Primary microcephaly (MCPH) is a neurodevelopmental disorder characterized by global reduction in cerebral cortical volume. The microcephalic brain has a volume comparable to that of early hominids, raising the possibility that some MCPH genes may have been evolutionary targets in the expansion of the cerebral cortex in mammals and especially primates. Mutations in \textit{ASPM}, which encodes the human homologue of a fly protein essential for spindle function, are the most common known cause of MCPH. Here we have isolated large genomic clones containing the complete \textit{ASPM} gene, including promoter regions and introns, from chimpanzee, gorilla, orangutan, and rhesus macaque by transformation-associated recombination cloning in yeast. We have sequenced these clones and show that whereas much of the sequence of \textit{ASPM} is substantially conserved among primates, specific segments are subject to high Ka/Ks ratios (nonsynonymous/synonymous DNA changes) consistent with strong positive selection for evolutionary change. The \textit{ASPM} gene sequence shows accelerated evolution in the African hominin clade, and this precedes hominid brain expansion by several million years. Gorilla and human lineages show particularly accelerated evolution in the IQ domain of \textit{ASPM}. Moreover, \textit{ASPM} regions under positive selection in primates are also the most highly diverged regions between primates and nonprimate mammals. We report the first direct application of TAR cloning technology to the study of human evolution. Our data suggest that evolutionary selection of specific segments of the \textit{ASPM} sequence strongly relates to differences in cerebral cortical size.

Introduction

The human brain, particularly the cerebral cortex, has undergone a dramatic increase in its volume during the course of primate evolution, but the underlying molecular mechanisms that caused this expansion are not known. One approach shedding light on the molecular mechanisms of brain evolution is the analysis of the gene mutations that lead to defects in brain development. Among the best examples of such defects is the human primary microcephaly syndrome. Primary microcephaly (MCPH) is an autosomal recessive neurodevelopmental disorder in which the brain fails to achieve normal growth. The affected individuals have severe reduction in brain size; however, the gyral pattern is relatively well preserved, with no major abnormality in cortical architecture (McCreaey et al. 1996; Mochida and Walsh 2001). Moreover, there are no recognizable abnormalities in the organs other than the central nervous system. The most common cause of MCPH appears to be mutations in the \textit{ASPM} gene (Roberts et al. 2002).

The \textit{ASPM} gene encodes a 10,434-bp-long coding sequence (CDS) with 28 exons, and spans 65 kb of genomic DNA at 1q31. \textit{ASPM} contains four distinguishable regions: a putative N-terminal microtubule-binding domain, a calponin-homology domain, an IQ repeat domain containing multiple IQ repeats (calmodulin-binding motifs), and a C-terminal region (Bond et al. 2002). Though the exact function of the human \textit{ASPM} in the brain needs to be clarified, the homologue in the fruit fly, \textit{Drosophila melanogaster}, abnormal spindle (\textit{asp}), is localized in the mitotic centrosome and is known to be essential for both the organization of the microtubules at the spindle poles and the formation of the central mitotic spindle during mitosis and meiosis. Mutations in \textit{asp} cause dividing neuroblasts to arrest in metaphase, resulting in reduced central nervous system development (Ripoll et al. 1985; do Carmo Avides et al. 2001; Riparbelli et al. 2001). In the mouse (\textit{Mus musculus}) brain, the \textit{Aspm} gene is expressed specifically in the sites of active neurogenesis. Expression in the embryonic brain was found to be greatest in the ventricular zone, which
is the site of cerebral cortical neurogenesis (Bond et al. 2002). This expression profile suggests a potential role for ASPM in regulating neurogenesis.

Interspecies comparisons of ASPM orthologs have shown their overall conservation, but also a consistent correlation of greater protein size with larger brain size (Bond et al. 2002). The increase in protein size across species is due mainly to the increased number of IQ repeats, suggesting that specific changes in ASPM may be critical for evolution of the central nervous system.

In an attempt to reconstruct the evolutionary history of the ASPM gene, we isolated large genomic clones containing the entire ASPM gene in several nonhuman primate species. Sequence analysis of these clones revealed a high conservation in both coding and noncoding regions, and showed that evolution of the ASPM gene might have been under positive selection in hominoids. These clones could also provide important reagents for the future study of ASPM gene regulation in its native sequence context.

