Alu-mediated 100-kb deletion in the primate genome: The loss of the agouti signaling protein gene in the lesser apes

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Agouti signaling protein (ASIP) is an endogenous antagonist of melanocortin receptors that controls a wide range of physiological functions. Its central role in regulation of the melanocortin system implied that ASIP has been relevant to the evolution of various physiological traits in primates. In this study, we have tried to determine DNA sequences of the ASIP gene (ASIP) of various simian species to find molecular evolutionary aspects of ASIP. Unexpectedly, we found that the whole coding region of ASIP was missing only from the gibbon genome; gibbons constitute a large group of hominoid species in Southeast Asia. Our analyses revealed that unequal homologous recombination mediated by two AluSx elements erased a ~100-kb region including ASIP from the gibbon genome. The data provide new evidence for the significant roles of Alu elements in the dynamic evolution of the primate genome.

[Supplemental material is available online at www.genome.org. The sequence data from this study have been submitted to GenBank under accession nos. AB236869–AB236883.]

Melanocortin receptors (MCRs) belong to the superfamily of G protein-coupled seven transmembrane receptors and participate in the regulation of a variety of physiological functions (Gantz and Fong 2003). Five subtypes of MCRs (MC1R–MC5R) are currently identified (Mountjoy et al. 1992). Proteolytic derivatives of proopiomelanocortin (POMC), including α-melanocyte stimulating hormone (MSH), γ-MSH, and adrenocorticotropic hormone, are endogenous agonists of MCRs; on the other hand, agouti signaling protein (ASIP) and agouti-related protein are endogenous antagonists of MCRs. Transcripts of the ASIP gene (ASIP) have been found in hair follicles of postnatal mice, whisker plate and ventral trunk in embryonic mice (Millar et al. 1995), and various organs in humans including ovary, heart, adipose tissue, liver, kidney, and foreskin (Kwon et al. 1994; Wilson et al. 1995).

ASIP was originally identified in a mouse mutant line, lethal yellow (A’), which shows yellow fur, diabetes, obesity, tumor susceptibility, and lethality of homozygous offspring (Bultman et al. 1992). Because of a 120- to 170-kb deletion, expression of ASIP in A’ mice is under the control of promoters of the hnRNP-associated with lethal yellow gene (RALLY, Fig. 1) that encodes ubiquitously expressed RNA-binding protein (Michaud et al. 1993; Miller et al. 1993; Duhl et al. 1994). The over- and ectopic expression of ASIP primarily deregulated signaling pathways that were mediated by MC1R and MC4R in hair follicles and hypothalamus, respectively, and consequently resulted in excessive production of yellowish pheomelanin and abnormal eating behavior (Yen et al. 1994; Klebigh et al. 1995; Ollmann et al. 1997). On the other hand, null mutants of ASIP showed black fur but did not show the other conspicuous traits (Bultman et al. 1992; Perry et al. 1995; Miltenberger et al. 2002).

Evolutionary homologs of ASIP have been isolated from numerous mammalian species including humans (Kwon et al. 1994; Väge et al. 1997, Leeb et al. 2000; Rieder et al. 2001; Eizirik et al. 2003; Kerns et al. 2004; Girardot et al. 2005), and, moreover, from chicken and fish (Klovins and Schiöth 2005). Genomic organization of ASIP is highly conservative in these mammalian homologs; protein-coding exons 2–4 were localized in a <10-kb region, and non-coding, alternatively transcribed exon 1, which may participate in transcriptional regulation, scattered in the larger upstream region (Bultman et al. 1992; Vrieling et al. 1994; Girardot et al. 2005). Associations between mutations in ASIP and body color variation suggest that ASIP participates in pigmentation via MC1R; details of the physiological roles of ASIP in these species remain unclear (Väge et al. 1997; Kanetsky et al. 2002; Eizirik et al. 2003; Kerns et al. 2004; Bonilla et al. 2005).

The main functions of ASIP, regulation of pigmentation and energy homeostasis, are of interest in developing medical treatments for obesity, diabetes, and pigmentation disorders. Therefore, it may be fruitful to characterize the molecular evolution of ASIP in mammals, especially in the evolutionary lineage leading to modern humans. For this purpose, we tried to isolate ASIP from various primate species. Unexpectedly, we have revealed that the gibbons (Family Hyllobatidae), a group of monophyletic hominoid species from Southeast Asia characterized by relatively small body size, monogamous society, and highly arboreal niches (Fleagle 1999), have lost the whole coding sequences of ASIP and adjacent genomic regions by AluSx-mediated unequal homologous recombination. Our data provide new evidence for the significant role of Alu-mediated deletion in the dynamic evolution of the primate genome.

