1 2	KRAB zinc finger protein diversification drives mammalian inter-individual methylation variability
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18 19	Major classification:
20	Biological Sciences
21	
22	Minor classification:
23	Genetics
24	
25	Keywords:
26	DNA methylation, endogenous retrovirus, KRAB zinc finger proteins, metastable epiallele,
27	VM-IAP
28	
29	
30	

31 Abstract

Most transposable elements (TEs) in the mouse genome are heavily modified by DNA methylation 32 and repressive histone modifications. However, a subset of TEs exhibit variable methylation levels 33 34 in genetically identical individuals and this is associated with epigenetically-conferred phenotypic differences, environmental adaptability, and transgenerational epigenetic inheritance. The 35 evolutionary origins and molecular mechanisms underlying inter-individual epigenetic variability 36 37 remain unknown. Using a repertoire of murine variably methylated intracisternal A-particle (VM-IAP) epialleles as a model, we demonstrate that variable DNA methylation states at TEs are 38 highly susceptible to genetic background effects. Taking a classical genetics approach coupled 39 40 with genome-wide analysis, we harness these effects and identify a cluster of KRAB zinc finger 41 protein (KZFP) genes that modifies VM-IAPs in *trans* in a sequence-specific manner. Deletion of the cluster results in decreased DNA methylation levels at these variably methylated alleles. An 42 analysis of ChIP-seq and RNA-seq datasets generated from KZFP-cluster mutants reveals that the 43 44 observed DNA methylation changes are accompanied by altered histone modifications and by dysregulation of neighbouring genes. We find that VM-IAPs cluster together phylogenetically and 45 46 that this is linked to differential KZFP binding, suggestive of an ongoing evolutionary arms race 47 between TEs and this large family of epigenetic regulators. These findings indicate that KZFP 48 divergence and concomitant evolution of DNA binding capabilities are mechanistically linked to 49 methylation variability in mammals, with implications for phenotypic variation and putative paradigms of mammalian epigenetic inheritance. 50

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54 Significance statement

Transposable elements (TEs) are repetitive sequences with potential to mobilise, causing genetic diversity. To restrict this, most TEs in the mouse are heavily epigenetically modified by DNA methylation. However, a few TEs exhibit variable methylation levels that differ between individuals and confer an epigenetic, rather than genetic, influence on phenotype. The mechanism underlying this remains unknown. We report the identification of a polymorphic cluster of KRAB zinc finger proteins (KZFPs) responsible for the epigenetic properties of these variably methylated TEs, with deletion of the cluster profoundly influencing their DNA methylation and expression of adjacent genes. We propose that rapid KZFP divergence underlies variable epigenetic states in mammals, with implications for epigenetically conferred phenotypic differences between individuals within and across generations.

75 Main text

76 Introduction

Complex genetic interactions contribute to evolutionary fitness, phenotypic variation, and disease risk. This is highlighted by comparative research across inbred mouse strains showing that genetic background not only influences basic fitness traits such as litter size and sperm count, but also modulates the penetrance and expressivity of numerous gene mutations (1, 2). Despite the extensive documentation of strain-specific epistatic effects in the mouse and their important implications for mechanistic insight and experimental reproducibility, the underlying modifier genes remain uncharacterised in most cases.

Studies on foreign DNA insertions in the mouse genome demonstrate that modifier genes can act via epigenetic pathways to drive genetic background-dependent phenotypes. A number of transgenes show predictable strain-specific DNA methylation patterns that are associated with transgene expression levels (3–6). Similar effects have been reported on the methylation state of endogenous retroviruses (ERVs), as exemplified by the MusD ERV insertion Dac^{1J} which is methylated in mouse strains that carry the unlinked *Mdac* modifier gene (7, 8). In strains lacking the *Mdac* allele, Dac^{1J} is unmethylated and the mice exhibit limb malformation.

Another example is provided by the *Agouti viable yellow* (A^{vy}) metastable epiallele, where a spontaneously inserted intracisternal A-particle (IAP) element influences the expression of the downstream coat-colour gene *Agouti* (9). IAPs are an evolutionarily young and highly active class of ERV (10). Variable DNA methylation of the A^{vy} IAP is established early in development across genetically identical mice and is correlated with a spectrum of coat colour phenotypes, which in turn display transgenerational inheritance and environmental sensitivity (11–13). Both the distribution and heritability of A^{vy} phenotypes are influenced by genetic background (14–16). 98 Therefore, the identification and characterisation of the responsible modifier genes can provide 99 insight into the mechanisms governing the early establishment of stochastic methylation states at 100 mammalian transposable elements.

101 We recently conducted a genome-wide screen for individual variably methylated IAPs 102 (VM-IAPs) in the C57BL/6J (B6) inbred mouse strain (17, 18). The screen yielded a robust set of experimentally validated regions to use as a model to investigate inter-individual epigenetic 103 104 variability. Most VM-IAPs belong to the IAPLTR1_Mm and IAPLTR2_Mm subclasses. Approximately half of them are full-length IAPs with an internal coding region flanked by near-105 identical long terminal repeats (LTRs); the other half are solo LTRs. While solo LTRs lack 106 107 autonomous retrotransposition potential, they are rich in regulatory sequences and thus have the ability to affect host gene expression. As observed for $A^{\nu y}$, methylation variability is re-established 108 at VM-IAPs from one generation to the next regardless of parental methylation states. Importantly, 109 VM-IAP methylation levels are consistent across all tissues of a single mouse, suggesting that 110 111 individual-specific methylation states are acquired in early development prior to tissue differentiation. The inter-individual variability suggests that the establishment of VM-IAP 112 113 methylation levels involves an early stochastic phase.

Here, we introduce genetic variation to the study of VM-IAPs. We report that half of the IAPs found to be variably methylated in B6 are present in 129 sub-strains while the vast majority are absent from the CAST/EiJ (CAST) genome. We find that a subset of the shared loci between B6 and 129 display variable methylation in both stains; the remainder are hypermethylated in 129. Further methylation quantification in reciprocal B6 x CAST F1 hybrids reveals pervasive maternal and zygotic genetic background effects. Through backcrossing and genetic mapping experiments, we identify a cluster of KRAB zinc finger proteins (KZFPs) on Chromosome 4 responsible for the strain-specific *trans*-acting hypermethylation of multiple B6 VM-IAPs. We show that deletion of the KZFP cluster leads to a decrease in DNA and H3K9 methylation, an increase in H3K4 trimethylation, and alterations in nearby gene expression at the targeted VM-IAPs. A phylogenetic sequence analysis demonstrates that genetic sequence plays a crucial role not only in the targeting of VM-IAPs by strain-specific KZFPs, but also in the establishment of methylation variability in a pure B6 context. Based on our findings, we propose that KZFP diversification is at the centre of the mechanism leading to variable epigenetic states within and across mouse strains.

