Mic genome...
complementary to the MO-repeat [MO-rpAS]. As shown in Figure 7B, scnRNAs complementary to the MO repeat were detected at early stages of conjugation [2–4 h post-mixing] and became undetectable at mid to late conjugation stages [after 6 h post-mixing], indicating that these MDS-complementary scnRNAs were produced once and then degraded. In contrast, scnRNAs complementary to the Tlr1-1 probe, which is part of the Tlr1 transposon-related IES element, persisted into the late conjugation stages [Fig. 7C], as observed previously [Aronica et al. 2008]. Taken together, our data indicate that the enrichment of IES-complementary scnRNAs is mainly achieved by selective degradation of scnRNAs complementary to MDSs.

Discussion

In this study, we addressed some of the key issues of the small RNA-directed epigenetic regulation of DNA elimination in *Tetrahymena*. As proposed in previous models [Fig. 1], we found that scnRNAs are produced exclusively from the Mic [Fig. 2] and a variety of Mic chromosomal locations [Fig. 3F] and that *EMA1*-dependent scnRNA selection occurs at a genome-wide level [Fig. 6; Supplemental Figs. S2, S3] by RNA turnover [Fig. 7]. However, in contrast to expectations, this study demonstrates that scnRNA production is strongly biased toward IESs [Fig. 3A–E] and occurs nonhomogeneously from the Mic chromosomes at the global level [Fig. 3F]. We further revealed that IES-biased scnRNA production is not caused mainly by the repetitiveness of IESs [Fig. 4], but is attributed to IES-biased transcription [Fig. 5; Supplemental Fig. S1].

The nonhomogeneous production of scnRNAs may explain position-dependent transgene elimination

The nonhomogeneous pattern of scnRNA production from the Mic genome [Fig. 3F] may address a previously unresolved question about the epigenetic regulation of DNA elimination: the position dependencies of transgene elimination. A neomycin resistance (neo) gene integrated into an MDS locus on a Mic chromosome is occasionally deleted from the genomes of new Macs during successive rounds of conjugation. The efficiency of neo elimination from different Mic loci varies significantly [Yao et al. 2003; Liu et al. 2005; Howard-Till and Yao 2007]. Because scnRNA production from MDSs is generally lower than that from IESs [Fig. 3A–E] and the production of scnRNAs from the Mic genome is not homogeneous [Fig. 3], the variable efficiencies of neo elimination from different Mic loci may be due to the amount of scnRNAs produced from the loci.

It has been proposed that scnRNA-mediated DNA elimination acts as a genome surveillance mechanism against transposons [for review, see Schoeberl and Mochizuki 2011]. However, the nonhomogeneous production of scnRNAs from the Mic genome indicates that scnRNAs cannot target transposons for DNA elimination if they are inserted into a Mic genome locus at which scnRNAs are not highly produced. The insertion of transposons may not immediately activate the DNA elimination pathway. Only after multiple insertions may...
some of the transposons integrate into scnRNA-producing loci, enabling the cells to produce scnRNAs complementary to the transposon to eliminate it. In this way, the nonhomogeneous production of scnRNAs may set a threshold for DNA elimination such that only transposons that can efficiently multiply in the Mic genome are targeted for DNA elimination.

**How does IES-biased scnRNA production occur?**

Our unexpected observation that scnRNA production is heavily biased for IESs at the transcriptional level has raised a new question: How is the IES-biased transcription of scnRNA precursors achieved? Theoretically, the biased transcription could be mediated by specific DNA sequences that reside in or around all >6000 IESs. However, we believe this is unlikely for two reasons. First, different IESs share no obvious consensus sequences. Second, the transcriptional start sites of the Mic transcripts seem to be distributed across broad regions of the genome [Fig. 5; Supplemental Fig. S1; Chalker and Yao 1996]. An alternative mechanism for the biased transcription is that scnRNAs are transported not only to the developing new Mac to target IESs for DNA elimination, but also to the new Mic to mark IES sites for future transcription. In this way, DNA elimination in the new Mac may be epigenetically and transgenerationally controlled not only by the genome contents of the parental Mac through the selective degradation of scnRNAs, but also by those of the grandparental Mac through transcriptional regulation of the Mic.

