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Regulation of alternative polyadenylation by genomic imprinting

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Maternally and paternally derived alleles can utilize different promoters, but allele-specific differences in co-transcriptional processes have not been reported. We show that alternative polyadenylation sites at a novel murine imprinted gene (*H13*) are utilized in an allele-specific manner. A differentially methylated CpG island separates polyA sites utilized on maternal and paternal alleles, and contains an internal promoter. Two genetic systems show that alleles lacking methylation generate truncated *H13* transcripts that undergo internal polyadenylation. On methylated alleles, the internal promoter is inactive and elongation proceeds to downstream polyadenylation sites. This demonstrates that epigenetic modifications can influence utilization of alternative polyadenylation sites.

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Transcription by RNA polymerase II (Pol II) requires multiple linked steps, including the assembly of an initiation complex, promoter release, elongation followed by splicing, polyadenylation (polyA), and dissociation of the polymerase complex from the template DNA. Each step provides an opportunity for the cell to regulate gene expression, either by changing the abundance or availability of mRNAs or by generating variant gene products. Epigenetic modifications to DNA and histones can influence transcription at the initiation stage by altering the accessibility of promoter sequences to initiation complex components (Kass et al. 1997a,b). Following initiation, regions of heterochromatin can subtly impede the progress of an elongating polymerase complex, reducing transcriptional output (Lorincz et al. 2004). However, it is not clear whether epigenetic modifications downstream from a promoter can lead to variant gene

products through alternative polyadenylation or splicing.

Polyadenylation, the addition of multiple adenylyl residues to the 3' end of a newly synthesized transcript, confers stability and is required for nuclear export (Huang and Carmichael 1996; Jacobson and Peltz 1996). This occurs following cleavage of the newly synthesized pre-mRNA, typically 15–30 nucleotides downstream from a conserved hexamer motif (usually AAUAAA or AUUAAA) (Proudfoot and Brownlee 1976; Zhao et al. 1999). A large proportion of human genes utilize more than one polyadenylation (polyA) site (Tian et al. 2005), making alternative polyadenylation a major source of transcriptional diversity. The mechanisms governing alternative polyA site selection in mammalian cells have been studied in detail at only a small number of loci; notably the immunoglobulin heavy chain (Takagaki and Manley 1998) and calcitonin/*CGRP* genes (Lou et al. 1998). In both cases, polyA site selection is cell type-specific and is regulated by changes in the concentration of diffusible RNA processing factors (Lou et al. 1998; Takagaki and Manley 1998).

Imprinted genes are differentially expressed on maternally and paternally derived alleles and are estimated to comprise 1%–2% of the mammalian transcriptome (Barlow 1995; Wood and Oakey 2006). These genes are typically clustered into regions of the genome containing multiple imprinted transcripts, which share common *cis*-regulatory elements. Such elements undergo differential CpG methylation when passing through the male and female germline. Allele-specific markings are maintained during embryonic development and can influence transcription of adjacent genes via *cis*-acting regulatory RNAs such as *Air* (Sleutels et al. 2002) or insulator factors such as CTCF (Hark et al. 2000).

Imprinted loci represent an attractive model for studying epigenetic mechanisms of transcriptional control, as both active and silent alleles are present in the same cellular environment. In the presence of an identical complement of *trans*-regulatory factors, allelic differences in transcription must be the result of epigenetic regulation. While the allele-specific use of alternative promoters has been well documented at imprinted loci (Blagitko et al. 2000; Holmes et al. 2003), allele-specific differences in cotranscriptional events such as splicing and polyadenylation have not been reported previously.

We undertook a computational screen for imprinted genes, identifying a novel imprinted domain on mouse Chromosome 2 containing at least two protein-coding genes: *H13* and *Mcts2* (Wood et al. 2007). *Mcts2* is a duplicate of the X-linked *Mcts1* gene that originated by retrotransposition of an mRNA from the X-linked locus into the fourth intron of *H13* (Wood et al. 2007). Unlike most retrotransposed gene copies, *Mcts2* has maintained the capacity to encode a protein and shares 94% amino acid identity with the X-linked paralog >65 million years after the gene duplication event. The promoter of *Mcts2* is associated with a CpG island that acquires DNA methylation in the female germline, and the retrogene is expressed primarily from paternally derived alleles in the developing embryo (Wood et al. 2007).