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129 **Results**

130 VM-IAPs exhibit strain-specific methylation states

131 To determine whether B6 VM-IAPs are variably methylated in other inbred mouse strains, we first 132 catalogued their presence or absence in the 129S1/SvlmJ (129) and CAST strains based on a previous analysis of polymorphic ERVs (19). The classification was verified, and at times 133 corrected, by visually assessing each locus in the 129 and CAST reference genomes (20). Out of 134 51 experimentally validated B6 VM-IAPs (18), 25 of the IAPs are present in 129 and 3 are present 135 136 in CAST (Fig. 1A). These numbers are consistent with our previous work showing that VM-IAPs are evolutionarily young IAPs (17), and were expected given the evolutionary relationship between 137 these three strains: B6 and 129 are classical inbred laboratory strains derived from several 138 139 subspecies while CAST is wild-derived and evolutionarily more distant.

We compared the methylation level of 19 loci conserved between B6 and 129 by bisulphite pyrosequencing. The probed cytosine-guanine dinucleotide (CpG) sites are comparable across loci and located at the most distal end of the 5' LTR of each element, close enough to the bordering

unique sequence to ensure amplification of a single product. As expected, all 19 regions exhibited 143 methylation variability across inbred B6 mice (i.e. more than 10% variability across individuals). 144 145 In contrast, only eight loci were variably methylated in 129 mice (Fig. 1B). Most of these displayed distinct methylation ranges compared to those observed in B6. The remaining 11 IAPs lacked 146 inter-individual variability and are therefore not VM-IAPs in the 129 strain (Fig. 1B). For the most 147 148 part, these elements were highly methylated, akin to the vast majority of the ~10,000 IAPs in the mouse genome. The susceptibility of VM-IAPs to genetic background effects provides an 149 opportunity to map the genetic determinants of inter-individual methylation variability. 150

Due to the repetitive nature of IAPs, it is difficult to rule out the possibility that the differences 151 152 in methylation between B6 and 129 are a result of sequence divergence within the elements themselves rather than a consequence of *trans*-acting modifiers. For example, a LINE element is 153 embedded in IAP-Rab6b in the 129 genome that is absent in the B6 genome (SI Appendix, Fig. 154 S3A). To avoid this confounder, we implemented a hybrid breeding scheme using B6 and CAST 155 156 mice (Fig. 2A). Because B6 VM-IAPs are largely absent from the CAST genome, F1 hybrid offspring inherit a single allele from their B6 parent. This property allowed us to assess whether a 157 158 haploid CAST genome is capable of inducing methylation changes at B6-specific VM-IAPs in 159 trans. Maternal and paternal transmission of these alleles was followed by crossing B6 females to 160 CAST males (BC) and CAST females to B6 males (CB), respectively. The reciprocal design 161 enabled the exploration of parent-of-origin effects in addition to genetic background effects for the 12 B6-specific VM-IAPs examined in this experiment. Furthermore, we used large sample sizes 162 163 to guarantee the detection of subtle shifts in the distribution of methylation levels at each locus, which additionally revealed that the frequency distributions of VM-IAP methylation levels in the 164

pure B6 population form skewed bell curves rather than normal distributions (*SI Appendix*, Fig.S1).

Two thirds of the assessed B6-specific VM-IAPs showed significant differences between BC 167 and CB methylation distributions (Fig. 2B and SI Appendix, Fig. S2). These effects were not 168 reciprocal, indicating they were not imprinting effects. For instance, at half of the loci, CB hybrids 169 showed hypermethylation of the paternally inherited B6 VM-IAP while the BC hybrids exhibited 170 171 levels comparable to pure B6. This suggests the presence of a CAST-specific maternally inherited modifier acting on the paternally inherited B6 allele (Fig. 2B and SI Appendix, Fig. S2). Paternal 172 transmission of the CAST modifier had no effect on the maternally inherited B6 VM-IAP, 173 174 consistent with a maternal effect. Additional experiments are required to better understand these 175 maternal genetic background effects, but strain-specific factors derived from the oocyte are likely involved. 176

177 In addition to genetic background-specific maternal effects, a subset of VM-IAPs exhibited 178 zygotic genetic background effects, defined as changes in VM-IAP methylation caused by the introduction of a CAST haploid genome regardless of parental origin. Four VM-IAPs displayed 179 180 significant shifts in methylation when BC and CB were compared to the B6 population (Fig. 2B 181 and SI Appendix, Fig. S2). IAP-Marveld2 was alone in showing a reduction in methylation in F1 hybrids compared to B6 individuals. The other three (IAP-Rab6b, IAP-Sema6d, and IAP-Fam78b) 182 were hypermethylated in F1 hybrids compared to B6 mice, suggesting that CAST-encoded 183 184 modifier(s) may be targeting these loci for repression. We note that IAP-Rab6b and IAP-Sema6d displayed both maternal and zygotic genetic background effects, while IAP-Gm13849 and 185 IAP-Slc15a2 displayed neither. The range of responses indicates that the mechanisms influencing 186 variable methylation at IAPs are not common across all loci. 187

188 Strain-specific IAP-*Rab6b* hypermethylation is driven by a single modifier locus

A successful genetic mapping experiment relies on an unambiguous phenotype. Unlike most of the VM-IAPs examined in hybrids, IAP-*Rab6b* (a solo LTR) exhibited non-overlapping B6 and F1 hybrid methylation distributions that were clearly distinguishable using a 60% methylation threshold (Fig. 2*B*). Due to the categorical nature of this 'methylation phenotype', IAP-*Rab6b* was selected to identify VM-IAP modifiers using B6/CAST hybrids.