If the latter hypothesis is true, we predict that [1] a protein other than Twi1p transports scnRNAs to the new Mic because Twi1p does not localize to the Mic at any stage [Mochizuki et al. 2002]; [2] IESs and MDSs in the Mic are differentially marked by some chromatin modifications or chromatin-associated proteins, and this difference is maintained during vegetative growth [one such candidate is histone H3 Lys 27 methylation, which is maintained in the Mic during vegetative growth] [Liu et al. 2007]; [3] an epigenetic “block” of DNA elimination through the ectopic introduction of an IES to the parental Mac [Chalker and Yao 1996] not only inhibits the elimination of the IES in the new Mac, but also inhibits Mic transcription, and thus scnRNA production, from the IES during the conjugation of the progeny. We await future research that would confirm or reject these predictions and reveal how ancestral genomes can epigenetically regulate the behavior of genomes of successive generations.

**Materials and methods**

**Small RNA cDNA library preparation and sequencing**

Wild-type *T. thermophila* strains B2086 and CU428, Chr4 nullisomic strains CU357 and CU367 [Bruins et al. 1983], TW1 complete [germ plus soma] knockouts [Mochizuki and Gorovsky 2004a] strains 20-1 and 20-4, and EMA1 knockout [Aronica et al. 2008] strains 7-17-1 and 8-6-1 were used for total RNA preparation. B2086, CU428, CU357, and CU367 were obtained from the *Tetrahymena* Stock Center at Cornell University. Cell mating was induced by mixing equal numbers of prestarved cells of two strains possessing complementary mating types, and asynchronous mating initiation was limited by adding 1/3 vol of 4× SPP at ~3 h post-mixing. RNA was isolated from 2.5 × 10^6 mating cells at the indicated time points post-mixing with the mirVana miRNA isolation kit [Ambion] or TRIzol reagent [Invitrogen]. For periodate oxidation of small RNAs [Akbergenov et al. 2006], the RNA pellet was dried and resuspended in 17.5 μL of borax buffer (4.375 mM borax, 50 mM boric acid at pH 8.6). Then, 2.5 μL of 0.2 M sodium periodate was added, and the reaction was incubated for 10 min at room temperature in the dark. Two microliters of glycerol was added, the solution was incubated for another 10 min at room temperature in the dark, and RNAs were purified by ethanol precipitation.

cDNA libraries of small RNAs were constructed as described [Hafner et al. 2008] with some modifications. In brief, RNA [with or without periodate oxidation] was mixed with radiolabeled 26- and 32-nt oligoribonucleotides [5'-GUGC GCCGG GAUAGGUU AACACUGU-3', 5'-AUUCUGUGGG UAGC GAAUAUGUUAACATGU-3']; Pmel restriction sites are underlined and fractionated in a sequencing gel, and the gel regions containing the two radiolabeled markers [plus a few bases on both sides] were excised. RNA was recovered from the gel and ligated to a 5'-adenylated 3'adapter oligonucleotide [5'-App-TTCGTATCCGCTTCTCTCGTTG-L-3', where L is the 3'-OH-blocking group; IDT] by T4 RNA ligase 2-truncated or T4 RNA ligase 2-truncated K227Q (New England Biolabs). After gel fractionation and purification as above, the product was ligated to the 5' adapter oligoribonucleotide [5'-GUUCAGAGUUUCACAG UCCGACGAUAC-3'] by T4 RNA ligase [Ambion]. The gel-purified final ligation product was then reverse-transcribed with the Sol-rv primer [5'-CAACGAGAACGCGCATAAGC-3'] and PCR-amplified with the Sol-rv primer and Sol-fw primer [5'-ATGTCATCG CGCACCCACAGCATTCAGTCTACAGTCCAG-3']. After phenol–chloroform extraction followed by ethanol precipitation, the PCR product was digested with Pmel [to cut the size markers] and fractionated by agarose gel electrophoresis, and the undigested PCR product was purified from the gel. The amplified cDNA libraries were mixed with 1/10 [by mass] of 4×X174 phage DNA, and single-read 36-base sequences were generated on the illumina GAII platform, resulting in ~17 million–27 million reads. The 4×X174 phage DNA was included because most scnRNAs begin with 5'U [HM Kurth and K Mochizuki, in prep.]; therefore, sequencing only scnRNAs disturbs the cluster recognition process during sequencing reactions. Raw sequencing data and processed data sets have been deposited at the NCBI Gene Expression Omnibus [http://www.ncbi.nih.gov/geo] as GSE38768.

