The evolution and consequences of snaR family transposition in primates

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Abbreviations: ASR, Alu/snaR related; CAS, catarrhine ancestor of snaR; CGβ, chorionic gonadotropin beta subunit; DHX34, DEAH box polypeptide 34; FLAM, free left Alu monomer; HAR1, human accelerated region 1; indel, insertion/deletion; LH, luteinizing hormone; LINE, long interspersed sequence; ncRNA, noncoding RNA; nt, nucleotide; NF90, nuclear factor 90; PCR, polymerase chain reaction; SINE, short interspersed sequence; snaR, small NF90 associated RNA; SNP, single nucleotide polymorphism

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The small NF90 associated RNA (snaR) family of small noncoding RNAs (ncRNA) appears to have evolved from retrotransposon ancestors at or soon after pivotal stages in primate evolution. snaRs are thought to be derived from a FLAM C-like (free left Alu monomer) element through multiple short insertion/deletion (indel) and nucleotide (nt) substitution events. Tracing snaR’s complex evolutionary history through primate genomes led to the recent discovery of two novel retrotransposons: the Alu/snaR related (ASR) and catarrhine ancestor of snaR (CAS) elements. ASR elements are present in the genomes of Simiiformes, CAS elements are present in Old World Monkeys and apes, and snaRs are restricted to the African Great Apes (Homininae, including human, gorilla, chimpanzee and bonobo). Unlike their ancestors, snaRs have disseminated by multiple rounds of segmental duplication of a larger encompassing element. This process has produced large tandem gene arrays in humans and possibly precipitated the accelerated evolution of snaR. Furthermore, snaR segmental duplication created a new form of chorionic gonadotropin β subunit (CGβ) gene, recently classified as Type II CGβ, which has altered mRNA tissue expression and can generate a novel short peptide.

The Biochemistry and Expression of snaR Non-Coding RNA

snaR non-coding RNAs were discovered in a screen of binding partners for the double-stranded RNA binding protein nuclear factor 90 (NF90).1,2 Two snaR family subsets (snaR-A and -B) were first identified, and subsequent analysis of human and chimpanzee genomes expanded the snaR family to 30 loci in human and 12 in chimpanzee,4 with approximately half in human (14) belonging to the snaR-A subset (Fig. 1; denoted in red). snaRs are typically ~120 nt long, terminate in a 3’ oligo A/oligo U tract, and are predicted to adopt stable secondary structures.2 In vitro transcription analysis of snaR-A constructs established that these genes contain an intragenic promoter and are transcribed by RNA polymerase III.1 Following its transcription, snaR-A appears to undergo processing and translocation to the cytoplasm, possibly through NF90-mediated transport.2,5 In the cytoplasm, the majority of snaR-A RNA co-sediments with ribosomes, raising the possibility that these non-coding RNAs function in translational control.5

snaR-A is induced upon cellular transformation and is expressed at high levels in most immortal cell lines, including HEK 293 cells from which they were first isolated.1,5 In contrast, RNA gel blot and quantitative RT-PCR analyses have found snaRs to have restricted and discrete human tissue expression. snaR-A is highly expressed in testis and notably the pituitary gland, while other snaR species are well-expressed in testis, but are differentially expressed in regions of the brain.5

The Multiple Stages of snaR Evolution Parallels Primate Evolution

A preliminary search of annotated genomes found snaR genes to be present in human and chimpanzee, but not in other mammals such as mouse or rhesus macaque.1 To determine their genetic
extent in primates and trace their molecular origin we exploited a genetic feature, namely that nearly all snaR genes are surrounded by highly conserved sequence. PCR was conducted on Great Ape genomic DNA using primers complementary to sequence encompassing snaR. All African Great Apes were found to contain snaR genes, but a shorter element (~100 nt) was present in the orangutan PCR duplicon. A search for the orangutan PCR duplicon discovered two orthologous segments separated by ~1.9 Kb on chromosome 19 of the rhesus macaque genome. Searches of the intervening macaque sequence revealed a triplicate tandem repeat of 1.9 Kb on chromosome 19 of the rhesus macaque genome. All African Great Apes were found to contain snaR genes, but a shorter element (~100 nt) was present in the orangutan PCR duplicon. A search for the orangutan PCR duplicon discovered two orthologous segments separated by ~1.9 Kb on chromosome 19 of the rhesus macaque genome. Searches of the intervening macaque sequence revealed a triplicate tandem repeat of 1.9 Kb on chromosome 19 of the rhesus macaque genome. 

