Human epidemiologic and animal model data indicate that susceptibility to adult-onset chronic disease is influenced by persistent adaptations to prenatal and early postnatal nutrition (1, 2, 13, 14); however, the specific biological mechanisms underlying such adaptations remain largely unknown. Cytosine methylation within CpG dinucleotides of DNA acts in concert with other chromatin modifications to heritably maintain specific genomic regions in a transcriptionally silent state (4). Genomic patterns of CpG methylation are reprogrammed in the early embryo and maintained thereafter (20). Because diet-derived methyl donors and cofactors are necessary for the synthesis of S-adenosylmethionine, required for CpG methylation (23), early nutrition may therefore influence adult phenotype via DNA methylation (26).

Accordingly, it is important to identify genomic regions that are likely targets for early nutritional influences on CpG methylation. Most regions of the adult mammalian genome exhibit little interindividual variability in tissue-specific CpG methylation levels. Conversely, CpG methylation is determined probabilistically at specific transposable element insertion sites in the mouse genome, causing cellular epigenetic mosaicism and individual phenotypic variability (19). Transposable elements (including retrotransposons and DNA transposons) are parasitic elements which are scattered throughout and constitute over 35% of the human genome (32). Most transposable elements in the mammalian genome are normally silenced by CpG methylation (32). The epigenetic state of a subset of transposable elements, however, is metastable and can affect regions encompassing neighboring genes. (19). We hypothesized that the epigenetic metastability of such regions renders them susceptible to nutritional influences during early development.

We tested this hypothesis in viable yellow agouti (Avy) mice. The murine agouti gene encodes a paracrine signaling molecule that signals follicular melanocytes to switch from producing black eumelanin to yellow phaeomelanin. Transcription is initiated from a hair cycle-specific promoter in exon 2 of the agouti (A) allele (Fig. 1A). Transient agouti expression in hair follicles during a specific stage of hair growth results in a subapical yellow band on each hair, causing the brown (agouti) coat color of wild-type mice (8). The nonagouti (a) allele was caused by a loss-of-function mutation in A (5); a/a homozygotes are therefore black. The Avy allele (Fig. 1A) resulted from the insertion of an intracisternal A particle (IAP) retrotransposon into the 5′ end of the A allele (8). Ectopic agouti transcription is initiated from a cryptic promoter in the proximal end of the Avy IAP. CpG methylation in this region varies dramatically among individual Avy mice and is correlated inversely with ectopic agouti expression. This epigenetic variability causes a wide variation in individual coat color (Fig. 2A), adiposity, glucose tolerance, and tumor susceptibility among isogenic Avy/a littermates (15).

Dietary methyl supplementation of a/a dams shifts the coat color distribution of their Avy/a offspring (30). Because Avy/a coat color correlates with Avy methylation status (15), it has been inferred that supplementation alters phenotype via Avy methylation (7). Nevertheless, no study has yet compared Avy methylation among the offspring of supplemented and unsupplemented dams (25). Therefore, to test our hypothesis that transposable elements are targets for early nutritional effects on epigenetic gene regulation, we had to determine if CpG methylation plays a role in diet-induced phenotypic alterations in Avy/a mice.

Agouti pseudoexon 1A (PS1A) was formed when a 4.1-kb genomic region containing exon 1A underwent duplication and inversion (Fig. 1A) (6). In mice that carry the light-bellied agouti (Avy) allele, exon 1A is oriented properly with respect to the agouti gene and drives agouti expression (and yellow pigmentation) throughout the hair growth cycle in ventral follicles. The orientation of the duplication is reversed in mice carrying the A allele (Fig. 1A). In these animals, exon 1A points away from agouti, causing a loss of ventral follicle-specific agouti expression (6). Early genetic analyses concluded...
that the \( A^{\alpha} \) IAP is located within \( agouti \) exon 1A (8); however, this conclusion has not been reevaluated since the subsequent characterization of the inverted repeat in the region (6).

In this study, after determining the actual location of the \( A^{\alpha} \) IAP, we showed that dietary methyl donor supplementation of \( a/a \) dams alters \( A^{\alpha}/a \) offspring phenotype by increasing CpG methylation at the \( A^{\alpha} \) locus. Furthermore, we demonstrated that the epigenetic metastability which confers this lability is due to the \( A^{\alpha} \) IAP.