Results

Comparison of Genomic Organization of the ASPM Genes in Primates

Homologues from chimpanzee (Pan troglodytes), gorilla (Gorilla gorilla), orangutan (Pongo pygmaeus), and rhesus macaque (Macaca mulatta) were isolated by transformation-associated recombination (TAR) cloning in yeast (Saccharomyces cerevisiae), the technique allowing direct isolation of a desirable chromosomal region or gene from a complex genome without constructing its genomic library (Kouprina and Larionov 2003). The method exploits a high level of recombination between homologous DNA sequences during transformation in the yeast. Since up to 15% divergence in DNA sequences does not prevent selective gene isolation by in vivo recombination in yeast (Noskov et al. 2003), for cloning purposes, a TAR vector was designed containing short human ASPM-gene-specific targeting hooks specific to the exon 1 and 3' noncoding regions (see “Materials and Methods”). The TAR cloning scheme for isolating the ASPM gene homologues from nonhuman primates is shown in Figure 1. The yield of ASPM-positive clones from chimpanzee, gorilla, orangutan, and rhesus macaque was the same as that from the human DNA, suggesting that most homologous regions from nonhuman primates can be efficiently cloned by in vivo recombination in yeast using targeting hooks developed from human sequences.

We have compared complete gene sequences from primate species with a 65-kb, full-size human ASPM gene. All the analyzed genes are organized into 28 exons encoding a 3,470-3,479-amino-acid-long protein. ASPM genes start with an approximate 800-bp-long CpG island, that harbors promoter sequences, 5' untranslated regions, and the first exon (Figure 2). ASPM sequences share a high degree of conservation (Figure 2H), and pairwise DNA identity ranges from 94.5% for macaque and gorilla to 99.3% for the human–chimpanzee comparison (Table 1). Multiple alignment of the genes revealed a low proportion of indels. Only ten insertions/delitions equal to or longer than 50 bp have been found, all of them located within introns (Figure 2B). Seven detected insertions were mainly associated with repetitive DNA: two (AT)$_n$ microsatellite expansions, three Alu insertions, including retroposition of AluY$_9$ in the orangutan–gorilla–chimpanzee–human clade, and retroposition of a new macaque-specific AluY subfamily similar to human AluY2. Analysis of eight different macaque individuals showed that this particular insertion is polymorphic in the macaque population (data not shown), and thus the insertion appears to be very recent. One macaque-specific 245-bp-long insertion is linked to expansion of a 49-bp-long, minisatellite-like array. The remaining macaque-specific insertion (50 bp) is nonrepetitive. A closer analysis suggests that the insert is not a processed pseudogene of known genes (data not shown).

Of the two detected deletions, the macaque-specific 72-bp-long deletion appears to be associated with nonrepetitive
Figure 2. Structure and Evolution of the ASPM Gene in Primates

The scale of all plots corresponds to the consensus sequence obtained based on a multiple alignment of five ASPM genes.

(A) Schematic representation of the alignment. Promoter regions, exons, and introns are marked in gray, red, and blue, respectively. White segments correspond to gaps.

(B) Positions of long (50 bp or longer) insertions/deletions. “O” denotes orangutan, “M” macaque, “OGCH” the orangutan–gorilla–chimpanzee–human clade, and “GCH” the gorilla–chimpanzee–human clade.

(C) Positions of polymorphic bases derived from the GenBank single nucleotide polymorphism (SNP) database.

(D) Positions of the CpG island. The approximately 800-bp-long CpG island includes promoter, 5’ UTR, first exon, and a small portion of the first intron.

(E) Location of an approximately 3-kb-long segmental duplication.

(F) Positions of selected motifs associated with genomic rearrangements in the human sequence. Numbers in parentheses reflect number of allowed differences from the consensus motif (zero for short or two ambiguous motifs, two for longer sites).

(G) Distribution of repetitive elements. The individual ASPM genes share the same repeats except of indels marked in (B).

(H) DNA identity and GC content. Both plots were made using a 1-kb-long sliding window with 100-bp overlaps. The GC profile corresponds to the consensus sequence; the individual sequences have nearly identical profiles.

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DNA. The second one, an 818-bp-long deletion in orangutan, was probably caused by homologous Alu–Alu recombination (see below and Figure S1). The remaining indels are related to expansion/contraction of a short minisatellite array. It was caused either by a 53-bp expansion in the gorilla–chimpanzee–human clade or by two independent deletions/contractions in the macaque and orangutan lineages.