**The Transposition of snaR Differs from that of its Ancestors**

Certain genetic properties of ASR and CAS strongly suggest they are novel retrotransposons: (1) ASR and CAS are derived from a retrotransposon; (2) the majority of their loci are flanked by short direct repeats, a hallmark of target site duplication and characteristic of retrotransposition; and (3) their loci are scattered randomly throughout the genomes of Simiiformes. All human CAS, and all but one chimpanzee CAS, have an ortholog in at least another species, whereas a substantial proportion of orangutan CAS and the majority of macaque CAS are species-specific. Therefore, CAS retrotransposition appears to have slowed or even stopped in the African Great Apes. By contrast, few snaR appear to be derived from retrotransposition; instead, most are flanked by sequence identical to the parent locus. This suggests that snaR have disseminated through duplication of a larger encompassing segment or ‘duplicon’. Segmental duplication leads to non-random, largely intrachromosomal distributions and is notably more active in the African Great Apes than in orangutan or lesser apes and monkeys. Indeed snaR display a non-random distribution, and in human are chiefly located on chromosome 19 in two large inverted tandem arrays (Fig. 1). To trace the dispersal of snaR from its parent locus, the ~1.9 Kb segment containing snaR-F on chromosome 19 (Fig. 2), we searched for this segment on the well annotated human genome.

![Figure 1](image_url)

**Figure 1.** Human snaR gene clusters. Shown is a region of human chromosome 19q13.32–33 (middle line). Expansions detail the snaR A/C and A/B/D clusters (top line) and the LG/CGβ cluster (bottom line). The locations and direction of transcription of snaR-A, B, C, D, E, F, G1 and G2 are denoted by colored arrow heads. CG31 and CG12 are denoted by hatched boxes, while DHX34 and LG/CGβ cluster genes are denoted by closed boxes and their direction of transcription by arrows. Distances are in megabases (M). Adapted from Parrott et al. with permission.
This gene is located at 52.56 Mb on chromosome 19q13.32–33, and two short duplications in the LH/CGβ gene cluster situated between the tandem arrays (Fig. 3). Each paralogous duplicon contained a snaR gene and a variable amount of flanking sequence. Based on the sequences flanking the duplicons, snaR genes appear to have diversified along two independent duplication pathways: a major pathway that gave rise to most snaRls, and a second pathway which impacted CGβ evolution (Fig. 3).10

The Major Pathway of snaR Duplication Led to its Accelerated Evolution

The major pathway of snaR gene duplication includes genes for snaR-A, -B, -C and -D present in the chromosome 19 clusters (Fig. 1), and snaR-H and -I on chromosomes 2 and 3. These appear to have undergone a series of genomic rearrangements involving insertion of a fragment of the snaR parental locus into a DEAH box polypeptide 34 (DHX34) gene, followed by segmental duplications (Fig. 3). All of these snaR genes retain flanking sequence derived from DHX34. This gene is located at 52.56 Mb on chromosome 19, -0.6 Mb downstream of the snaR cluster near 53 Mb (Fig. 1). We infer that a fragment of the parental snaR locus was inserted into a copy of a region of the DHX34 gene, giving rise to a hypothetical intermediate (Fig. 3, step i). The insertion point is in long interspersed element (LINE) sequence (hatched in Fig. 3), dividing the DHX34 duplicate into ‘5DX’ (2.1 kbp) and ‘3DX’ (1.2 kbp) fragments. This intermediate presumptively served as the source of H- and I-duplicons, which retain 5DX sequence, and of the A/B/C/D-duplicon which retains 3DX sequence (Fig. 3). All these duplicons contain a ‘Core’ sequence consisting of a snaR gene surrounded by -0.5 Kb of parent sequence (Fig. 3). The Core, which has 5ꞌ Alu sequence and a 3ꞌ end 6 bp upstream of that of the I-duplicon (Fig. 3), is reminiscent of ‘Duplication Cores’ identified as the foci of complex interspersed duplication blocks13 and as a source of rapidly evolving genes undergoing positive selection.9,12,13 It is not known whether snaR acts as the ‘driver’ or is merely a ‘passenger’ of this duplication process.