**MATERIALS AND METHODS**

**Animals and diets.** \( A^{\alpha} \) mice were obtained from the colony at the Oak Ridge National Laboratory (29). The \( A^{\alpha} \) mutation arose spontaneously in the C3H/HeJ strain. Mice carrying the mutation were backcrossed with C57BL/6J mice for one to three generations before being propagated by sibling mating. These animals therefore include 6.25% to 25% of the C3H/HeJ genome and 75% to 93.75% of the C57BL/6J genome (31). This congenic colony has been propagated by sibling mating and forced heterozygosity for the \( A^{\alpha} \) allele for over 200 generations, resulting in an essentially invariant genetic background.

Virgin \( a/a \) females, 8 weeks of age, were assigned randomly to NIH-31 diet or NIH-31 supplemented with the methyl donors and cofactors folic acid, vitamin B12, choline chloride, and anhydrous betaine (Harlan Teklad) (30). All ingredients were provided by Harlan Teklad except for the anhydrous betaine (Finnsgur Bioproducts). The diets were provided for 2 weeks before the females were mated with \( A^{\alpha}/a \) males and throughout pregnancy and lactation. Upon weaning to a stock maintenance diet at age 21 days, the \( A^{\alpha}/a \) offspring were weighed, tail tipped, photographed, and rated for coat color phenotype (Fig. 2A). Animals in this study were maintained in accordance with all relevant federal guidelines, and the study protocol was approved by the Duke University Animal Care and Use Committee.

**FIG. 1.** IAP insertion site in \( A^{\alpha} \) allele. (A) Exon 1A of the murine \( agouti \) gene lies within an interrupted 4.1-kb inverted duplication (shaded block arrows). The duplication gave rise to pseudoexon 1A (PS1A). On the \( A \) allele, PS1A is located \( \sim \)100 kb upstream of exon 2 and \( \sim \)15 kb downstream of the contraoriented exon 1A (6). The \( A^{\alpha} \) mutation was caused by a contraoriented IAP insertion (striped bar; tall arrowhead shows direction of IAP transcription). A cryptic promoter within the long terminal repeat proximal to the \( agouti \) gene (short arrowhead labeled \( A^{\alpha} \)) drives ectopic \( agouti \) expression in \( A^{\alpha} \) animals. In \( A \) and \( a \) animals, transcription starts from a hair cycle-specific promoter in exon 2 (short arrowhead labeled \( A, a \)). Small arrows show the positions of PCR primers used to selectively amplify the exon 1A and PS1A regions. (B) Agarose gel showing products of long-range PCR of \( A^{\alpha}/a \) genomic DNA. Forward (F) and reverse (R) primers are described relative to the direction of the inverted duplicate regions and are shown in A. The same forward primer was used in all reactions. Three different reverse primers specific to the PS1A region (lanes 5 to 7) amplified fragments of \( \sim \)1.4 kb (the \( a \) allele) and \( \sim \)6 kb (the \( A^{\alpha} \) allele, including the IAP insert). Three different reverse primers specific to the exon 1A region (lanes 2 to 4) amplified only the smaller fragments (\( \sim \)1.6 kb). The exon 1A fragments are larger than the PS1A \( a \) fragments due to the more distal location of the exon 1A reverse primers.
Phenotypic classification. The coat color phenotype of $A^v/a$ mice was assessed at 21 and 100 days of age. A single observer classified coat color by visual estimation of the proportion of brown fur: yellow ($\leq5\%$ brown), slightly mottled ($\geq5\%$ but less than half), mottled (about half), heavily mottled (greater than half but $\leq95\%$), and pseudoagouti ($>95\%$). The term pseudoagouti is used to describe $A^v/a$ animals in which ectopic agouti expression was silenced (or nearly silenced) by CpG methylation, recapitulating the brown agouti phenotype of an $A^v$ mouse.