We first investigated whether the low methylation state (<60%) could be rescued in a subsequent generation by backcrossing F1 hybrids to B6 mice. Low methylation was reacquired in approximately half of N1 backcrossed offspring, irrespective of parental origin (Fig. 3*A* and *SI Appendix*, Fig. S3*B*). This is indicative of limited redundancy in IAP-*Rab6b*-targeting modifiers in the CAST genome. The segregation of methylation states was not attributable to IAP-*Rab6b* copy number, as hemi- and homozygous individuals were represented in both the highly and lowly methylated groups (*SI Appendix*, Fig. S3*C*).

We conducted an additional round of B6 backcrossing to further characterise the inheritance 201 202 pattern of IAP-Rab6b methylation states. N2 offspring generated from highly methylated N1 males recreated the 1:1 ratio of high-to-low methylation observed in N1 offspring, while N2 offspring 203 generated from lowly methylated N1 males were all lowly methylated (Fig. 3A and B). This 204 205 Mendelian inheritance pattern indicates that a single dominant CAST-derived locus causes the hypermethylation of B6-derived IAP-Rab6b in trans. This was confirmed by crossing highly 206 methylated N2 males to B6 females, once again producing roughly equal numbers of highly and 207 lowly methylated N3 offspring (Fig. 3A and B). 208

We next mapped the modifier locus using the Giga Mouse Universal Genotyping Array (GigaMUGA), a 141,090 single nucleotide polymorphism (SNP) microarray designed to capture

the genetic diversity found across mouse strains (21). Due to the evolutionary distance between 211 B6 and CAST, a majority of the probed SNPs are informative between the two strains. DNA 212 samples from 47 N3 individuals were analysed on the array. SNP calls were filtered to identify 213 heterozygous SNPs shared by all 23 highly methylated individuals and absent from all 24 lowly 214 methylated individuals. The resulting SNPs all mapped to a 7.3 Mb interval on distal Chromosome 215 216 4 (Fig. 3C; SI Appendix, Fig. S4; Dataset S1: Table S1). We separately analysed 22 N2 individuals using the lower-resolution MiniMUGA array and independently identified the same genomic 217 region (Dataset S1: Table S2). Combining both mapping experiments delimited a 6.4 Mb window 218 on Chromosome 4 containing the IAP-Rab6b modifier(s) (GRC38/mm10, chr4:141,964,197-219 148,393,136). Of note, this locus was found to exhibit high heterozygous SNP density in a study 220 comparing the genomes of sixteen different laboratory mouse strains (20). 221

Assessment of the genes within the mapped interval revealed a cluster of KZFPs (Fig. 3D). 222 Present in the hundreds, KZFPs make up the largest and most diverse transcription factor family 223 224 in higher vertebrate genomes (22, 23). They are best known for their role in sequence-dependent transposable element repression. Their C₂H₂ zinc finger arrays recognize and bind DNA motifs 225 226 with high specificity and their KRAB domain recruits the scaffold protein KAP1, which in turn 227 induces heterochromatin (24). The rapid evolutionary expansion of murine KZFPs sets them apart 228 from other more conserved epigenetic regulators. We therefore hypothesised that this KZFP cluster, designated Chr4-cl, contains the strain-specific modifier(s) of IAP-Rab6b. The B6, CAST, 229 and 129 variants of this cluster are henceforth referred to as Chr4-cl^{B6}, Chr4-cl^{CAST}, and Chr4-cl¹²⁹, 230 231 respectively.

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234 The CAST-derived modifier locus targets multiple VM-IAPs in a sequence-specific manner

IAP sequences are highly repetitive in the mouse genome due to the evolutionary youth and 235 retrotransposition potential of IAP elements (10). In view of the sequence-specificity of KZFP-236 237 induced epigenetic repression, we reasoned that VM-IAPs with sequence similarity to IAP-Rab6b may also be targeted by the same modifier locus. Six solo LTR VM-IAPs with more than 90% 238 sequence identity to IAP-Rab6b were selected as potential targets along with IAP-Sema6d and 239 240 IAP-Fam78b, which had exhibited hypermethylation in F1 hybrids (Fig. 2B). Methylation was quantified in N2 individuals, half of which were highly methylated at IAP-Rab6b (i.e. 241 heterozygous carriers of the CAST modifier locus) and half of which were lowly methylated at 242 IAP-Rab6b (i.e. non-carriers of the CAST modifier locus). We found that individuals that were 243 highly methylated at IAP-*Rab6b* were also highly methylated at six out of the eight assessed loci 244 - IAP-Tmprss11d, IAP-Pink1, IAP-Rps12, IAP-Trbv31, IAP-Ect2l, and IAP-Sema6d - and vice 245 versa for the lowly methylated individuals (Fig. 4A). This result suggests that these regions are 246 247 additional modifier targets. In contrast, IAP-Gm20110 and IAP-Fam78b methylation levels were not concordant with IAP-Rab6b methylation levels (Fig. 4A). A sequence alignment of the nine 248 249 solo LTR VM-IAPs revealed a single region that distinguished IAP-Gm20110 and IAP-Fam78b 250 from the other six IAPs (Fig. 4B and SI Appendix, Fig. S5). The 28 bp DNA segment, containing 251 an insertion and various SNPs in IAP-Gm20110 and IAP-Fam78b, is a likely binding site for the CAST-specific modifier. 252

The cross-locus comparison highlights the sequence dependence of the modifiers of these epialleles and provides support for a KZFP-mediated mechanism. Interestingly, our earlier observations in pure 129 mice showed variable methylation at IAP-*Gm20110* and IAP-*Fam78b*, and hypermethylation at the other six IAPs (Fig. 1*B*), suggesting that Chr4-cl¹²⁹ shares VM-IAP modifier allele(s) with Chr4-cl^{CAST} that are absent from Chr4-cl^{B6}.

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259 The KZFP cluster on Chromosome 4 modifies VM-IAP methylation states

The unique clustered organisation of KZFPs in the mouse genome stems from segmental duplications, resulting in high sequence similarity among adjacent KZFPs and low-quality cluster reference sequences (22). To circumvent the technical difficulties and potential functional redundancy associated with generating single-KZFP knockouts (KOs), we examined the consequences of deleting the entire Chr4-cl using a previously generated Chr4-cl KO mouse line (25).