Following the removal of the adapter sequences from the sequence reads, we removed reads containing any non-A/C/G/T bases; reads from tRNAs, rRNA, snRNAs, and the mitochondrial genome; and reads from mixed cDNA libraries of small RNAs were constructed as described [Hafner et al. 2008] with some modifications. In brief, RNA [with or without periodate oxidation] was mixed with radiolabeled 26- and 32-nt oligoribonucleotides [5'-GUGC GCCGG GAUAGGUU AACACUGU-3', 5'-AUUCUGUGGG UAGC GAAUAUGUUAACATGU-3']; Pmel restriction sites are underlined and fractionated in a sequencing gel, and the gel regions containing the two radiolabeled markers [plus a few bases on both sides] were excised. RNA was recovered from the gel and ligated to a 5'-adenylated 3'adapter oligonucleotide [5'-App-TTCGTATCCGCTTCTCTCGTTG-L-3', where L is the 3'-OH-blocking group; IDT] by T4 RNA ligase 2-truncated or T4 RNA ligase 2-truncated K227Q (New England Biolabs). After gel fractionation and purification as above, the product was ligated to the 5' adapter oligoribonucleotide [5'-GUUCAGAGUUUCACAG UCCGACGAUAC-3'] by T4 RNA ligase [Ambion]. The gel-purified final ligation product was then reverse-transcribed with the Sol-rv primer [5'-CAACGAGAACGCGCATAAGC-3'] and PCR-amplified with the Sol-rv primer and Sol-fw primer [5'-ATGTCATCG CGCACCCACAGCATTCAGTCTACAGTCCAG-3']. After phenol–chloroform extraction followed by ethanol precipitation, the PCR product was digested with Pmel [to cut the size markers] and fractionated by agarose gel electrophoresis, and the undigested PCR product was purified from the gel. The amplified cDNA libraries were mixed with 1/10 [by mass] of 4×X174 phage DNA, and single-read 36-base sequences were generated on the illumina GAII platform, resulting in ~17 million–27 million reads. The 4×X174 phage DNA was included because most scnRNAs begin with 5'U [HM Kurth and K Mochizuki, in prep.]; therefore, sequencing only scnRNAs disturbs the cluster recognition process during sequencing reactions. Raw sequencing data and processed data sets have been deposited at the NCBI Gene Expression Omnibus [http://www.ncbi.nih.gov/geo] as GSE38768.

Following the removal of the adapter sequences from the sequence reads, we removed reads containing any non-A/C/G/T bases; reads from tRNAs, rRNA, snRNAs, and the mitochondrial genome; and reads from mixed 4×X174 phage DNA. Then, sequences from 26- to 32-nt RNAs were used as scnRNA sequences. The first 25 nt of each scnRNA sequence was mapped to the annotated Mic genome, the Cal locus, the LMR locus, or the selected 218 Mac genome scaffolds. After mapping scnRNAs to the Cal or LMR locus, the repetitiveness of the genome was normalized by either dividing the numbers of mapped scnRNAs by their sequence frequencies in the draft Mic genome ("weighted") or excluding scnRNAs that were complementary to more than two positions in the Mic genome ("uniquely mapped").

**Annotation of the draft Mic genome**

The draft Mic genome sequence (version 2, supercontigs) was obtained from the *Tetrahymena* Comparative Sequencing Project [Broad Institute of Harvard and Massachusetts Institute of
GRO-seq

Nuclear fractions were isolated from wild-type strains B2086 and CU428 at 2 h post-mixing as described previously (Sweet and Allis 2006) with some modifications. All fractionation steps were performed at 4°C or on ice. Cells (1.2 × 10^7) were suspended in 100 mL of medium B and blensed for 30 sec at the highest speed (Waring 700G commercial blender). Samples were divided among two 50-mL conical tubes and centrifuged with a swing rotor at 1750g, 2750g, 3300g, and 4200g for 5 min each and twice at 4500g for 10 min. The supernatants of each step were blended as above for 20 sec and then centrifuged at the subsequent speed. Each pellet was washed once in modifed nuclei resuspension buffer [NRB, 50 mM Tris-Cl at pH 8.0, 30% glycerol, 5 mM MgCl₂, 0.1 mM EDTA], resuspended in 200 μL of NRB [per 50-mL tube], and stored at −80°C. A portion of the pellet was removed for DAPI staining, and the ratios between the Mac and Mic and the concentrations of nuclei were examined on a hemocytometer. The Mic-enriched nuclear fraction used for the experiments in Figure 5 and Supplemental Figure S1 consisted of −99.5% Mic and −0.5% Mac, as judged by the sizes of the nuclei.