Long-range PCR. The Expand Long-Template PCR system (Roche) was used per the manufacturer's instructions. Primers were designed to amplify either the PS1A or exon 1A region (22): exon 1A forward, 1A-F (TCAGATTCTGGAGTGACAGATAGATCC) and reverse, 1A-R1 (TTCCAG FIG. 2. Maternal dietary methyl supplementation and coat color phenotype of $A^v/a$ offspring. (A) Isogenic $A^v/a$ animals representing the five coat color classes used to classify phenotype. The $A^v$ alleles of yellow mice are hypomethylated, allowing maximal ectopic agouti expression. $A^v$ hypermethylation silences ectopic agouti expression in pseudoagouti animals (15), recapitulating the agouti phenotype. (B) Coat color distribution of all $A^v/a$ offspring born to nine unsupplemented dams (30 offspring; shaded bars) and 10 supplemented dams (39 offspring; black bars). The coat color distribution of supplemented offspring is shifted toward the pseudoagouti phenotype compared to that of unsupplemented offspring ($P = 0.008$).
RESULTS AND DISCUSSION

To determine if the IAP insertion causes the epigenetic metastability in the A<sup>av</sup> region, we needed first to determine the IAP location in the A<sup>av</sup> allele. Exploiting sequence dissimilarities between exon 1A and PS1A (6) (Fig. 1A), long-range PCR was used to amplify the sequence bracketing the consensus IAP insertion site of both regions. This demonstrated clearly that the 4.5-kb IAP insert is contained within PS1A and not in exon 1A, as previously reported (8) (Fig. 1B).

To distinguish between the A<sup>av</sup> and a alleles and thus enable A<sup>av</sup>-specific quantitation of PS1A methylation in A<sup>av</sup>/a mice, we sequenced the PS1A region downstream from the consensus IAP insertion site in a/a and A<sup>av</sup>/A<sup>av</sup> homozygotes. We identified and exploited a single-nucleotide polymorphism within an A/I consensus sequence to cleave the PS1A region of the a allele while leaving the A<sup>av</sup> allele intact. Hence, by digesting genomic DNA with A/I and employing reverse primers specific to PS1A, bisulfite sequencing (9) was used to quantify site-specific CpG methylation of the A<sup>av</sup> PS1A in A<sup>av</sup>/a mice. Importantly, because each A<sup>av</sup>/a cell contains only one copy of the A<sup>av</sup> allele, our assay quantitates the percentage of cells in which each A<sup>av</sup> CpG site examined is methylated.

Dietary supplementation of a/a dams throughout the reproductive cycle did not affect litter size or offspring body weight at age 21 days (data not shown). Supplementation did shift the coat color distribution of A<sup>av</sup>/a offspring toward the brown (pseudogoat) phenotype (Fig. 2B). To determine if this phenotypic change was caused by increased A<sup>av</sup> CpG methylation, we quantitated PS1A CpG methylation at seven CpG sites (~600 bp downstream from the IAP insertion site) in tail tip DNA from all A<sup>av</sup>/a offspring born to nine unsupplemented and 10 supplemented dams. We chose to measure CpG methylation in this region for two reasons: (ii) amplification of bisulfite-treated DNA was more reliable in this region than in the IAP long terminal repeat, and (ii) the correlations between average percent methylation and coat color phenotype are comparable in the two regions (IAP long terminal repeat r<sup>2</sup> = 0.82, downstream of PS1A r<sup>2</sup> = 0.85), indicating that measurements of CpG methylation 600 bp downstream of the IAP insertion site are representative of average methylation levels throughout the PS1A region encompassing the A<sup>av</sup> transcription start site.

Percent methylation in A<sup>av</sup> PS1A was distributed bimodally in unsupplemented A<sup>av</sup>/a offspring (Fig. 3A), suggesting a probabilistic epigenetic switch that tends to assume one of two methylation states (19). Maternal supplementation caused a general increase in methylation at each site (Fig. 3A). We examined the relationships among supplementation, CpG methylation, and coat color by mediational regression analysis (3). The highly significant effect of supplementation on coat coloration vanished when A<sup>av</sup> methylation was included in the model (Fig. 3B). This provides the first experimental evidence that A<sup>av</sup> CpG methylation mediates the effect of supplementation on A<sup>av</sup>/a coat color.

To determine if the nutritional effect on A<sup>av</sup> PS1A methylation in tail DNA extends to other tissues, average percent methylation of A<sup>av</sup> PS1A was also measured in liver, kidney, and brain samples from animals representing the five coat color phenotypes. PS1A methylation in the tail correlated

GATTTCATCAATAATGGCT), 1A-R2 (AGGTAGATTTAAACACCCGGT), and 1A-R3 (GCCGAGAGACCTTAAATATTGCCTG); PS1A forward, 1A-F, and reverse, PS1A-R1 (AACCATTTTTGGAGAAACCCATGAATC), PS1A-R2 (GAAGAAAACCAGATACGACAAGGGTTAG), and PS1A-R3 (TGAATCAGAAAGGATTTAGTAAAATGGCTC).