We first assessed DNA methylation effects caused by the loss of Chr4-cl in a pure B6 genetic 266 background. Compared to wild type (WT) mice, which exhibited the expected inter-individual 267 methylation variability at all loci, Chr4-cl KO mice showed significantly lower methylation levels 268 at IAP-Rab6b, IAP-Pink1, IAP-Ect2l, and IAP-Rps12 (Fig. 4C). The effect was particularly 269 pronounced at IAP-Rab6b, where all KO mice were completely unmethylated. This result shows 270 that Chr4-cl^{B6} is necessary for the acquisition of variable methylation at IAP-*Rab6b* and reveals 271 an important mechanistic role for KZFPs in the stochastic methylation of retrotransposons. Of 272 note, the other VM-IAPs targeted by the CAST-specific modifier did not show a reduction in 273 methylation in the absence of Chr4-cl^{B6}. Given the extensive redundancy displayed KZFPs in the 274 mouse genome (25), it is possible that the variable methylation observed at these regions in B6 275 mice is conferred by KZFP(s) located in other cluster(s). 276

We next asked whether Chr4-cl can mediate the strain-specific hypermethylation of VM-IAPs 277 using the 129 Chr4-cl locus. Homozygous B6 Chr4-cl KO mice were crossed to WT 129 mice, 278 which harbour Chr4-cl¹²⁹ as well as most VM-IAPs of interest (Fig. 1A, Fig. 4D). F1 mice were 279 backcrossed to 129 followed by two rounds of heterozygous intercrosses (Fig. 4D). VM-IAP 280 methylation was assessed in the resulting Chr4-cl KO and WT mice of mixed B6/129 genetic 281 282 background. In this instance, all of the predicted Chr4-cl targets from our cross-locus comparison in Fig. 4A exhibited significantly lower methylation levels in Chr4-cl KO mice compared to their 283 WT counterparts, often reflecting a return to the variable levels observed at these regions in pure 284 B6 mice (Fig. 4*E*). WT methylation levels were largely consistent with the pure 129 data from Fig. 285 1B despite the use of different 129 sub-strains. These results indicate that Chr4-cl is the 286 functionally relevant segment of the 6.4 Mb interval identified in our mapping experiment and 287 288 demonstrate that KZFPs are strain-specific VM-IAP modifiers.

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290 Loss of Chr4-cl alters the chromatin and transcriptional landscape near targeted VM-IAPs

291 The recruitment of KAP1 and subsequent H3K9 trimethylation by the methyltransferase SETDB1 are characteristic of epigenetic silencing by KZFPs. To gain insight into the mechanism by which 292 strain-specific Chr4-cl KZFP(s) target VM-IAPs, we analysed previously generated ChIP-seq 293 294 datasets that profiled histone modifications in Chr4-cl WT and KO embryonic stem (ES) cells of mixed B6/129 background (25). Visual inspection of H3K9me3 ChIP-seq tracks at Chr4-cl-295 targeted VM-IAPs revealed a modest decrease in H3K9me3 enrichment upon loss of Chr4-cl 296 (Fig.5A and SI Appendix, Fig. S6A and B). More striking, however, was a marked increase in 297 H3K4me3 in KO cells at Chr4-cl targets, with levels equivalent to those observed at neighbouring 298 299 gene promoters (Fig. 5A, C and SI Appendix, Fig. S6A). No increase in H3K4me3 was observed

at non-targets IAP-*Gm20110* and IAP-*Fam78b* (Fig. 5*B*). Therefore, the acquisition of H3K4me3
 represents the default chromatin state at these loci, which is partially impeded either directly or
 indirectly by Chr4-cl KZFPs in early development.

The H3K4me3 mark is associated with transcriptional activity and localises to gene promoters, with greatest enrichment in the region immediately downstream of the transcription start site (TSS) (26). In line with this, the increase in H3K4me3 in Chr4-cl KO cells at targeted VM-IAPs was exclusively found at their 3' end, downstream of the TSS embedded in the solo LTRs (Fig. 4*B*, Fig. 5*A*, *C* and *SI Appendix*, Fig. S6*A*).

308 Next, we explored whether the remodelled chromatin landscape at VM-IAPs in Chr4-cl KO 309 ES cells was associated with altered expression of neighbouring genes. RNA-seq datasets 310 generated from the same Chr4-cl WT and KO ES cells revealed differences in gene expression near IAP-Pink1 and IAP-Rab6b. Pink1 and Rab6b, located 1 kb upstream and 3 kb downstream of 311 312 IAP-Pink1 and IAP-Rab6b, respectively, were upregulated in Chr4-cl KO ES cells (but only Pink1 313 reached statistical significance) (Fig. 5D and SI Appendix, Fig. S6C). Slco2a1, located 20 kb upstream of IAP-Rab6b, was significantly downregulated in Chr4-cl KO ES cells. These data 314 315 indicate that the loss of Chr4-cl results in a range of transcriptional disruptions. While 316 transcriptional changes were not observed near the other Chr4-cl-targeted VM-IAPs, our analysis does not rule out longer range transcriptional effects. 317

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319 Methylation variability at IAPLTR2_Mm elements is sequence dependent

The seven VM-IAPs that we identified as Chr4-cl targets are all solo LTRs of the IAPLTR2_Mm subclass. To determine how VM-IAPs compare to other solo LTRs of this subclass from an

evolutionary perspective, we built a neighbour-joining tree of all solo IAPLTR2 Mm elements in 322 the B6 genome. Consistent with a KZFP-mediated mechanism, we found that VM-IAPs of this 323 324 subclass cluster together phylogenetically (Fig. 6A). This is in agreement with our previous analysis on IAPs of the IAPLTR1_Mm subclass (17) and reinforces the concept that genetic 325 sequence is instructive in the establishment of inter-individual methylation variability. We selected 326 327 five epigenetically uncharacterised IAPs in the VM-IAP-enriched subtree to test whether members of this clade are in fact unidentified VM-IAPs. All five candidates failed to display methylation 328 variability, highlighting that other determinants such as genomic context likely also play a role in 329 the acquisition of methylation variability at IAPs (Fig. 6B). Given that murine IAPs are almost 330 invariably highly methylated, it follows that the VM-IAP-enriched subtree has escaped epigenetic 331 repression, at least partially. Notably, this clade was enriched in H3K4me3 in WT ES cells and 332 showed a greater increase in H3K4me3 in Chr4-cl KO ES cells compared to other IAPLTR2_Mm 333 solo LTRs (Fig. 7A). 334