An approximately 3-kb-long intronic segment between exons 4 and 5 is present in several copies in the genome (Figure 2E; Figure S2). Closer analysis of the human genome confirmed that copies of this region are homologous to 24 segmental duplications located mainly in telomeric regions of Chromosomes 1–8, 10, 11, 16, 19, 20, and Y. Based on the sequence similarity and the presence of an L1P4 LINE insertion at the 5′ end, the most closely related are three duplications at 7q11–13. The most similar copy is located on Chromosome 7 and shares 93% identity with the ASPM intronic segment. Five duplications are located on Chromosome 1; the closest copy is found 27 Mb away from the ASPM gene.

We looked for several common motifs associated with genomic breakpoints in cancers (Abeyesinghe et al. 2003). Figure 2F shows the positions of such potentially unstable oligonucleotides. Interestingly, the orangutan-specific deletion (Figure 2B) has its 5′ breakpoint located just 1 bp upstream of a sequence 100% identical to the chi-like consensus motif GCWGGWGG (see Figure S1). The chi motif is recognized by the RecBCD-mediated recombination pathway in prokaryotes and seems to be associated with rearrangements in the human genome (Dewyse and Bradley 1991; Chuzhanova et al. 2003). Both deletion breakpoints in the orangutan deletion are located within 5′ parts of two Alu sequences, suggesting that the deletion was created by homologous Alu–Alu recombination. Similar homologous recombinations with breakpoints located near chi-like motifs in 5′ regions of Alu sequences were described previously (Chen et al. 1989; Rudiger et al. 1995).

In summary, despite the presence of a few indels, coding and noncoding regions of ASPM homologues show a marked degree of conservation.

### Evolution of the ASPM Protein

We have analyzed ASPM CDSs from six primate species: human, chimpanzee, gorilla, orangutan, rhesus macaque, and African green monkey (Cercopithecus aethiops). Except for orangutan and rhesus macaque, two or more ASPM CDSs were used for analysis. ASPM proteins showed the same overall length and domain structure (Figure 3A). The IQ repeat domain contains the same number of repeats, suggesting that their expansion occurred in early primate evolution. The CDSs are, as expected, more conserved than the complete gene sequences with promoter and intronic regions (Table 2; Table 3). Only six short indels were identified (Figure 3B).

From the DNA and protein conservation profiles (Figure 3I), ASPM segments evolve differently along the length of the CDS. N- and C-terminal regions and the region corresponding to exons 5–15 are conserved. In contrast, exons 3 and 4 and the complete IQ repeat domain (positions 1,267–3,225)

### Table 1. Pairwise Identity of Aligned Primate ASPM Genes

|        | Macaque | Orangutan | Gorilla | Chimpanzee | Human |
|--------|---------|-----------|---------|------------|-------|
| Macaque| 94.74   | 94.7      | 94.82   | 94.84      | 94.84 |
| Orangutan| 88.84  | 97.57     | 97.7    | 97.7       | 97.7  |
| Gorilla| 90.17   | 94.99     | 98.76   | 98.76      | 98.76 |
| Chimpanzee| 90.23  | 95.08     | 98.12   | 99.3       | 99.3  |
| Human  | 90.27   | 95.14     | 98.13   | 98.91      | 98.91 |

The pairwise identities were calculated for five complete ASPM genes and therefore include all promoter regions, introns, and exons. The values above the diagonal show DNA identities (in percent) calculated after removing indels. Under the diagonal are values for comparisons with gaps.

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### Table 2. Pairwise Identity of ASPM CDSs

|        | Green Monkey | Macaque | Orangutan | Gorilla | Chimpanzee | Human |
|--------|--------------|---------|-----------|---------|------------|-------|
| Green monkey| 99.02      | 97.29   | 97.17     | 97.32   | 97.29      |
| Macaque | 98.89       | 97.37   | 97.31     | 97.45   | 97.43      |
| Orangutan| 97.04       | 97.06   | 98.68     | 98.7    | 98.69      |
| Gorilla | 97.09       | 97.16   | 98.45     | 99.14   | 99.14      |
| Chimpanzee| 97.26      | 97.33   | 98.5      | 99.11   | 99.51      |
| Human  | 97.23       | 97.3    | 98.49     | 99.11   | 99.51      |

The pairwise identities were calculated for six CDSs. The values above the diagonal show DNA identities (in percent) calculated after removing indels. Under the diagonal are values for comparisons with gaps.