Results

Loss of ASIP in gibbons

We determined the nucleotide sequences of ASIP from various simian species including great apes, Old World monkeys, and New World monkeys; however, no specific PCR products were obtained for the gibbons. We then performed genomic Southern
DNA are also indicated (see Methods). Restriction enzymes used in the fragmentation of genomic S. oedipus N. leucogenys (Gibbons) H. lar Chemiluminescence images of genomic Southern blotting for ASIP are also shown. DNA samples loaded in each lane are as follows: 165-kb region around human subfamilies, while Young Alu some 20 (∼1) and human chromo-
left Old and New World Monkeys (H. pileatus, and H. agilis, S. syndactylus, and S. oedipus. Restriction enzymes used in the fragmentation of genomic DNA are also indicated (see Methods).

blotting analyses (Fig. 1). Probes designed based on human ASIP exon 2, which encodes an evolutionarily highly conserved leader peptide region, did not detect specific DNA fragments in the gibbon genome, while a probe for RALY, which is 200 kb away from ASIP on human chromosome 20, detected a specific DNA fragment in the gibbon genome. We thus considered that the gibbons might have lost ASIP from their genome.

The genomic organization of human chromosome 22q11, in terms of the relative position of ASIP and its neighbors including RALY, the eukaryotic translation initiation factor2 subunit 2β gene (EIF2S2), the S-adenosylhomocysteine hydrolase gene (AHHCY), and the itchy homolog E3 ubiquitin protein gene (ITCH), was highly conserved in the mouse chromosome 2 counterpart (Fig. 1) (Deloukas et al. 2001; Waterston et al. 2002). A noticeable difference between the two homologous regions was the presence of high-density short interspersed elements, especially Alu elements, in the human counterpart. In the 165-kb region around human ASIP, 81% of Alu elements belonged to the old Alu subfamilies (Fig. 1), which experienced the peak of expansion in the primate genome 35–55 million years ago (Mya) (Shen et al. 1991; Kapitonov and Jurka 1996; Batzer and Deininger 2002). These Alu elements were considered to have existed in the primate genome prior to the diversification of the hominoids.

Alu elements act as a source of genomic instability among primates via several different mechanisms (Batzer and Deininger 2002). We considered that the Alu elements mediated the loss of ASIP in the gibbon genome. To identify chromosomal break points, we estimated the span of the putative deletion in the gibbon genome by using PCR and direct sequencing. A total of 36 amplicons, including five human Sequence Tagged Sites, were examined for Hylobates lar DNA. We found a missing region in the gibbon genome covering from 94 kb upstream to 2 kb downstream of the three coding exons of ASIP in the human genome (Fig. 2A).

We subsequently examined the PCR amplification of AL035458.35 10008U and AL356229.16 3491L (Fig. 2B,C; Supplemental Tables 3, 4). This PCR failed to yield a specific product from human DNA since the distance between these two primers on the human genome was too large for PCR (>100 kb). We obtained a 4- to 5-kb fragment only from the gibbon DNA but not from human DNA (Fig. 2B,C). As a result of further multiple PCR assays described in Figure 2, we consequently obtained 1.0-kb and nested 0.8-kb fragments from the gibbon DNA that most likely contained the junction of the chromosomal break points. The amplification of 1.0-kb and 0.8-kb fragments was not observed in other primate species including humans, great apes, Old World monkeys, and New World monkeys (Fig. 2C).