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Chr4-cl KZFPs target IAPLTR2_Mm elements

To determine whether Chr4-cl KZFPs are capable of recognising IAPLTR2_Mm elements, the 337 IAPLTR2_Mm consensus sequence was queried for binding motifs previously assigned to 16 338 339 Chr4-cl KZFPs (25). Two of the query hits, ZFP989 and Gm21082, exhibited ChIP-seq enrichment at IAPLTR2_Mm solo LTRs (Fig. 7B). Both ZFP989 and Gm21082 appear to bind the same region 340 of the LTR and are thus likely the product of a gene duplication event within Chr4-cl. Gm21082 341 was previously reported to target IAPLTR2 Mm elements along with one other KZFP, ZFP429, 342 which is located in a KZFP cluster on Chromosome 13 (Chr13-cl) (25). Interestingly, ZFP989 and 343 344 Gm21082 showed reduced enrichment at VM-IAPs compared to other IAPLTR2_Mm solo LTRs,

whereas ZFP429 exhibited strong preferential binding at IAPs in the VM-IAP-enriched subtree(Fig. 7*B*).

It is feasible that the CAST and 129 alleles of ZFP989 or Gm21082 are responsible for the 347 strain-specific hypermethylation of VM-IAPs, while ZFP429 may be involved in the establishment 348 of inter-individual methylation variability in B6 mice. This would explain why DNA methylation 349 at some of the VM-IAPs targeted by Chr4-cl¹²⁹ were unaffected by the loss of Chr4-cl in a pure 350 351 B6 background (Fig. 4C). We note that while we have shown that multiple KZFPs are capable of binding LTRs of the IAPLTR2_Mm subtype, none of the candidate Chr4-cl KZFPs appear to 352 recognise the predicted binding site identified in Fig. 4E. This is unsurprising considering that the 353 354 B6 allele(s) of the Chr4-cl KZFP modifier(s) are not expected to strongly bind VM-IAPs, and the ChIP-seq datasets used for this analysis were generated through stable expression of epitope-355 tagged B6 KZFPs. 356

357

358 Discussion

Variable methylation of murine IAPs across genetically identical individuals was reported more 359 360 than two decades ago (27), yet the underlying mechanisms and evolutionary origins of this phenomenon have remained elusive. In this study, we identify widespread genetic background-361 specific modification of VM-IAPs and exploit these to investigate the genetic determinants of 362 mammalian epigenetic stochasticity. We demonstrate that a polymorphic KZFP cluster on 363 Chromosome 4 promotes the sequence- and strain-specific hypermethylation of multiple VM-IAPs 364 in *trans*, the loss of which alters the chromatin and transcriptional landscape of the modified loci 365 and their surrounding genetic environment. We expect our classical genetics approach using inbred 366 367 mouse strains to be generalisable to other variably methylated regions in the mouse genome.

The identification of Chr4-cl KZFP(s) as strain-specific VM-IAP modifier(s) is consistent 368 with the literature documenting KZFPs as products of rapidly evolving genes with critical 369 370 functions in transposable element repression (reviewed in Ecco et al., 2017). The large number of species-specific KZFPs in the mouse compared to most other higher vertebrates suggests that 371 murine KZFPs have undergone particularly rapid amplification (23). It is thought that this 372 373 expansion reflects an active evolutionary arms race following ERV invasion events (22, 28). Our finding that Chr4-cl contains strain-specific modifier(s) of certain IAP elements indicates that 374 murine KZFPs are evolving rapidly enough to detect significant divergence within the mouse 375 species with important epigenetic and transcriptional ramifications. IAPs, which are murine-376 specific and represent the most mutagenic ERV class in the mouse (10), have likely played a major 377 role in this process. Thus, comparative research across mouse strains is uniquely suited to the study 378 of KZFP gene evolution. 379

A significant technical hindrance in taking full advantage of cross-strain mouse genetics in 380 381 this context relates to the extensive redundancy exhibited by KZFPs both within and across clusters (22, 23, 25). The current mouse strain reference genomes have large gaps in KZFP clusters, 382 rendering a cross-strain comparison of Chr4-cl sequences currently unfeasible (20). In fact, while 383 384 we expect significant KZFP sequence differences in Chr4-cl across mouse strains, we 385 acknowledge that we have not excluded the possibility that the strain-specific effects we have 386 identified are driven by differences in KZFP gene regulation rather than allelic variation of KZFPs themselves. Indeed, a recent study showed that the NOD and B6 mouse strains exhibit differential 387 388 T cell gene expression and 3D chromatin organisation in Chr4-cl, with implications for diabetes 389 phenotypes (29). The improvement of genetic engineering tools for repetitive gene families and the generation of high-quality mouse strain reference genomes with full coverage over KZFP
 clusters will be crucial in addressing this issue.

We have shown that in addition to mediating the strain-specific hypermethylation of 392 393 VM-IAPs, KZFPs play an important role in the establishment of inter-individual methylation variability in a pure B6 background, as evidenced by the complete loss of DNA methylation at 394 IAP-Rab6b in B6 Chr4-cl KO mice. The specificity of KZFP binding relies on four amino acids 395 396 within each zinc finger, so mutations in these key residues or in the DNA sequence of their binding 397 sites have important implications for target site binding kinetics (30). We propose that stochastic methylation arises when VM-IAP sequences are weakly recognised by KZFPs during early 398 399 preimplantation development. Consistent with this model, a large number of murine KZFPs are 400 highly expressed in ES cells, including most of the KZFPs in Chr4-cl (22, 23, 25, 31). Furthermore, we previously documented a lack of methylation co-variation across VM-IAPs within an 401 individual mouse (17), which is in agreement with low-affinity binding interactions occurring 402 403 independently between a KZFP and multiple VM-IAP targets. The phylogenetic clustering of VM-IAPs provides further support for this mechanism given that KZFP function is driven by DNA 404 405 sequence recognition. It is noteworthy that ZFP429 preferentially binds solo LTRs in the VM-IAP-406 enriched subtree compared to other IAPLTR2_Mm solo LTRs, perhaps reflecting a host adaptive 407 response to elements that have escaped epigenetic repression. Alternatively, it is possible that interindividual methylation variability is an early sign of transposable element domestication. 408

While this framework predicts that the sequence of an IAP and that of its KZFP modifier(s) are the prime drivers of inter-individual methylation variability, additional factors are expected to influence the probability of a binding event occurring. This is illustrated by the presence of highly methylated IAPLTR2_Mm elements within the VM-IAP-enriched subtree. Potential influencing factors include chromatin accessibility of VM-IAP insertion sites and the number and expression level of KZFPs targeting a particular locus. Importantly, we envisage that other IAP-binding proteins interfere with binding kinetics between KZFPs and VM-IAPs. In fact, VM-IAPs are enriched for CCCTC-binding factor (CTCF), a methylation-sensitive DNA binding protein that may act as an antagonist to methylation-promoting KZFP modifiers (17, 18).