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are more variable. Indeed, nonsynonymous substitutions in hominoid primates (Figure 3C) and in ancestral lineages (Figure 3D) and nonsynonymous polymorphism (Figure 3E) are nearly absent in the conserved central (exons 5–15) and C-terminal regions. This pattern indicates different rates of evolution along the ASPM protein, visualized by plots of synonymous Ks and nonsynonymous Ka rates (Figure 3F) and supported by phylogenetic analysis (see below and Figure 4). It is notable that the comparison of the primate and mouse proteins also revealed the same pattern of conservative and nonconservative regions along ASPM protein (Figure S3).

Analysis of the nonsynonymous/synonymous substitution ratio (\(\omega = \text{Ka}/\text{Ks}\)) revealed an elevated value in the human branch (Figure 4A). According to the likelihood ratio test, the human \(\omega\) rate is significantly different from the rate in the rest of the tree (\(p < 0.05\)). Also, the model that the complete gorilla–chimpanzee–human clade is evolving at one \(\omega\) rate different from that in the rest of the tree is well supported (\(p < 0.01\)). Because ASPM consists of regions with different degrees of sequence conservation (see Figure 3), we separately analyzed a conserved region (exons 5–15 plus a small part of exon 16) and a variable IQ repeat domain. As can be seen (Figure 4B) the conserved region has all branches shorter, indicating overall a slower rate of evolution. In the human lineage, the \(\omega\) ratio equals zero; however, the test for whether the human branch has a different (lower) \(\omega\) rate than the rest did not yield significant values. In contrast, the tree based on the variable IQ repeat domain exhibits \(\omega\) values greater than one for the human and gorilla branches (Figure 4C). The likelihood ratio test supports the model in which human and gorilla lineages evolved under a significantly higher \(\omega\) ratio than the rest of the tree. Similar results were obtained for exon 18 with additional sequences from two New World monkeys (Figure 4D). As seen from Figure 4A–4D, different sequences from African green monkey, gorilla, and chimpanzee individuals result in different \(\omega\) values for their corresponding terminal branches. One chimpanzee sequence also produced an \(\omega\) ratio greater than one for exon 18 (Figure 4D). It is worth noting that neither codon bias nor selection on third codon positions seemed to influence the synonymous rate Ks strongly (Table S1). Therefore, the high Ka/Ks ratios in human and gorilla are likely to be products of adaptive evolution.

Sequencing of two CDSs in African green monkey, three in gorilla, and three in chimpanzee allowed us to look for ASPM polymorphism in those species (see Figure 3E). Human polymorphism data from ASPM mutant haplotypes are not representative of wild-type variation so were not used in these comparisons. For African green monkey, five synonymous and five nonsynonymous changes were found between two sequences. The gorilla and chimpanzee CDSs in particular showed an apparently high degree of replacement polymorphism. Gorilla polymorphism included 35 point mutations (15 silent mutations and 21 replacements). Chimpanzee sequences differed in five synonymous and 11 nonsynonymous sites. In order to interpret this seemingly high level of observed polymorphism, intraspecific diversity was compared to interspecific diversity using the McDonald and Kreitman test (McDonald and Kreitman 1991). In the case of chimpanzee polymorphism compared to divergence with human, we could not reject the null hypothesis that polymorphism and divergence between species were significantly different (William’s adjusted G statistic = 0.083, chi-square with 1 d.f., not significant; values based on PAML-generated Ka and Ks values using the free ratio model). Gorilla polymorphism was compared to divergence between the gorilla common ancestor and the human–chimpanzee common ancestor. In this case we can reject the null hypothesis (William’s adjusted G statistic = 122.45, chi-square with 1 d.f., \(p < 0.001\)) to conclude that the pattern of gorilla polymorphism is therefore different from the divergence pattern. Indeed gorilla polymorphism is less than variation resulting from divergence: within species, the \(\omega\) ratio is 1.43 for gorillas compared to 2.2 for the divergence between the gorilla common ancestor and the human–chimpanzee common ancestor. Intraspecific variation, although seemingly unusual in showing so many replacement substitutions in both chimpanzee and gorilla, is less than or in line with what we have observed for ASPM divergence between species. Therefore, relaxation of selection cannot explain the high nonsynonymous/synonymous substitution ratios among African hominoids, further supporting the idea that adaptation has occurred in ASPM.