We identify five polyA sites for *H13* and show that at least three are utilized in an allele-specific manner. This is likely to depend on allele-specific DNA methylation

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at the internal *Mcts2* promoter, which is established during maternal gametogenesis and serves to promote functional *H13* products from maternally derived alleles in the embryo. These findings provide strong evidence that epigenetic modifications can influence alternative polyadenylation, and pave the way for future studies to determine precisely how this influence can be exerted.

Results and Discussion

Alternative polyadenylation at the *H13* locus

H13 encodes signal peptide peptidase; an intramembrane aspartic protease responsible for the dislocation of signal peptides from the endoplasmic reticulum membrane (Weihsen et al. 2002). The ORF is spread over 12 exons (Fig. 1A) and encodes a protein consisting of seven transmembrane helices (Fig. 1B). EST evidence suggests a number of transcript variants. Northern blots (Fig. 1C) and 5' and 3' RACE were performed in material extracted from whole brain tissues of neonatal mice, revealing one promoter and five distinct polyA sites (Fig. 1A). Al-

though alternative *H13* promoters cannot be absolutely excluded, all transcript species identified by Northern analysis (Fig. 1C) can be accounted for by different 3' RACE products. Three polyA sites are clustered within a single block of 3' untranslated region (UTR) situated 20 kb downstream from the monoexonic *Mcts2* gene (*H13a-c*). The alternatively skipped penultimate exon of the 2-kb *H13a* transcript contains a premature stop codon that removes a C-terminal KKXX membrane retention signal from full-length signal peptide peptidase proteins, altering subcellular localization patterns (Urny et al. 2006). Two internal polyA sites lie 0.5 kb (*H13d*) and 3 kb (*H13e*) upstream of the *Mcts2* transcriptional start site (Fig. 1A). Biochemical characterization of the protein product of *H13* has previously demonstrated that the catalytic residues essential for signal peptide peptidase activity lie within the region of the ORF downstream from the *H13d* and *H13e* polyA sites (Weihsen et al. 2002). Hence, only transcripts extending beyond *Mcts2* and the CpG island (*H13a-c*) can encode functional signal peptide peptidase proteins (Fig. 1B). *H13d*