The most prominent difference in chromatin structure that we observed between Chr4-cl WT 418 419 and KO ES cells was a substantial increase in H3K4me3 in Chr4-cl KO cells at the 3' end of targeted VM-IAPs. This suggests that KZFP binding in early development prevents the 420 accumulation of H3K4me3 at VM-IAP TSSs, potentially enabling subsequent DNA methylation. 421 422 In line with this, the ADD domain of *de novo* DNA methyltransferases DNMT3A and DNMT3B 423 (and of their cofactor DNMT3L) specifically binds unmethylated H3K4 (32, 33). In the absence of modifying KZFPs, other transcription factors and histone modifying enzymes have increased 424 access to VM-IAP sequences, which may in turn contribute to the dysregulation of VM-IAP-425 426 neighbouring genes. Interestingly, a recent study using the BXD recombinant inbred mouse panel 427 identified six major *trans*-acting dominant suppressors of H3K4me3 in male germ cells, all of 428 which were mapped to KZFP clusters (34).

Our work is consistent with previous research on strain-specific modifiers (35). A growing number of ERV-derived mouse mutations whose phenotypic penetrance is dependent on genetic background have been associated with modifier genes located in an interval on Chromosome 13 containing a KZFP cluster (Chr13-cl). This region harbours *Mdac*, modifier of the dactylaplasiacausing *Dac*^{1J} insertion, as well as modifiers of IAPs shown to mediate a range of strain-specific phenotypes (8, 36–38). Moreover, two polymorphic KZFPs in Chr13-cl, SNERV1 and SNERV2, were recently found to influence ERV expression in lupus-susceptible mouse strains (39).

Interestingly, the Chr4-cl KZFP Zfp979 is responsible for the strain-specific methylation of the 436 HRD transgene (40), indicating that polymorphic Chr4-cl KZFPs target a variety of foreign DNA 437 438 sequences. Notably, the genes whose expression we found to be altered in Chr4-cl KO mice near Chr4-cl-targeted VM-IAPs have been implicated in human disease. Mutations in *PINK1*, which 439 codes for a kinase involved in mitochondrial quality control, are associated with Parkinson's 440 441 disease (41); mutations in *SLCO2A1*, a prostaglandin transporter, cause chronic enteropathy (42). It is possible that Chr4-cl KZFP diversification influences related susceptibilities in mice. Taken 442 together, a picture emerges of KZFPs as fine-tuners of evolution within the mouse species, 443 whereby KZFP divergence across strains leads to strain-specific epigenetic landscapes with 444 important phenotypic implications. Moreover, it is possible that KZFP divergence across human 445 populations contributes to variable phenotypic outcomes via the differential recognition of 446 endogenous (and perhaps even exogenous) retroviruses. 447

We have shown that most VM-IAPs are susceptible to maternal effects and posit that these 448 449 too are driven by strain-specific KZFPs. A number of KZFPs have been characterised as maternaleffect genes expressed in the oocyte (43-45). Maternally-derived ZFP57, for example, is required 450 451 for the maintenance of DNA methylation at imprinted regions during global post-fertilisation 452 methylation erasure (43). It is intriguing that, on a B6 background, A^{vy} and IAP-Gm13849 only 453 exhibit epigenetic inheritance upon maternal transmission (12, 17). We speculate that certain paradigms regarded as mammalian transgenerational epigenetic inheritance may in actual fact be 454 the product of post-fertilisation re-targeting of epigenetic states by germline-derived KZFPs. 455

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459 Materials and Methods

All mouse work was conducted in compliance with the Animals (Scientific Procedures) Act 1986 Amendment Regulations 2012 following ethical review by the University of Cambridge Animal Welfare and Ethical Review Body (Home Office project license # PC213320E). DNA methylation was quantified using bisulphite pyrosequencing and genetic mapping was carried using the GigaMUGA SNP genotyping array (21). Data availability and details of mouse experiments, molecular techniques, and computational analyses performed in this study are described in *SI Appendix*, Materials and Methods.

467

468 **Supporting information**

- 469 SI Appendix: Materials and Methods; Supplemental Figures S1-S5.
- 470 Dataset S1: Supplemental Tables S1-S6.
- 471 Dataset S2: Unprocessed MiniMUGA SNP calls.
- 472 Dataset S3: Unprocessed GigaMUGA SNP calls.
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475 Acknowledgements

- 476 This study was supported by the Wellcome Trust, United Kingdom (210757/Z/18/Z to A.C.F.-S.),
- 477 the BBSRC, United Kingdom (BB/R009996/1 to A.C.F.-S.), the Medical Research Council,
- 478 United Kingdom (MR/R009791/1 to A.C.F.-S.), the National Institutes of Health, U.S.
- 479 (1ZIAHD008933 to T.S.M.), and by PhD scholarships from the Cambridge Trust and the BBSRC
- to T.M.B. and J.L.E., respectively. We thank Michael Imbeault for valuable advice on our work