Mic-enriched nuclear fractions were used to identify nascent transcription by applying the GRO-seq technique (Core et al. 2008) with some modifications. Briefly, ~1.3 × 10^7 nuclei of the Mic-enriched fraction were used for the run-on reaction, which consisted of 500 μM ATP, 500 μM GTP, 500 μM UTP, and 0.33 μM α-32PCTP. After DNase I treatment, the total RNA was isolated with TRIzol LS reagent (Invitrogen) and resuspended in 20 μL of water. Base hydrolysis was performed for 20 min on ice to digest RNA products to ~30–250 nt. The RNAs were treated with DNase I again, and RNA was purified twice through P30 spin columns (Bio-Rad). BrdU-incorporated RNA was enriched on anti-BrdU beads (Santa Cruz Biotechnology) for 2 h at 4°C and resuspended in 21 μL of water. A 20-μL aliquot of the RNA was end-repaired and resuspended in 10.5 μL of water.

cDNA was synthesized as described above in the section “Small RNA cDNA Library Preparation and Sequencing” with some modifications. For 5′ adapter ligation, 10 μL of the end-repaired RNAs was incubated with 1 μL of 50 μM 5′ adapter, 0.5 μL of Ribolock [Fermentas], 2.15 μL of ATP-free T4 RNA ligase buffer, 6 μL of 50% PEG-8000, and 2 μL of T4 RNA ligase 2-truncated K227Q (New England Biolabs) overnight at 16°C. The RNAs were enriched on anti-BrdU beads as above and resuspended in 10 μL of water. For 5′ adapter ligation, 10 μL of 3′ adapter-ligated RNAs was incubated with 1 μL of 50 μM 5′ adapter, 0.5 μL of Ribolock, 2.2 μL of T4 RNA ligase buffer, 1 μL of 10 mM ATP, 6 μL of 50% PEG-8000, and 2 μL of T4 RNA ligase [Ambion] overnight at 16°C. The RNAs were purified again with anti-BrdU beads and resuspended in 10 μL of water.

The adapter-ligated RNA was incubated with 2 μL of 10 μM Sol-rv primer and 1 μL of 10 mM dNTP mix for 5 min at 65°C and for 2 min on ice. Then, 4 μL of 5 × first strand buffer, 1 μL of 0.1 M DTT, 1 μL of Ribolock, and 1 μL of SuperScript III reverse transcriptase [Invitrogen] were added. The mixture was incubated for 30 min at 50°C, for 30 min at 55°C, and for 15 min at 70°C. Then, 5 μL of RNase H [Ambion] was added, followed by incubation for 10 min at 37°C. The cDNA was amplified by subjecting 9.5 μL of the reaction to 17 cycles of PCR with KAPA HiFi Hot Start DNA polymerase [Kapa Biosystems] and Sol-rw and Sol-rv primers. The amplified cDNA library was fractionated in a 2% agarose gel; ~100- to 200-base pair (bp) products were extracted, and single-read 50-base sequences were generated on a HiSeq 2000 platform [llumina] to obtain ~200 million reads. Raw sequencing data and processed data sets have been deposited at the NCBI Gene Expression Omnibus [http://www.ncbi.nlm.nih.gov/geo] as GSE38768. Reads containing any non-A/C/G/T bases and reads derived from rRNAs, tRNAs, snRNAs, and the mitochondrial genome were removed. The first 25 nt of each GRO-seq read was used to uniquely map the representative Mic loci.
RNA staining and Northern hybridization

Total RNA was isolated with TRizol [Invitrogen]. The total RNA from 1 × 10^6 cells was separated in a 15% sequencing gel, and RNAs were stained with GelRed (Gentaur). RNA was then blotted onto a nitrocellulose membrane and probed with 32P-labeled oligo DNA probes as described previously (Aronica et al. 2008). The probes used in this study were Tlr1-1 (Aronica et al. 2008), which is complementary to the Tlr1 transposon-like IES, and MO-repeat (5′-TTTTACCAAAGATAAAGTGCAATTTGAAGATTAGTCCAATTGCTAA-3′; the MO repeat is underlined), which is complementary to the MDS-limited MO repeat. The MO repeat (5′-ACTAATCTCTCAAATTTGACACTTT-3′) was identified as the most repetitive 25-nt sequence that only existed in the annotated MDSs.

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Biased transcription and selective degradation of small RNAs shape the pattern of DNA elimination in *Tetrahymena*

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