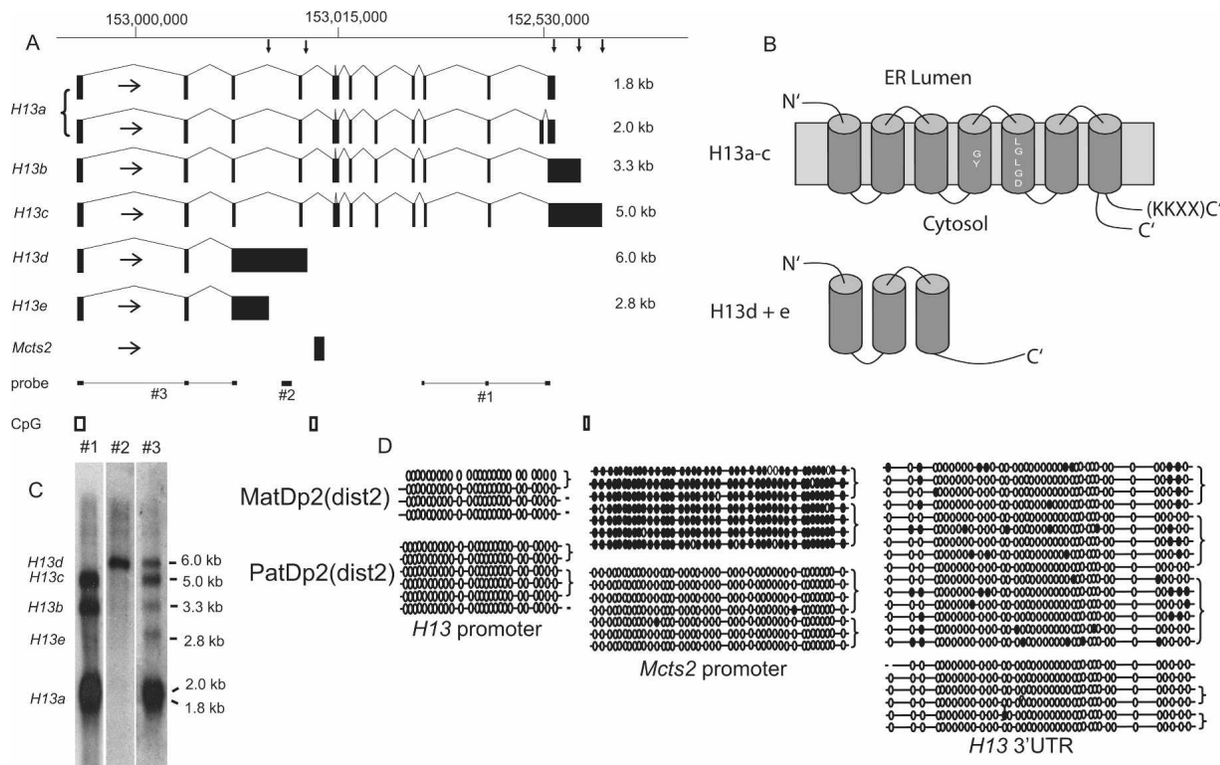


Figure 1. Gene structure and CpG methylation at the murine *H13/Mcts2* locus. (A) Transcript map of *H13* isoforms, determined by 5' and 3' RACE, Northern, and GenBank EST evidence. The bar indicates nucleotide position on mouse Chromosome 2 [Mm. build 37]. Representative GenBank accession numbers are as follows: *H13a* 1.8 kb, AJ345032; *H13a* 2.0 kb, AK049101; *H13b*, NM_010376; *H13c*, F830002G13; *H13d*, 4021402J09; *H13e*, AK004690; *Mcts2*, NM_025543. Exons are black rectangles and splice patterns are indicated. Vertical arrows indicate polyA sites, and horizontal arrows show the direction of transcription. The greater abundance of *H13a* prevented the simultaneous detection of *H13a*, *H13b*, and *H13c* in a single 3' RACE reaction. Consequently, we did not determine whether the alternatively skipped penultimate exon is also included in *H13b* or *H13c* transcripts. The positions of Northern probes for *C* are shown at the bottom, as are the three CpG islands at which methylation was investigated in *D*. (B) Predicted structure of proteins translated from *H13* transcript isoforms. Cylinders represent transmembrane (TM) helices. Residues critical for signal peptide peptidase activity are indicated within transmembrane helices 4 and 5. Distinct C termini for *H13a-c* protein isoforms arise from the inclusion or exclusion of a penultimate exon (see the text). Membrane topology for the larger protein was determined by Weihsen et al. (2002). The proteins putatively translated from *H13d* and *H13e* have not been described previously; hence, membrane topology is not shown. (C) Northern blots performed with neonatal brain mRNA identify transcripts utilizing all five polyA sites identified by 3' RACE. (D) DNA methylation at three CpG islands: the *H13* promoter, the *Mcts2* promoter, and the *H13* downstream. Horizontal lines denote individual DNA strands on which filled circles represent methylated CpG dinucleotides; open circles are unmethylated CpGs. Genomic DNA was extracted from whole brain of neonatal mice carrying uniparental partial duplications for distal Chromosome 2 [MatDp(dist2) and PatDp(dist2)].