and Noah Kessler, Amir Hay, Jessi Becker, and other members of the Ferguson-Smith lab for 481 useful discussions. We are grateful to Jesse Loilargosain for technical assistance, Ali Robinson for 482 mouse husbandry support, Gernot Wolf for communications on NGS datasets, Melania Bruno and 483 Sherry Ralls for sample preparation and shipment, and Anastasiya Kazachenka for preliminary 484 data on VM-IAP methylation levels in 129 mice. 485 486 References 487 488 J. R. Shorter, et al., A diallel of the mouse collaborative cross founders reveals strong 489 1. strain-specific maternal effects on litter size. G3 Genes, Genomes, Genet. 9, 1613–1622 490 (2019). 491 F. Odet, et al., The founder strains of the collaborative cross express a complex 492 2. combination of advantageous and deleterious traits for male reproduction. G3 Genes, 493 Genomes, Genet. 5, 2671–2683 (2015). 494 3. C. Sapienza, J. Paquette, T. H. Tran, A. Peterson, Epigenetic and genetic factors affect 495 transgene methylation imprinting. Development 107, 165–168 (1989). 496 497 4. N. D. Allen, M. L. Norris, M. A. Surani, Epigenetic control of transgene expression and imprinting by genotype-specific modifiers. Cell 61, 853-861 (1990). 498 499 5. P. Engler, et al., A strain-specific modifier on mouse chromosome 4 controls the methylation of independent transgene loci. Cell 65, 939-947 (1991). 500 A. Schumacher, P. A. Koetsier, J. Hertz, W. Doerfler, Epigenetic and genotype-specific 501 6. effects on the stability of de novo imposed methylation patterns in transgenic mice. J. 502 Biol. Chem. 275, 37915-37921 (2000). 503 7. K. R. Johnson, P. W. Lane, P. Ward-Bailey, M. T. Davisson, Mapping the mouse 504 dactylaplasia mutation, Dac, and a gene that controls its expression, mdac. Genomics 29, 505 457-464 (1995). 506 H. Kano, H. Kurahashi, T. Toda, Genetically regulated epigenetic transcriptional 507 8. 508 activation of retrotransposon insertion confers mouse dactylaplasia phenotype. Proc. Natl. Acad. Sci. 104, 19034–19039 (2007). 509 510 9. D. M. J. Duhl, H. Vrieling, K. A. Miller, G. L. Wolff, G. S. Barsh, Neomorphic agouti mutations in obese yellow mice. Nat. Genet. 8, 59-65 (1994). 511 10. L. Gagnier, V. P. Belancio, D. L. Mager, Mouse germ line mutations due to 512 retrotransposon insertions. Mob. DNA 10, 15 (2019). 513 11. G. L. Wolff, R. L. Kodell, S. R. Moore, C. A. Cooney, Maternal epigenetics and methyl 514

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- 626 Figure legends
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628 Fig. 1. Inter-individual methylation variability at IAPs is strain-specific. (A) B6 VM-IAPs are polymorphic across inbred mouse strains. All experimentally validated B6 VM-IAPs (18) were 629 scored for presence (navy blue rectangles) or absence (light blue rectangles) in the 129S1/SvlmJ 630 631 and CAST/EiJ reference genomes. Instances in which a classification could not be made with 632 confidence due to gaps in the reference sequences are shown in white. VM-IAPs are colour-coded according to their structure (full-length IAPs, blue; truncated IAPs, orange; solo LTRs, pink). LTR 633 634 subclass annotation, as defined by RepeatMasker, is indicated above each VM-IAP. VM-IAPs are 635 named based on their closest coding gene. (B) DNA methylation levels in B6 (grey) and 129S2/Sv 636 (purple) inbred mice of IAPs shared between the two strains. Some IAPs exhibit variable 637 methylation (>10 % variance across individuals) in both strains (left of dotted line); others are only variably methylated in B6 mice (right of dotted line). Methylation levels of the distal-most CpGs 638 639 of the IAP 5' LTRs were quantified from genomic DNA using bisulphite pyrosequencing. Each dot represents the average methylation level across CpGs for one individual. 640

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642 Fig. 2. VM-IAP methylation levels are subject to maternal and zygotic genetic background effects (GBEs). (A) Reciprocal hybrid breeding scheme. BC F1 hybrids (green diamonds) were generated 643 by breeding B6 (black) females with CAST (brown) males. CB F1 hybrids (yellow diamonds) 644 645 were produced from the reciprocal cross of CAST females and B6 males. (B) VM-IAPs classified based on their susceptibility to maternal GBEs (upper left), maternal zygotic GBEs (upper right), 646 zygotic GBEs (lower left), or neither (lower right). Violin plots represent the B6 (grey), BC 647 (green), and CB (yellow) F1 offspring methylation distributions. Dotted and dashed lines show the 648 649 distribution quartiles and median, respectively. Faint hollow circles represent individual-specific

methylation levels, quantified from genomic DNA and averaged across the distal CpGs of the
VM-IAP 5' LTR. B6, BC, and CB methylation levels were compared for each VM-IAP using the
Kruskal-Wallis test followed by Dunn's post hoc multiple comparison test (**** p<0.0001; ns:
not significant). Sample sizes are shown below each graph. Graphs for the 8 additional VM-IAPs
analysed in this experiment can be found in *SI Appendix*, Fig. S2.

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Fig. 3. Strain-specific IAP-*Rab6b* hypermethylation is driven by a single dominant modifier locus 656 657 on Chromosome 4. (A) Genetic backcrossing uncovers Mendelian inheritance pattern of 658 IAP-Rab6b methylation states. F1 BC males were backcrossed to B6 females to produce the first backcrossed generation (N1). Three highly methylated (red) and three lowly methylated (grey) N1 659 660 males were backcrossed to B6 females to produce the N2 generation, and highly methylated N2 males were once again backcrossed to B6 females to produce the N3 generation. Average percent 661 662 CAST DNA remaining in the genome at each generation is indicated under the graph. 60% 663 methylation was used as the cut-off value to classify individuals as highly (red) or lowly (grey) methylated. (B) Pedigree illustrating the inheritance patterns of IAP-Rab6b methylation states. 664 Percentages reflect the data in panel A. (C) Genetic mapping of the modifier locus to a 7.3 Mb 665 interval on distal Chromosome 4 using the GigaMUGA SNP microarray. A map is shown of 666 heterozygous SNPs along the chromosome that are informative between B6 and CAST in 20 N3 667 individuals (full set of individuals shown in SI Appendix, Fig. S4). Heterozygous SNPs shared 668 amongst all highly methylated N3 individuals and absent from all lowly methylated N3 individuals 669 are shown in blue. The corresponding mapped region is highlighted in yellow. (D) Expanded view 670 671 of the 2.5 Mb KZFP cluster located within the mapped interval. Sequence gaps in the current reference genome (GRCm38/mm10) are displayed as black boxes above the annotated genes. 672

573 Stripped region represents the portion of DNA excluded by our independent analysis of N2 574 individuals using the MiniMUGA SNP microarray. KZFP genes are bolded. Annotations were 575 lifted from the UCSC Gencode V24 track.