Discussion

In this study, we applied TAR cloning technology to investigate molecular evolution of the ASPM gene, which is involved in determining the size of the human brain and in which mutations lead to MCPH. The ASPM homologue in the fruit fly is essential for spindle function, suggesting a role for this gene in normal mitotic divisions of embryonic neuroblasts. Complete gene homologues from five primate species were isolated and sequenced. In agreement with the predicted critical role of ASPM in brain development, both coding and noncoding regions of ASPM homologues showed a marked degree of conservation in humans, other hominoids, and Old World monkeys. The differences found in noncoding regions were small insertions/deletions and lineage-specific insertions of evolutionarily young \(Alu\) elements into introns.

Analysis of nonsynonymous/synonymous substitution ratios indicates different rates of evolution along the ASPM protein: part of ASPM evolved under positive selection while other parts were under negative (purifying) selection in human and African ape lineages. Such “mosaic” selection has been previously described for other proteins (Endo et al. 1996; Crandall et al. 1999; Hughes 1999; Kreitman and Comeron 1999). When our work was completed, the paper by Zhang supporting accelerated evolution of the human ASPM was issued (Zhang 2003). However, because the author did not analyze the gorilla gene homologue, he concluded that accelerated sequence evolution is specific to the hominid lineage. Our finding that selection on ASPM begins well before brain expansion suggests that the molecular evolution of ASPM in hominoids may indeed be an example of a molecular “exaptation” (Gould and Vrba 1982), in that the originally selected function of ASPM was for something other than large brain size.

In the case of ASPM, rapidly evolving residues are mainly concentrated in the IQ repeat domain containing multiple IQ motifs, which are calmodulin-binding consensus sequences. While there is no direct evidence yet, it is likely that the function of human ASPM is modulated through calmodulin
The scale of all plots corresponds to the 3,480-amino-acid-long protein alignment; positions in the CDS were scaled accordingly.

(A) Structure of the human ASPM CDS and protein. The first scheme shows positions of major domains in the ASPM protein (Bond et al. 2002). The putative microtubule-binding domain is in gray, the calponin-homology domain in orange, IQ repeats in blue, and the terminal domain in black. Positions of exons in the CDS are drawn in the second block. To separate individual exons, odd numbered exons are colored in black and even numbered ones in white.

(B) Positions of insertions/deletions in the protein sequences. Coordinates correspond to the human protein sequence. “O” denotes orangutan, “G” gorilla, “M” macaque, “Gm” African green monkey, and “OGCH” the orangutan–gorilla–chimpanzee–human clade.

(C) Substitutions in hominoid CDSs relative to the common ancestor. The expected ancestor CDS was derived using ML codon reconstruction implemented in PAML. African green monkey and rhesus macaque were outgroups. Nonsynonymous/synonymous (κ = Ka/Ks) ratios were free to vary in all branches. Positions marked in green correspond to synonymous changes relative to the ancestral sequence; the red bars indicate nonsynonymous changes.

(D) Synonymous (red) and nonsynonymous (green) changes in ancestral lineages leading to human. aOGCH–aGCH is the ancestral lineage from the orangutan divergence to the gorilla divergence; aGCH-aCH represents the lineage from the gorilla divergence to the chimpanzee common ancestor. aCH–human corresponds to the human lineage after the chimpanzee divergence. There are seven synonymous and 19 nonsynonymous human-specific substitutions. Methods and description are the same as in (C).

(E) Positions of polymorphic bases for different CDSs of African green monkey, gorilla, chimpanzee, and human. Positions marked in green correspond to synonymous polymorphisms, and the red bars indicate nonsynonymous sites. Numbers of compared sequences are in parentheses;
Table 3. Pairwise Identity of ASPM Proteins

|             | Green Monkey | Macaque | Orangutan | Gorilla | Chimpanzee | Human |
|-------------|--------------|---------|-----------|---------|------------|-------|
| Green Monkey| 98.36        | 95.81   | 95.56     | 95.76   | 95.59      |       |
| Macaque     | 98.36        | 96.1    | 95.85     | 96.02   | 95.88      |       |
| Orangutan   | 95.57        | 95.8    | 97.72     | 97.86   | 97.75      |       |
| Gorilla     | 95.48        | 95.71   | 97.49     | 98.41   | 98.3       |       |
| Chimpanzee  | 95.71        | 95.91   | 97.67     | 98.38   | 99.02      |       |
| Human       | 95.54        | 95.77   | 97.55     | 98.27   | 99.02      |       |