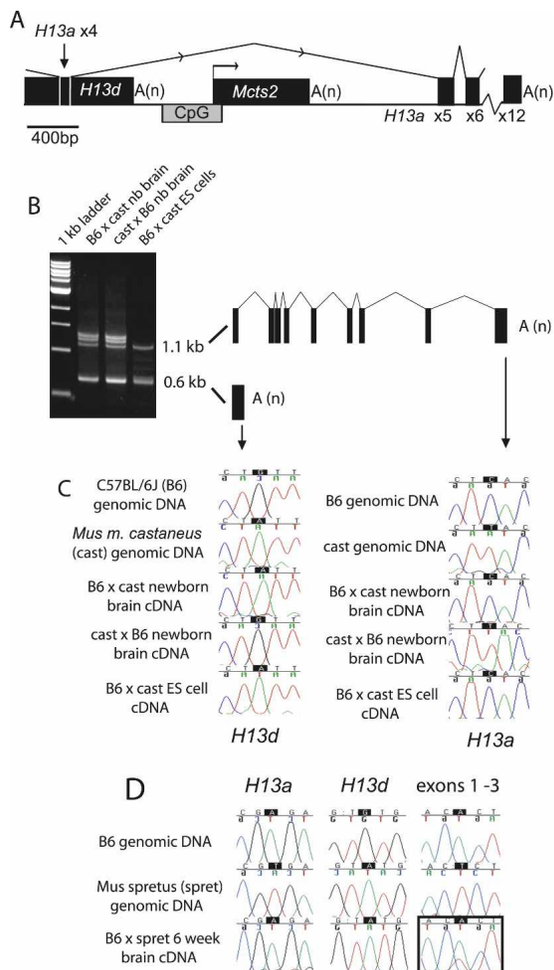


Figure 2. Allele-specific utilization of *H13* polyA sites. (A) Enlarged schematic showing the fourth intron of *H13* including *Mctc2*, the differentially methylated CpG island, and the *H13d* polyA site. Exons are black rectangles, and splice patterns are indicated with arrows depicting the direction of transcription. The fourth exon of *H13a-c* is shown entirely contained within the 3'UTR of *H13d*. The 3' RACE primer used in *B* anneals within this region. (B) 3' RACE was performed using a gene-specific primer within the fourth exon of *H13a-c*, in material from F1 hybrids of the C57BL/6J (B6) and *Mus musculus castaneus* (cast) substrains. The major products utilized either the *H13a* (1.1 kb) or *H13d* (0.6 kb) polyA sites. Products corresponding to *H13b* and *H13c* are not seen, presumably due to their lower abundance and larger size reducing the efficiency of amplification. Additional splice variants of *H13a* are seen in whole newborn brain but not in ES cells. In all figures, the maternal strain is given first for hybrid material. (C) RACE products were sequenced over regions containing SNPs between B6 and cast alleles. For whole-brain material, three biological replicates were performed with similar results (data not shown). (D) RT-PCR products covering exons 1–3 (Fig. 1A, probe #3) originate from both parental alleles. No SNP was identified between B6 and cast, so B6 × *Mus spretus* (spret) material was used. The black box indicates the hybrid sequence trace showing biallelic expression.

and *H13e* have the potential to encode a protein of unknown function, consisting of the three N-terminal transmembrane helices of signal peptide peptidase with 51 unique C-terminal residues (Fig. 1B).

Allele-specific CpG methylation

Three CpG islands lie in the region: one at the intronic *Mctc2* promoter, another at the *H13* promoter, and a

third separating the *H13b* and *H13c* polyA sites (Fig. 1A). To investigate the allele-specific methylation profile of these three regions, we sequenced bisulphite-modified genomic DNA from whole neonatal brains of mice carrying uniparental partial disomies for distal Chromosome 2 [MatDp(dist2) and PatDp(dist2) methods]. At the *Mctc2* promoter, methylation is present on maternally derived alleles but absent on paternally derived alleles (Fig. 1D). We have shown previously that this region undergoes methylation during oogenesis but is unmethylated in sperm (Wood et al. 2007), suggesting that the somatic methylation pattern is inherited from the gametes. The CpG island at the *H13* promoter is predominantly unmethylated on both alleles in neonatal brain (Fig. 1D), as is the CpG island downstream from the *H13c* polyadenylation site. It is not presently clear whether transcription initiates from the 3' CpG island.