**Genomic sequencing.** The PS1A and exon 1A regions were amplified from genomic DNA (22) from two A<sup>av</sup>/A<sup>av</sup> and two a/a animals. The primers used were 1A-F and 1A-R1 in exon 1A and 1A-F and PS1A-R1 in PS1A (see above). The PCR products were gel purified and sequenced independently (Perkin Elmer dye terminator cycle sequencing system). Regions of identical sequence among iso- genetic homoyogotes were assembled into contigs with Genecode II software (Biosoft).

**Methylation assay.** A G/A single-nucleotide polymorphism (position 1092 in accession number AF540972, present in both exon 1A and PS1A) that allowed the a and A<sup>av</sup> alleles to be cleaved selectively by A/I and SacI, respectively, was identified. We digested 0.5 μg of genomic DNA (22) overnight with 5 U of either A/I or SacI. Bisulfite modification was performed by a procedure adapted from the recently optimized protocol of Grunau et al. (9). Specifically, each digested genomic DNA sample was precipitated with ethanol, washed, and denatured in 50 μl of 0.3 M NaOH (20 min at 37°C). Deamination was initiated by addition of 450 μl of a solution of saturated sodium bisulfite (Sigma) and 10 mM hydroquanine (Sigma), pH 5.0. Each sample was overlaid with mineral oil and incubated for 4 h at 55°C in the dark. The samples were desalted with the Wizard DNA clean-up system (Promega) and resolved in 50 μl 1 mM Tris-Cl, pH 8.0. DNA samples were desalted by addition of 5.5 μl of 3 M NaOH and incubation at 37°C for 20 min, then ethanol precipitated, washed in 75% ethanol, and suspended in 10 μl of 1 mM Tris-Cl, pH 8.0. Because of the possible differential stability of methylated and unmethylated DNA under long-term storage conditions, all studies were conducted with freshly isolated genomic DNA.

Bisulfite-modified DNA (4 μl) was PCR amplified in 50-μl reactions with Platinum Taq DNA polymerase (Invitrogen) per the manufacturer’s instructions (40 cycles). PCR bands were agarose gel purified (GenElute Minus EBr Spin Columns; Sigma) and sequenced manually (Thermo Sequenose radiolabeled terminator cycle sequencing kit; USB Corporation) according to the manufacturer’s instructions. Sequencing products were resolved by polyacrylamide gel electrophoresis, with blank lanes between the C and T lanes to avoid signal overlap. Percent methylation at each CpG site was quantitated by phosphor imaging (percent methylation = 100 × ([C volume]/[C volume + T volume])). Sequencing confirmed that endonuclease digestion was complete. The seven Cs performed studies. For the exon 1A region, we examined forward primer 1ABF3 (ATTTTGTTGGATATAAGTTAGTATAG), reverse primer 1ABR3 (TAAACTAAACTAATACAAACCC), and sequencing primer 1ABF4 (ATTCTGGAATCTAAAACTATGTTTG), and the PS1A region, we used forward primer 1ABF3, reverse primer PS1A-BR2 (CCATAAAATCTA AAAATAATTTAAATTAAC), and sequencing primer 1ABF4. For the A<sup>av</sup> IAP, we used forward primer 1APF3 (ATTTTTTAGAAAGGAGTAGAAG AGTAG), reverse primer IAPR4 (TAATTTCTCAATAAACATTTTTCAACATTA), and sequencing primer IAPFS (ATATTTTTGAAGTTGaTTATG TATGG).

**Statistical analysis.** Because the supplements were provided to the dams, they were the appropriate units of analysis. Therefore, all analyses used within-litter averages for 9 litters (30 a/a (offspring) or 10 litters (39.4/a) (offspring) for the unsupplemented and supplemented groups, respectively. Group comparisons of litter size and day 21 weights were performed by t test. Percent methylation data were not distributed normally and were therefore transformed dichotomously (<20% = 0, ≥20% = 1) before analysis. Relationships among supplementation, A<sup>av</sup> methylation, and coat color were analyzed by mediational regression analysis (3) with SAS software.