The pairwise identities were calculated for six protein sequences. The values above the diagonal show DNA identities (in percent) calculated after removing indels. Under the diagonal are values for comparisons with gaps. 
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A surprising number of cerebral cortical neurons are aneuploid (Rehen et al. 2001). Perhaps directed selection of specific domains of ASPM helps ensure faithful chromosome segregation to allow a larger number of cerebral cortical neurons to be formed without an unduly high incidence of chromosome aneuploidy.

Functional genomics studies are clearly needed to elucidate the exact nature of the molecular mechanisms affected by ASPM gene evolution in hominoids. Here, we have demonstrated the utility of TAR cloning for evolutionary sequence comparisons among humans and other primates. In addition, the ASPM TAR clones isolated in these studies could provide valuable reagents for studying ASPM gene regulation in its natural sequence context. Overall, we anticipate this technology will be extremely useful in studying the evolution of other genes that may be responsible for uniquely human traits.

Note
The related paper by Evans et al. (2004) was published in Human Molecular Genetics shortly after this paper was submitted.

Materials and Methods

TAR cloning of the ASPM gene homologues by in vivo recombination in yeast. To isolate the full-size ASPM gene from the human (Homo sapiens), chimpanzee (Pan troglodytes), gorilla (Gorilla gorilla),...
orangutan (Pongo pygmaeus), and rhesus macaque (Macaca mulatta) genomes, a TAR vector containing two unique hooks was constructed. Two targeting sequences were designed, 151 bp 5' and 151 bp 3', from the available human genomic sequence of ASPM (positions 133,758–133,888 and 132,922–132,931) in the BAC RP11-32D17 (GI:16972838). The targeting sequences were PCR amplified from the human genome DNA using two specific primers (Table S2). PCR products were cloned into a polylinker of the basic TAR vector pVC604 as Apal–SalI and Sac–XhoI fragments. Before transformation experiments, the TAR cloning vector was linearized with SacI to release the TAR element. Genomic DNA samples were prepared from chimpanzee, gorilla, orangutan, and rhesus macaque fibroblast cell lines (Coriell Institute for Medical Research, Camden, New Jersey, United States) in agarose plugs. Spheroplast transformation experiments were carried out as previously described in Kourprina and Larionov (1999). To identify clones positive for ASPM, yeast transformants were examined by PCR using diagnostic primers specific for exon 2 and exon 27 of ASPM (Table S2). Integrity of yeast artificial chromosomes (YACs) and the issue of their stability during propagation in yeast were examined. DNA was isolated from ten subclones carrying the ASPM YACs for each primate, and their size was analyzed by NotI digestion followed by CHEF. Each subclone carried a YAC of similar size, indicating that these clones were stable in yeast. Alu profiles of the clones were checked by TaqI digestion of 1 µg of total yeast DNA isolated from transformants. Samples were run by electrophoresis, transferred to a nylon membrane, and hybridized with an Alu probe. YACs were retrofitted into bacterial artificial chromosomes (BACs) by homologous recombination in yeast using a BACNeo8 retrovector, BRV1, and then transformed into a reaDH10B E. coli strain (Kourprina and Larionov 1999). Before sequencing, the integrity of inserts in BACs was confirmed by NotI, HindIII, and SacI digestion, and Southern blotting. Rhesus, chimpanzee, gorilla, orangutan, and rhesus macaque (approximately 3.2 kb) and exon 18 of the red-chested mustached tamarin (Saguinus labiatus) and black-handed spider monkey (Ateles geoffroyi) (approximately 4.7 kb) were PCR amplified using a pair of specific primers (Table S2) from primate genomic DNA (Coriell Institute for Medical Research) and then TA-subcloned for further sequencing.