Allele-specific polyadenylation in intersubspecies hybrid mice

The differentially methylated region (DMR) at the *Mctc2* promoter lies ~200 base pairs (bp) downstream from the *H13d* polyA site (Fig. 2A), and ~3.5 kb downstream from *H13e* (Supplemental Fig. 1). The proximity of these two sites to the DMR raised the possibility that they could be differentially utilized on maternally and paternally derived alleles. To address this, 3' RACE was performed in neonatal brain material from intersubspecies hybrid mice. Following gel purification, 3' RACE products were sequenced over regions containing SNPs between the parental strains, revealing the relative utilization of each polyA site by each parental allele (Fig. 2B,C; Supplemental Fig. 1). Transcripts terminating upstream of the *Mctc2* DMR (*H13d* and *H13e*) originate preferentially from paternally derived alleles (Fig. 2C; Supplemental Fig. 1). In contrast, the *H13a* polyA site downstream from the DMR is utilized predominantly on maternally derived alleles (Fig. 2C). This is observed in cDNA prepared from whole neonatal brains and from embryonic stem (ES) cells (Fig. 2C). RT-PCR amplification of exons 1–3 yields products that originate from both parental alleles (Fig. 2D), consistent with biallelic transcriptional initiation from the unmethylated *H13* promoter (Fig. 1C). The allele-specific utilization of polyA sites within a single cell type indicates that polyadenylation can be regulated by epigenetic mechanisms.

Disrupted polyadenylation following uniparental inheritance

To confirm these findings in an independent genetic system, polyadenylation at *H13* was studied in mice carrying uniparental partial disomies for distal Chromosome 2. This allowed the expression of *H13* transcripts to be compared in mice that inherit two copies of the *H13* locus from their mothers [MatDp(dist2)] versus mice that inherit two paternal copies of this genomic region [PatDp(dist2)]. Northern blots were generated from total brain RNA and hybridized with a probe to the common 5' exons (Fig. 1A, probe #3). This confirmed the preferential maternal expression of transcripts utilizing the *H13a* and *H13c* polyA signals, but *H13b* transcripts were not detectable on this genetic background [(C3H × 101)F1] (Fig. 3A). Transcripts utilizing both the *H13d* and *H13e* polyA signals are present at greater levels following pa-

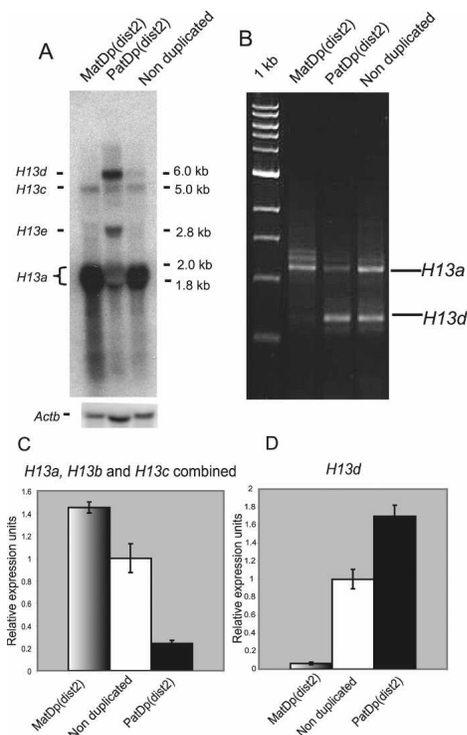


Figure 3. Disrupted polyadenylation in mice carrying uniparental partial disomies. (A) Northern generated with total RNA extracted from neonatal brains of mice carrying uniparental partial duplications for distal Chromosome 2. Nonduplicated mice are either homozygous, heterozygous, or noncarriers of the T11H translocation but in all cases will inherit distal Chromosome 2 biparentally. The blot was hybridized with probe #3 from Figure 1A. One biological replicate was performed. Note that *H13a* transcripts are not completely absent following paternal inheritance, but *H13d* transcripts are not detected in *MatDp(dist2)*. (B) Qualitative 3' RACE assay, using the same gene-specific primer employed in Figure 2B. Two biological replicates gave similar results (data not shown). (C) Combined expression of transcripts extending through the DMR (*H13a-c*) in neonatal brains measured by QRT-PCR. Data are presented as ratios of the mean *Ct* value in relation to the mean *Ct* value for β -actin for each genotype. Microarray expression data confirmed that this endogenous control gene did not change significantly between *MatDp(dist2)* and *PatDp(dist2)* samples (Supplemental Fig. 2). *Ct* values are transformed as described (Pfaffl 2001) and are presented in relation to nonduplicated controls that are set to a value of 1. Amplifications were performed in quadruplicate ($n = 4$). Error bars represent the standard error of the mean. The amplified region spanned exons 6 and 7. (D) Expression of *H13d* transcripts measured by QRT-PCR. The amplified region lies within 500 bp of the *H13d* polyA site. Data are presented as in C.

ternal versus maternal inheritance. Qualitative 3' RACE assays conducted from *MatDp(dist2)* and *PatDp(dist2)* neonatal brain material further demonstrate that uniparental transmission results in disrupted 3' end formation (Fig. 3B).