Structure of the break point

A BLAST search was performed on 960 bp of the 1.0-kb fragment against the human genome sequences. Nucleotide positions (n.p.) 1–617 showed high homologies with n.p. 11215–11567 and n.p. 11587–11832 of the human genomic contig AL035458.35 (E = 1 × 10−170, 1 × 10−76, respectively), while n.p. 512–945 showed high homology with n.p. 491–925 of the human genomic contig AL356299.16 (E = 0.0). The overlapping portion (n.p. 500–630 of the 1-kb fragment) contained a partial region of AluSx elements in AL350458.35 and AL356299.16 (AluSx™ and AluSx™ in Fig. 3, respectively). The 0.8-kb fragment amplified from H. lar DNA also showed identical chimeric sequences (data not shown). A 26-bp Alu core sequence, which was considered a hotspot of recombination (Rudiger et al. 1995), was highly conserved in AluSx™ in the gibbon chimeric sequence, AluSx™ and AluSx™ (Fig. 3). In addition, the chromosomal break point was located near the Alu core sequence (Fig. 3). These results indicated that the chimeric sequence obtained from H. lar DNA was the junction sequence created by unequal homologous recombination between evolutionary homology of AluSx™ and AluSx™ in the ancestral gibbon genome. AluSx™ and AluSx™ are located at 32,218,520–32,218,810 bp and 32,322,430–32,322,700 bp, respectively, in the assembled human chromosome 20 sequence, and the extent of the region that was deleted by unequal homologous recombination was estimated to be ∼100 kb. The 100-kb deletion involved the whole coding sequence of ASIP and pseudogene 1 of exportin (XPOPTP1). The junction sequences were successfully obtained for the six subjected gibbon species. The features of the junction sequence were conserved among them; therefore, it was revealed that the origin of the 100-kb deletion was older than the diversification of three genera, Nomascus, Symphalangus, and Hylobates.
**Discussion**

*Alu* elements have amplified in the primate genome through retropropagation and exist with an extremely high copy number in the human genome (~500,000 per haploid genome). *Alu*-mediated recombination in the human germ line is known to be responsible for the deletions that are found in association with several inherited diseases, such as α-thalassaemia, Duchenne muscular dystrophy, and Tay-Sachs disease (for review, see Deininger and Batzer 1999). *Alu* elements also abundant in the genomes of other primate species may have contributed to the evolution of their genome organization through such recombination; however, *Alu*-mediated recombination was reported to facilitate small deletions involving few exons in the tropoelastin gene (Szabo et al. 1999). The 100-kb deletion presently identified in the gibbon genome provides new evidence for the participation of *Alu*-mediated recombination in the evolution of the primate genomes.

An *Alu*-mediated 161-kb deletion in the human breast cancer 1 gene (Gad et al. 2003) has suggested that *Alu*-mediated recombination has, indeed, a potential to cause deletions >100 kb. The *Alu* elements causing the 100-kb deletion belong to a member of the *AluX* subfamily, which is one of the oldest and most abundant *Alu* subfamilies in the primate genomes (Batzer and Deininger 2002). It has generally been found that homologous recombination occurs more efficiently in the presence of longer stretches of identical sequences (Hasty et al. 1991). *AluX* elements have accumulated a lot of random substitutions, insertions, and deletions during primate evolution (as seen in Fig. 3), and now in the human genome, their role in homologous recombination should be smaller than in other younger *Alu* subfamilies that maintain stringent homology. However, it has been reported that, in the human genome, *AluX* elements still work to create genetic diversity through unequal homologous recombination (Koda et al. 2000; Ringpfeil et al. 2001; Rossetti et al. 2004). In addition to homologous recombination, gene conversion and retrotransposition should be considered as alternative mechanisms of *Alu*-mediated deletion in the primate genomes; however, gene conversion and retrotransposition mediating much smaller deletions (Hayakawa et al. 2001; Callinan et al. 2005) may not likely fit to the *ASIP* deletion in the gibbon genome.

It is obvious that the 100-kb deletion is specific to the present gibbon species, although the dating of this deletion remains unclear. Fossil evidence suggested that gibbon ancestors emerged in the early middle Miocene (Fleagle 1984), and a recent molecular phylogenetic study has proposed a comparable estimate of 15–18 Mya for the divergence of the gibbons from other hominoid lineages (Raum et al. 2005). Hence, the three genera of gibbons are considered to have shared a substantial period of common ancestry, 8–10 Myr prior to their divergence (Cronin et al. 1984; our unpublished data). The 100-kb deletion likely occurred in the period of the common ancestry.