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Fig. 4. The KZFP cluster Chr4-cl modulates the methylation state of multiple VM-IAPs. (A) Cross-677 locus comparison of N2 methylation states. Methylation levels were quantified at IAP-*Tmprss11d*, 678 IAP-Pink1, IAP-Ect2l, IAP-Rps12, IAP-Trbv31, IAP-Sema6d, IAP-Gm20110, and IAP-Fam78b 679 680 in 20 N2 individuals. IAP-Rab6b methylation level had previously been determined to be high 681 (>60 %, red) or low (< 60%, grey). Red dashed and grey dotted lines connect the average 682 methylation values of N2 individuals across regions. Percent sequence identity to IAP-Rab6b, as 683 determined by the BLAT alignment tool (46), is shown for each locus above the x-axis. (B)Alignment of VM-IAP LTR sequences in the region displaying divergence between Chr4-cl 684 685 targets and non-targets. Contraoriented elements were reverse-complemented (rc) prior to generating the alignment. Dots represent conserved bases, dashes indicate lack of sequence, and 686 divergent bases are shown in blue. The full-length alignment can be found in SI Appendix, Fig. S5. 687 (C) Methylation quantification of genomic DNA extracted from Chr4-cl WT mice (grey circles) 688 and Chr4-cl KO mice (hollow grey circles) on a pure B6 genetic background. (D) Diagram of 689 Chr4-cl KO location (Chr4:145383918-147853419, GRCm38/mm10) and breeding scheme used 690 for the data presented in panels D and E. The Chr4-cl KO was generated in B6 mice, which were 691 subsequently backcrossed to the 129X1/SvJ strain. (E) Methylation quantification of genomic 692 DNA extracted from Chr4-cl WT mice (purple circles) and Chr4-cl KO mice (hollow purple 693 694 circles) on a mixed B6/129 F2 genetic background. IAP-Sema6d was excluded from this analysis because it is absent from the 129 genome (Fig. 1A). 695

Statistics for panels D and E: unpaired t tests with False Discovery Rate of 5% computed using the two-stage step-up method of Benjamini, Krieger and Yekutieli (* q < 0.05; ** q value < 0.01; *** q < 0.001; **** q < 0.0001).

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Fig. 5. Chr4-cl influences the chromatin and transcriptional landscape at and near targeted 700 VM-IAPs. (A) H3K9me3 and H3K4me3 ChIP-seq signal at Chr4-cl target VM-IAPs in Chr4-cl 701 WT (black) and KO (blue) ES cells of mixed B6/129 genetic background. BAM coverage tracks 702 703 were generated and visualised in IGV. VM-IAPs are shown in red and directionality is indicated 704 with a white arrow. NCBI37/mm9 genome coordinates and neighbouring annotated genes are displayed above and below the ChIP-seq tracks, respectively. (B) As in panel A, but for non-target 705 706 VM-IAPs. (C) Mean H3K4me3 ChIP-seq signal over the seven confirmed Chr4-cl targets (upper 707 panel) and over all solo LTRs of the IAPLTR2_Mm subclass in the mouse genome (lower panel). 708 Dotted lines represent mean signal and shaded regions represent error estimates (standard error 709 and 95% confidence interval). Plots were generated using SeqPlots software (47). (D) RNA-seq signal from Chr4-cl WT (black) and KO (blue) ES cells of mixed B6/129 genetic background for 710 711 the genes Pinkl (upstream of IAP-Pinkl), Slco2Al (upstream of IAP-Rab6b), and Rab6b 712 (downstream of IAP-Rab6b). Two biological replicates per genotype are shown. Datasets were downloaded from the GEO database (accession numbers listed in Dataset S1: Table S6). 713

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Fig. 6. Methylation variability at IAPLTR2_Mm elements is sequence driven. (*A*) Neighbourjoining tree of all solo LTR IAPs of the IAPLTR2_Mm subclass in the B6 genome between 200 and 800 bp in length. Solo LTR sequences were aligned using MUSCLE software and the neighbour-joining tree was built using Geneious Prime software. Navy blue and orange nodes represent experimentally validated VM-IAPs and non-variable IAPs, respectively. The VM-IAPenriched subtree, containing all known IAPLTR2_Mm VM-IAPs (navy), is shown in greater
resolution and labelled with GRC38/mm10 genomic coordinates and strandedness. (*B*)
Methylation quantification of genomic DNA from eight B6 individuals at five solo LTRs in the
VM-IAP-enriched subtree (orange). Percent sequence identity to IAP-*Rab6b* is shown above the
x-axis for each IAP.

725

726 **Fig. 7.** The VM-IAP-enriched subtree exhibits H3K4 trimethylation and distinct KZFP binding. 727 (A) Heatmaps of H3K4me3 ChIP-seq coverage in Chr4-cl WT (left) and KO (right) ES cells of mixed B6/129 genetic background over all solo LTRs of the IAPLTR2_Mm subclass (n = 556). 728 729 VM-IAPs and IAPs belonging to the VM-IAP-enriched subtree were clustered for the analysis. 730 All solo LTRs are anchored from their 5' start to their 3' end, with a pseudo-length of 500 bp. The 731 analysis was extended to 500 bp up- and downstream of each element. Average read coverage is 732 plotted above each heatmap. Dotted lines represent mean signal and shaded regions represent error estimates (standard error and 95% confidence interval). Plots and heatmaps were created using 733 734 SeqPlots (47). (B) Heatmaps of overexpressed ZFP989-HA (left), Gm21082-FLAG (middle), and 735 ZFP429-HA (right) ChIP-seq coverage in F9 EC cells over all solo LTRs of the IAPLTR2_Mm subclass (n = 556). Plotting settings as in panel A. 736





A





Avg. % Methylation

20

N:

B6

300

(6 VM-IAPs)









115





(2 VM-IAPs)





2.5 Mb





В

D





²nd HET x HET generations













IAP-Fam78b