Quantitative RT-PCR (QRT-PCR) assays were designed to quantify allele-specific differences in the utilization of polyA sites at *H13*. Collectively, *H13a*, *H13b*, and *H13c* transcripts are generated at approximately sevenfold greater levels from maternal versus paternal alleles (Fig. 3C). The *H13d* transcript is produced at ~30-fold greater levels following paternal versus maternal inheritance (Fig. 3D). These results are consistent with semiquantitative microarray measurements over the same exonic regions (Supplemental Fig. 2).

Collectively, these data indicate that polyA sites situated 0.2 kb (*H13d*) and 3 kb (*H13e*) upstream of the *Mcts2* DMR are utilized primarily on unmethylated paternal alleles. Maternal alleles preferentially generate *H13* transcripts that elongate through the methylated and transcriptionally inactive *Mcts2* promoter to the three polyA sites located ~20 kb downstream (*H13a-c*).

Alternative polyadenylation is controlled by maternal germline methylation

To further investigate the role of allele-specific methylation in this phenomenon, embryos lacking maternally derived methylation imprints were generated by crossing *Dnmt3l*^{-/-} mothers with *cast* fathers (Bourc'his et al. 2001). Heterozygous offspring lack allele-specific DNA methylation at the *Mcts2* promoter (Fig. 4A), and this is associated with activation of the normally silent maternal allele (Fig. 4B). Qualitative 3' RACE for the *H13* gene demonstrates that loss of intronic methylation is associated with increased utilization of the *H13d* polyA site in *Dnmt3l*^{-/+} embryos (Fig. 4C). This is entirely consistent with *PatDp(dist2)* individuals, which also exhibit biallelic hypomethylation at this region (Fig. 3; data not shown). Despite up-regulation of *H13d* relative to *H13a*

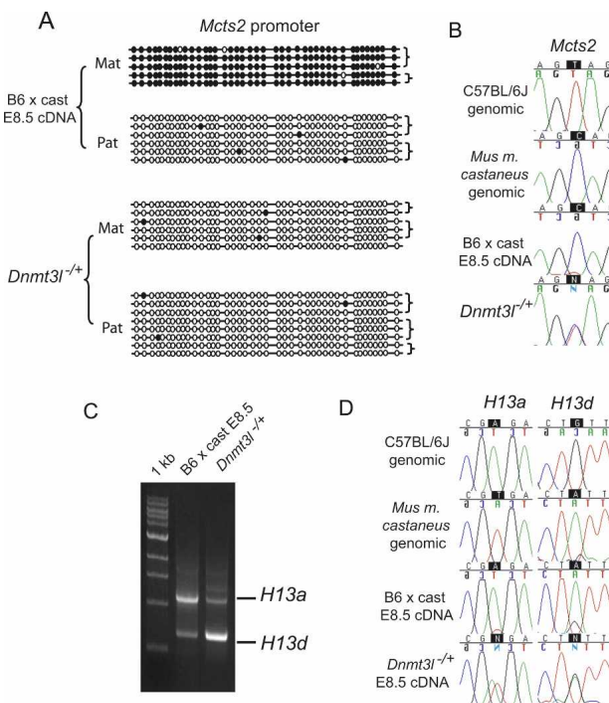


Figure 4. Disrupted polyadenylation in maternal imprint-free embryos. (A) Methylation status of the *Mcts2* CpG island in pooled embryos ($n = 5$) derived from *Dnmt3l*^{-/-} dams and wild-type cast sires (*Dnmt3l*^{-/+}) and control embryos, determined by sequencing bisulphite-modified genomic DNA. Parental strands were distinguished on the basis of a SNP between *Mus domesticus* (B6) and *Mus musculus* (*cast*) alleles. Filled circles are methylated, and closed circles are unmethylated. Strands derived from the same PCR amplification are joined by braces on the right. (B) Allele-specific RT-PCR sequencing assay for *Mcts2* in *Dnmt3l*^{-/+} hybrid embryos (pooled, $n = 5$). (C) Qualitative 3' RACE in cDNA from pooled *Dnmt3l*^{-/+} hybrid embryos. (D) 3' RACE products from C were excised and sequenced over regions containing SNPs between the two parental strains.