**Nucleotide sequence accession numbers.** The following sequence data were submitted to GenBank: 4<sup>th</sup> PS1A (accession number AF540972), 6<sup>th</sup> PS1A (accession number AF540973), 4<sup>th</sup> exon 1A (accession number AF540974), and 1A exon 1A (accession number AF540975).
highly with that in the other tissues (Fig. 4A). The tissues studied were derived from the three germ layers of the early embryo: endoderm (liver), mesoderm (kidney), and ectoderm (brain). These data thus indicate that Avy methylation is determined in the early embryo and maintained with high fidelity throughout development. This is consistent with previous studies showing high agouti expression in all tissues of yellow but not pseudoagouti Avy/a mice (31). Hence, the nutritional effect

**FIG. 3.** CpG methylation within the Avy PS1A of Avy/a offspring from unsupplemented and methyl-supplemented dams. (A) Percentage of cells methylated at each of seven CpG sites in the Avy PS1A in all Avy/a offspring of nine unsupplemented and 10 supplemented dams. DNA was isolated from tail tips at weaning. The seven CpG sites studied are located ~600 bp downstream from the Avy IAP insertion site. Percent methylation is distributed bimodally in unsupplemented offspring, with less than 20% of the cells being methylated at each site in most animals. Maternal methyl supplementation increases mean methylation at each site, generating a more uniform distribution. Dotted lines show the average percent methylation across the seven sites in all Avy/a offspring according to coat color phenotype. (B) Mediational regression analysis (3) of supplementation, Avy methylation, and coat color. Supplementation significantly affects offspring coat color (top), but this relationship is nullified when Avy PS1A methylation is included in the regression model (bottom). This indicates that Avy CpG methylation is solely responsible for the effect of supplementation on coat color.
on $A^\nu$ methylation likely occurs during early embryonic development and affects all tissues.

Coat color phenotype and $A^\nu$ methylation also persisted to adulthood. Independent classification of coat color phenotype at age 100 days agreed with the day 21 classification in 48 of 50 $A^\nu/a$ animals (data not shown). Similarly, mean $A^\nu$ PS1A methylation in day 21 tail DNA predicted that in day 100 liver DNA ($r^2 = 0.95$). Open triangles, unsupplemented offspring; solid triangles, supplemented offspring. Neither group departed significantly from the line of identity (shown). Hence, $A^\nu$ PS1A methylation is maintained with high fidelity into adulthood.

Alleles such as $A^\nu$, whose epigenetic marks are determined probabilistically but subsequently maintained stably, have been termed metastable epialleles (19). Metastable epialleles are commonly associated with transposable elements (19), which can interfere with the expression of neighboring genes (27). Nonetheless, it remains unclear if the epigenetic metastability of the $A^\nu$ allele is due to the IAP insert or if some unique characteristic of the PS1A sequence contributes to its probabilistic variability in CpG methylation. Because the exon 1A region is nearly identical to the PS1A region (Fig. 1A), it provides an ideal cis negative control region by which to examine the IAP’s influence on epigenetic metastability. Similarly, the PS1A region on the $a$ allele of $A^{a}/a$ heterozygotes provides a negative control region in trans.

We therefore measured site-specific methylation of seven homologous CpG sites within exon 1A and PS1A of the $a$ and $A^\nu$ alleles in $A^{a}/a$ animals of divergent phenotypes (Fig. 5). The extreme interindividual variability of $A^\nu$ PS1A ($P < 0.0001$ by analysis of variance; Fig. 5B) was not observed in consensus sites within PS1A and exon 1A on the $a$ allele (Fig. 5C and D). Instead, methylation in those regions was tightly regulated in a site-specific fashion and did not differ significantly among individuals. Notably, the interindividual coefficient of variation in percent methylation at each site was only slightly greater than that obtained by serial digestion and bisulfite sequencing of replicate DNA samples from a single individual (data not shown). Percent methylation at individual CpG sites was quantitatively similar between the two regions on $a$. Methylation of $A^\nu$ exon 1A, which lies approximately 15 kb upstream of the IAP, was also not significantly different among individual mice (Fig. 5A), but was hypermethylated relative to consensus sites on the $a$ allele ($P < 0.0001$). Clearly, the epigenetic metastability that renders $A^\nu$ nutritionally labile is associated with the IAP insertion.