It is a challenge to infer a phenotypic consequence of the 100-kb deletion in the gibbons. As *ASIP* regulates lipid metabolism via MC4R in humans (for review, see Voisey and van Daal 2002), there is room to presume a possible contribution of the 100-kb deletion to the evolution of a unique phenotype of the gibbons: remarkably small body mass that is related to energy homeostasis. The 100-kb deletion is expected for the primates other than *Gorilla* and *Pan*; however, *Gorilla* is the closest living relative to the early hominids. Hence, 200 Mya for the divergence of the gibbons from other hominoid lineages (Fleagle 1984), there is room to presume a possible contribution of the 100-kb deletion to the evolution of a unique phenotype of the gibbons: remarkably small body mass that is related to energy homeostasis.
sion no. AI241007.1), where the functions of these alternative transcripts are currently unknown. Impacts of the 100-kb deletion on the expression of the neighboring genes wait to be investigated. Finally, embryonic lethality of homozygous *Ay* and lethal nonagouti mice was shown to be caused by mutations in *RALY* and *AHCY*, respectively (Michaud et al. 1993; Miller et al. 1994); these facts may have limited the boundaries of the 100-kb deletion in the gibbon genome.

In summary, we describe here that AluSx-mediated homologous recombination deleted the 100-kb region including the whole coding sequence of *ASIP* from the gibbon genome. Our data provide new evidence for the significant role of Alu elements in the dynamic evolution of the primate genome.

### Methods

#### Samples


#### Sequencing of ASIP

DNA samples were prepared from blood specimens or lymphoblastoid cell lines using the NaI method (Wang et al. 1994) or the standard phenol/chloroform method. Three coding exons (exons 2–4) of *ASIP* were sequenced using PCR and the following direct sequencing technique. Primers for the PCR amplification are listed in Supplemental Table 1. Pairwise combinations of two sets of upper and lower primers were examined for each exon. PCR reactions contained 1.5 mM MgCl2, 0.2 mM each dNTP, 0.4 µM forward and reverse primers, and 0.025 unit per µL AmpliTaq Gold (Applied Biosystems). Thermal cycle was 9 min at 95°C, followed by 40 cycles of 30 sec at 96°C, 30 sec at 50°C–65°C (depending on primers sets and templates), 1 min at 72°C, and a final extension for 5 min at 72°C. The PCR products were purified using Microcon PCR (Millipore). Nucleotide sequences of the PCR products were determined using a BigDye Terminator Cycle Sequencing Ready Reaction kit version 1.1 and an ABI PRISM 310 Genetic Analyzer or an ABI PRISM 3100-avant Genetic Analyzer (Applied Biosystems). Sequences of each exon were combined into continuous protein coding sequences of *ASIP* by referring to sequences of intron–exon boundaries of human *ASIP*.

![Figure 3](https://www.genome.org/content/1/3/488/Figure3.jpg)
Southern blotting analysis

Hybridization probes for exon 2 of human ASIP and human RALY were synthesized using a PCR-DIG labeling Kit (Roche Applied Science). The primers for the PCR-DIG labeling are listed in Supplemental Table 1. DNA (5 µg) was digested with EcoRI or BglII (New England Biolabs) for 16 h at 37°C. Digested fragments were separated by TAE-buffered 0.8% agarose gel electrophoresis for 10 h at 20 V, and then transferred to Hybond N° (Amersham Pharmacia Biotech) using the standard capillary transfer method. After 20 h transfer, DNA was fixed on the membrane by baking for 30 min at 120°C. Pre-hybridization and hybridization were done in DIG-Easy Hyb (Roche Applied Science). Detection of signal was performed using anti-DIG alkaline phosphatase-Fab fragments (Roche Applied Science) and CDP-Star (Roche Applied Science). The chemiluminescent images were recorded by using an LAS-1000 (Fujifilm).

Localization of the chromosomal break points

The profiles of repetitive sequences in a region spanning 32,165–32,330 megabases (Mb) of human chromosome 20 were analyzed by using RepeatMasker. The extent of the deletion involving gibbon ASIP was roughly estimated by using PCR and direct sequencing. The PCR primer sets are listed in Supplemental Table 1. DNA (5 µg) was digested with EcoRI or BglII (New England Biolabs) for 16 h at 37°C. Digested fragments were separated by TAE-buffered 0.8% agarose gel electrophoresis for 10 h at 20 V, and then transferred to Hybond N+ (Amersham Pharmacia Biotech) using the standard capillary transfer method. After 20 h transfer, DNA was fixed on the membrane by baking for 30 min at 120°C. Pre-hybridization and hybridization were done in DIG-Easy Hyb (Roche Applied Science). Detection of signal was performed using anti-DIG alkaline phosphatase-Fab fragments (Roche Applied Science) and CDP-Star (Roche Applied Science). The chemiluminescent images were recorded by using an LAS-1000 (Fujifilm).

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