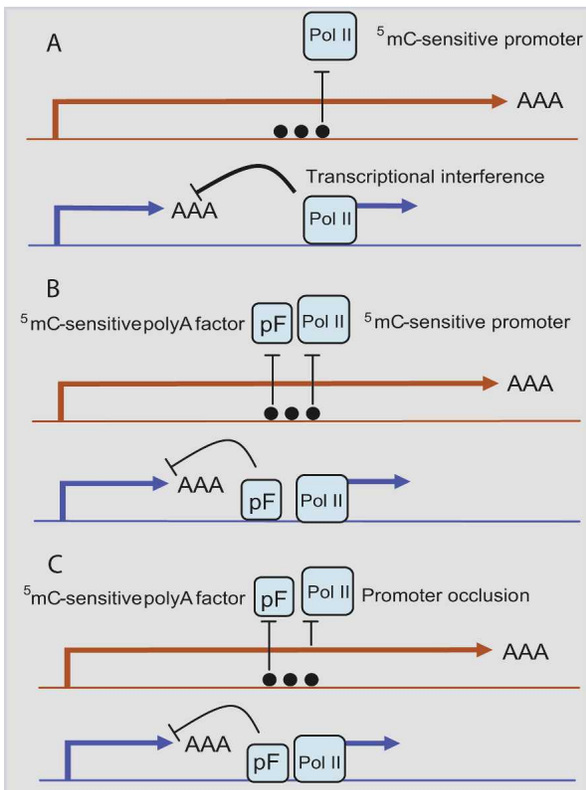


Figure 5. Models for epigenetic regulation of polyadenylation at *H13*. Maternal alleles are shown in red, and paternal are in blue. Filled circles represent CpG methylation (5mC), and Pol II represents the RNA polymerase II initiation complex. pF denotes a hypothetical polyadenylation factor(s). (A) Methylation-sensitive formation of transcriptional initiation complexes at the *Mcts2* promoter results in transcriptional interference specifically on paternally derived alleles. In this model, transcription factors bound at the *Mcts2* promoter act indirectly as polyadenylation factors. (B) Methylation-sensitive binding of transcriptional initiation complexes and distinct polyadenylation factors within the DMR. In this model, allele-specific utilization of the *H13d + e* polyA sites and *Mcts2* transcription occur independently and do not require common factors. (C) Methylation prevents the binding of polyadenylation factors on maternally derived alleles, and the elongation of *H13* transcripts through the maternal *Mcts2* promoter brings about silencing by promoter occlusion (Shearwin et al. 2005).

in the *Dnmt3l*^{-/-} embryos, both products were generated at levels sufficient to assess their allele-specific expression pattern by sequencing. The *H13a* and *H13d* polyA sites show reciprocal allele-specific biases in wild-type embryonic day 8.5 (E8.5) embryos but are utilized at approximately equal levels on each allele in *Dnmt3l*^{-/-} individuals (Fig. 4D). Thus, alternative polyadenylation in the embryo is controlled by DNA methylation marks that are established during oogenesis. Furthermore, biallelic hypomethylation in *Dnmt3l*^{-/-} embryos and *PatDp*(dist2) neonates decreases the output of functional signal peptide peptidase transcripts (*H13a-c*) (Figs. 4C, 3C).