Conversely, proximity to PS1A apparently destabilizes IAP methylation. We performed bisulfite sequencing of the IAP/PS1A junction to quantify methylation at nine CpG sites within the cryptic promoter of the IAP (data not shown). Each animal’s average percent methylation in the PS1A region was predicted by that in the neighboring IAP regardless of maternal diet ($r^2 = 0.92, n = 22$ animals). Hence, whereas most IAPs in the mouse genome are heavily methylated (24), methylation at the $A^\nu$ IAP correlates with that in the neighboring PS1A region and varies dramatically among individuals. Therefore, epigenetic metastability at the $A^\nu$ locus occurs via a mutual interaction between a transposable element and its specific genomic region.

These results indicate that epigenetic metastability caused by juxtaposition of transposable elements and genomic promoter region DNA renders a subset of mammalian genes epigenetically labile to the effects of nutrition and other environmental influences during early development. Our findings have important implications for humans because transposable elements constitute over 35% of the human genome (32) and are found within about 4% of human genes (16). Furthermore, many human genes are transcribed from a cryptic promoter within the L1 retrotransposon (17), analogous to ectopic agouti transcription originating in the $A^\nu$ IAP. It has been proposed that transposable elements in the mammalian genome cause considerable phenotypic variability, making each individual mammal a “compound epigenetic mosaic” (27). Our results provide compelling evidence that the specific composition of each individual’s “epigenetic mosaic” is influenced by early nutrition.

FIG. 4. $A^\nu$ PS1A methylation as a function of tissue type and animal age. (A) Average percent methylation of seven CpG sites in the $A^\nu$ PS1A in tail (T), liver (L), kidney (K), and brain (B) samples from five $A^{a}/a$ animals representing the five coat color classes shown in Fig. 2A. $A^\nu$ methylation in the tail correlates highly with that in other tissues ($r^2 > 0.98$ for all comparisons). (B) Average percent methylation of $A^\nu$ PS1A in day 100 liver versus that in day 21 tail DNA. Percent methylation in day 21 tail predicts that in day 100 liver ($r^2 = 0.95$). Open triangles, unsupplemented offspring; solid triangles, supplemented offspring. Neither group departed significantly from the line of identity (shown). Hence, $A^\nu$ PS1A methylation is maintained with high fidelity into adulthood.
Our findings are also important in the context of epigenetic inheritance at the \textit{Avy} locus. When \textit{Avy}/a animals inherit the \textit{Avy} allele maternally, agouti expression and coat color phenotype are correlated with maternal phenotype in that yellow dams produce fewer pseudoagouti offspring than do pseudoagouti dams (28, 30). This phenotypic inheritance was originally attributed to a maternal effect on metabolic differentiation (28). A recent study (15), however, suggests that this parental effect...
is caused by incomplete erasure of epigenetic marks at the $A^{v}$ locus in the female germ line. Our findings show that early nutrition can influence the establishment of epigenetic marks at the $A^{v}$ locus in the early embryo, thereby affecting all tissues, including, presumably, the germ line. Hence, incomplete erasure of nutritionally induced epigenetic alterations at $A^{v}$ provides a plausible mechanism by which adaptive evolution (10) may occur in mammals.

The moderate nature of the nutritional treatment used in these studies further underscores their relevance to humans. Whereas severe methyl donor deficiency has been demonstrated to induce gene-specific DNA hypomethylation in rodents (12), we show here that merely supplementing a mother's nutritionally adequate diet with extra folic acid, vitamin B$_{12}$, choline, and betaine can permanently affect the offspring's DNA methylation at epigenetically susceptible loci. This finding supports the conjecture that population-based supplementation with folic acid, intended to reduce the incidence of neural tube defects, may have unintended influences on the establishment of epigenetic gene-regulatory mechanisms during human embryonic development (21).

It is increasingly evident that epigenetic gene regulatory mechanisms play important roles in the etiology of human diseases (11, 18). Our findings demonstrate that mammalian metastable epialleles associated with transposable elements enable early environmental influences, including nutrition, to persistently affect these important regulatory mechanisms. Hence, epigenetic alterations at metastable epialleles are a likely mechanistic link between early nutrition and adult chronic disease susceptibility.

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