RT-PCR of ASPM coding regions. RNAs were extracted from primate cell lines (African green monkey (Cercopithecus aethiops) kidney, COS-7 [American Type Culture Collection, Manassas, Virginia, United States], chimpanzee peripheral lymphoblast, EB176 [JC], and gorilla peripheral lymphoblast, EB [JC] [European Collection of Cell Cultures, Wiltshire, United Kingdom]) using TRizol reagent (Invitrogen, Carlsbad, California, United States). Reverse transcription and 3' and 5' RACE reactions were performed using SMART RACE cDNA Amplification Kit (BD Biosciences, San Jose, California, United States).

Sequencing. Chimpanzee, gorilla, orangutan, and rhesus macaque TAR clone containing full-size ASPM genes were directly sequenced from BAC DNAs (Polushin et al. 2001). Forward and reverse sequences of the promoter and exon 18 as well as fragments of coding regions of the ASPM homologues were run on a PE-Applied Biosystem 3100 Automated Capillary DNA Sequencer (Applied Biosystems, Foster City, United States). Primer pairs for cDNA sequencing were designed based on the human ASPM mRNA sequence. Primer sequences are available upon request. All sequences were named and numbered according to the clone/accession identifier.

Sequence analysis. Genomic sequences were aligned using MAVID (http://baboon.math.berkeley.edu/mavid/) (Bray and Pachter 2004); protein-coding and protein-coding DNA sequences were aligned by DIALIGN2.1 (http://www-bibis.ebi.ac.uk/dialign/) (Morgenstern 1999). Alignments were manually edited in the SEAVIEW editor (http://pbil.univ-lyon1.fr/software/seeawiew.html) (Galtier et al. 1996). We have used a number of programs from the EMBOSS package (http://www.hgmp.mrc.ac.uk/Software/EMBOSS/) for sequence analysis. Short nucleotide patterns associated with genomic rearrangements were searched using FUZZNCL (EMBOSS). We searched for the following recombinogenic motifs: chi-like octamer (CGWWGGWGG), immunoglobulin heptamer (GATAGTG), translin (ATGCAGN(0,4)GCCCWSW and GCNCWSCTN(0,2)GCCCWSSW), topoisomerase II (RNYNNCNNGYNGKNNY), topoisomerase IId (GTNWAYATTNATNR), topoisomerase Iii (YYNTASYGKYTYNTNC), and V(D)J recombina (CAGACTG1(12/ 23)ACAAAAAC). For short or highly ambiguous patterns (topo- isomerase IId matches were ignored; for long Alu repeats, translin, V(D)J recombination) up to two mismatches were permitted. Prediction of CpG islands was performed by CPGLPLOT (EMBOSS) with default parameters (length ≥ 200; CpG/GpG ≥ 0.6; GC ≥ 0.5). CENSOR (http://www.girinst.org/Censor_Server-Data-Entry_Forms.html) (Jurka et al. 1996) and REPEATMASKER (http://repeatmasker.genome.washington.edu/cgi-bin/RepeatMasker; developed by A.F.A. Smit and P. Green) were used for identification of repetitive elements. Minisatellites were detected by TANDEM REPEAT FINDER (Benson 1999). ASPM segmental duplications in the orangutan and rhesus macaque (Table S2) were analyzed by cloning genomic DNA samples from BAC DNAs (Polushin et al. 2001). Forward and reverse sequencing were designed based on the human ASPM coding sequence. Synonymous and non synonymous substitutions were detected by SNAP (http://www.hiv.lanl.gov/content/hiv-db/SNAP/WEBSNAP/SNAP.html), Codon maximum likelihood (ML) in CO- DEML in PAML v. 3.13 (http://abacus.gen.ucl.ac.uk/software/ paml cana.html) (Yang 1997). In the construction of phylogenetic trees, reconstruction of ancestral sequences, and detection of positive selection. Branch lengths and ancestral sequences were reconstructed using a free Ω ratio for individual branches. The methodology of likelihood ratio tests is described elsewhere (Yang 1998). For large alignments, the initial input trees for PAML were estimated by ML implemented in PHYLO_WIN (http://pbl.univ-lyon1.fr/software/phylowin.html) (Galtier et al. 1996). Segmental duplications were clustered by a neighbor-joining method implemented in the same program. Distance measurements for the identification of conspecific specific diversification calculated in PAUP (Swoford, D. L. 2003. PAUP v. 4.0b10; Sinauer Associates, Sunderland, Massachusetts, United States; http://paup.csit.fsu.edu/index.html) and corrected for multiple substitutions using the Tamura-Nei algorithm.