In this report, we identified multiple polyA sites for the murine signal peptide peptidase gene (*H13*) and showed that at least three are utilized in an allele-specific manner. Previous *in vivo* studies of alternative polyadenylation mechanisms in mammals have focused on tissue-specific variation, which can be explained on the basis of tissue-specific differences in the concentra-

tion of *trans*-regulatory factors. Such explanations are insufficient to account for polyA site selection at the *H13* locus, as maternally and paternally derived alleles in ES cells are potentially exposed to the same array of diffusible molecules. Hence, allele-specific epigenetic differences must play a role. Consistent with this conclusion, a differentially methylated CpG island and promoter were identified in the region separating maternally and paternally utilized polyA sites. Hypomethylation within this region correlates with the use of polyA sites 0.5 and 4 kb upstream, whereas hypermethylated alleles preferentially utilize sites between 20 and 25 kb downstream. Several possible explanations could account for these observations, three of which are outlined in Figure 5.

On paternally derived alleles, silencing of *H13a-c* is not complete. Transcriptional output was approximately sevenfold lower relative to maternal alleles (Fig. 3C), but still readily detectable by Northern blot (Fig. 3A). In a previous report, we examined the allele-specific expression of *H13a-c* (exons 3–12) using qualitative methodology and found some degree of expression from the paternal allele in whole embryo and placental tissues (Wood et al. 2007). It will be important to determine whether these observations reflect cell type-specific imprinting or low level expression from the paternal allele across all cell types of these complex tissues. Silencing of *H13d* on maternal alleles is more absolute, showing 30-fold lower expression levels than paternal alleles in neonatal brain.

In summary, we identified a novel mechanism of imprinted gene regulation acting at the level of alternative polyadenylation. The imprinted *H13* locus has the potential to provide novel insights into this important transcriptional phenomenon. Further work is now required to elucidate the precise mechanism by which identical DNA sequences can generate alternatively polyadenylated transcripts within a single cellular environment.

Materials and methods

Tissue sources

Mice with uniparental partial disomies (or uniparental duplications) were generated by intercrossing heterozygotes for the T(2:9)11H reciprocal translocation (Cattanach and Beechey 1997). Mice carrying maternal duplications for distal Chromosome 2 *MatDp*(dist2) also carry paternal duplications for distal Chromosome 9, and vice versa. Maternal imprint-free embryos were generated as described previously (Bourc'his et al. 2001). RNA from the hybrid ES cell line used in Figure 2 was a gift from Wolf Reik.

DNA methylation

RNA and DNA were prepared using the RNeasy and DNeasy kits (Qiagen) and quantified using an Agilent 2100 bioanalyser. Bisulphite conversion was performed as described (Wood et al. 2007). All strands presented in Figures 1 and 4 differed either in epigenotype or in the presence of unconverted cytosines, or were derived from separate PCR amplifications. All strands showed >95% conversion.

Gene expression

5' and 3' RACE-ready cDNAs were synthesized using the SMART RACE kit (Clontech), with 0.5 μ g of polyA⁺ mRNA per 100 μ L of final volume. Details of all RACE and PCR primers, cycling conditions, and SNPs are listed in the Supplemental Material. Northern blots were performed using either 10 μ g of total or 2 μ g of polyA⁺ RNA. Microarrays (Affymetrix u430_v2) were hybridized according to standard Affymetrix protocols, and data were analyzed using the GCOS/MAS5 algorithm.

QRT-PCR was performed using custom TaqMan assays designed by Applied Biosystems. cDNA was prepared using the SuperScript II kit (Invitrogen) primed with oligo dT. Three microliters of total RNA were used for a 50- μ L final RT reaction volume, of which 2 μ L were used per 10 μ L of QPCR reaction. Expression of *H13* isoforms was measured relative to

β -actin, which did not change significantly between MatDp(dist2) and PatDp(dist2) samples measured by microarray (Supplemental Fig. 2). Sample values were determined using the relative expression ratio mathematical model (Pfaffl 2001), with *Ct* values transformed into expression ratios according to the formula

$$\text{Ratio} = 2^{\left[\left(\text{control}_{\text{sample}} - \text{target}_{\text{sample}} \right) - \left(\text{control}_{\text{baseline}} - \text{target}_{\text{baseline}} \right) \right]}$$

where $\text{target}_{\text{baseline}}$ denotes the mean of the *Ct* values measured for the target gene in the baseline sample, $\text{control}_{\text{baseline}}$ is the equivalent value for the endogenous control gene, and $\text{target}_{\text{sample}}$ and $\text{control}_{\text{sample}}$ are the mean *Ct* values determined in the nonbaseline sample.

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