Supporting Information

Commentary. Selection operating on codon usage may increase the Ω ratio by lowering the rate of synonymous substitutions (Sharpe and Li 1987, 1989). Therefore, we tested the correlations between the CAI (Sharpe and Li 1987) and the rate of synonymous substitutions (Ks). We found no significant association between the tested variables. Moreover, interspecies comparisons disclosed that CAI is nearly identical for all compared species, and no CAI increase over other species was detected for human or gorilla (data not shown). On the other hand, there was a significant negative correlation between CAI and both protein and DNA identity. A partial correlation analysis revealed that the significant positive linear correlation between Ka and CAI was merely caused by the strong negative correlation of Ka with DNA and protein identity. When we controlled for identity, the correlation between Ka and CAI disappeared (data not shown). These results may indicate that at positively selected sites, protein changes are prefered over optimization of codon usage, and thus mutations causing amino acid replacements do not immediately produce optimal codons. It should be noted that selection on codon usage seems to be generally relaxed in mammals (Durret and Mouchiroud 2000). Mammalian codon usage as well as the rate of non synonymous substitutions can be potentially biased by selection favoring a high GC content (or even saturation by G and C) at the third codon positions (GC3) (Bernardi and Bernardi 1985; Aota and Ikemura 1986). However, ASPM is an AT-rich gene (GC content 36.4%–37%) and, as expected (Bernardi and Bernardi 1985; Aota and Ikemura 1986), the third codon positions are also AT-rich (GC3 content, 30.6%–31.4%) and thus far from saturation. In summary, neither the codon bias nor selection on the third codon seems to strongly influence the synonymous rate Ks. Therefore the high Ka/Ks ratio in human and gorilla is likely to be the product of adaptive evolution.

Figure S1. Recombination Breakpoints in the Orangutan-Specific 818-bp-Long Deletion

Both orangutan breakpoints are located within 5' portions of two Alu elements. The sequence conservation is marked by different shades of gray. Both Alu elements are compared to their corresponding AluSp and AluXc subfamily consensus sequences. Gorilla, chimpanzee, and human sequences located 1 bp downstream of the 5' breakpoint share a perfect match with the chi-like octamer consensus sequence GCWGWWGG (first box, positions matching the chi consensus are shown in black). On the other hand, the 3' breakpoint sequences are diverged from the chi consensus (second box). Both Alu elements in the alignment are shown from the first position and end at the same
position, and thus positions in one element correspond to positions in the other Alu copy. As can be seen, the breakpoint position in the first AluSp repeat exactly corresponds to the breakpoint position within the second AluS element, suggesting homologous recombination between the two repeats.

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Figure S2. Segmental Duplications of the Fourth Internal Intron
From left to right: phylogeny, chromosomal position, band name, identity to ASPM segment (percent same), and a schematic alignment of segmental duplications. The ASPM segment (black) shares similarity with 24 segmental duplications that contain additional sequences and are present on several human chromosomes. The ASPM copy and three duplications on Chromosome 7 share the same L1P4 terminal insertion, which is absent from all other duplications. The tree on the left shows evolutionary relationships among the duplications estimated by the neighbor-joining method.

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Figure S3. Comparison of Mouse and Human ASPM Proteins
The amino acid identity in the conserved regions is 85.44%, 49.39%, and 68.74% for exon 3, exon 4, and the IQ domain, respectively. In Figure S3.

CDS and Protein Correlations
All correlations were obtained for the same 100-amino-acid/300-nucleotide-long, nonoverlapping windows. The first value shows the correlation coefficient; p-value is in parentheses. The section over the diagonal is calculated using the Pearson (linear) correlation coefficient; under the diagonal are correlations obtained using the Spearman’s rank coefficient—nonparametric). Nontrivial or interesting significant correlations are shown in bold and italics. The CAI represents the mean for all species (the CAI values are nearly identical for individual species). The ω ratio, Ka, and Ks (rows/columns 2, 3, and 4) correspond to all branches of the phylogenetic tree. They were obtained using a ML model with one fixed ω ratio for all branches.

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Table S1. Primers Used in This Work
Upper case letters indicate sequences homologous to ASPM and lower case letters indicate cloning sites.

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Table S2. CDS and Protein Correlations

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