The A/B/C/D-duplicons that encompass the snaR-A, -B, -C and -D genes consist of the Core linked to variably sized fragments of 3DX and additional chromosome 19 sequence, termed R19 (Fig. 3). Both the 3DX and R19 sequences are flanked by Alu. Initial insertion of the Core-3DX composite (Fig. 3, step ii) appears to have been followed by its tandem duplication with R19, and both events were possibly facilitated by Alu-Alu-mediated recombination (Fig. 3, step iii), thought to be a common recombination mechanism in primates.14 Uniform tandem repeats are evident in the A/C cluster (Fig. 1). In the A/B/D cluster the uniformity of the tandem repeats is altered, especially at the 5ꞌ end (55.29–55.30 Mb) as a result of variability in the lengths of their constituent 3DX (~0.7–1.2 Kb) and R19 (~0.7–3.6 Kb) sequences.

Expansion through segmental duplication often confers genetic freedom on the resultant homologous genes, allowing them to undergo accelerated evolution.11 Interestingly, the most abundant human snaR genes, snaR-A and snaR-B/C, form two distinct subsets and appear to have diverged rapidly from each other.5,6 snaR-A is present in human and gorilla, arguing that they originated in the common ancestor of African Great Apes followed

Figure 2. The step-wise evolution of snaR in the parent locus. Left: Schematic of the major deletions (δ1 and δ2 in red) and expansions (ε1 and ε2 in green) relating FLAM-C, ASR, CAS and snaR genes. These molecular events apparently occurred contemporaneously with major events in primate evolution (dashed horizontal gray lines), allowing their timing to be estimated. Pol III A and B boxes are represented by gray and black boxes, respectively. snaR-12 is an intermediate species between CAS and snaR. Right: Schematic illustrating the insertion of a progenitor FLAM C element (green arrow) into chromosome 19 sequence in primates (open box), and its stepwise evolution to ASR, CAS and snaR. The dissemination of ASR and CAS by retrotransposition (dark blue arrows) and of snaR by segmental duplication (orange arrows) from the parent locus is depicted. Adapted from Parrott and Mathews4 with permission.

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by loss in the genus *Pan*; on the other hand, *snaR*-B and -C are unique to human and presumably evolved after the *Homo-Pan* species divergence. *snaR*-A underwent copy number expansion (Fig. 1) giving rise to 14 paralogous alleles in human and to an unknown number in gorilla. The distribution of *snaR*-B and -C, interspersed with *snaR*-A in the two *snaR* clusters in a tandem repeat pattern, strongly suggests that *snaR*-B/C evolved recently from redundant *snaR*-A copies (Fig. 1). The A/B/C/D-duplication contains multiple SINE and LINE elements and the average substitution rate outside of the *snaR* locus is ~3.7 nt per 120 nt. In contrast, there are ~17 nt changes between human *snaR*-A (121 nt) and *snaR*-C (119 nt) species (14.0–14.2% difference), and consequently these two subsets are predicted to fold into distinctly different structures. This rate of substitution approaches that observed between the human and chimpanzee orthologs of Human Accelerated Region 1 (*HAR1*) small non-coding RNA (18 substitutions in 118 nt, 15.3% difference), which is considered to be one of the most rapidly evolved RNAs in humans. Single nucleotide polymorphisms (SNPs) also demonstrate recent genetic mutation; interestingly both *snaR*-G2 (Human Genome Diversity Project, rs3810177) and *snaR*-I (rs13323015) contain SNPs that display transition mutations between human ethnic groups.

**snaR Transposition Impact on Chorionic Gonadotropin Genes**

Chorionic gonadotropin (CG) is a glycoprotein hormone essential to primate reproduction. It is a heterodimer composed of an α subunit common to luteinizing hormone (LH) and other gonadotropins and a unique β subunit. In human there are 6 *CGβ* genes and a single *LHβ* gene arranged in a ~40 Kb gene cluster between the two large *snaR*
arrays on chromosome 19q13.33 (Fig. 1). The \( CGβ \) genes of African Great Apes can be classified as Type I and Type II. Type I \( CGβ \) genes are thought to have arisen through duplication and minor mutation of the \( LHβ \) gene. The Type II \( CGβ \) genes, \( hCGβ1 \) and \( hCGβ2 \), are unique to African Great Apes and evolved through substitution of ancestral Type I sequence to African Great Apes and evolved through duplication and minor mutation of the \( LHβ \) gene. The Type II \( CGβ \) genes, \( hCGβ1 \) and \( hCGβ2 \), are unique to African Great Apes and evolved through duplication and minor mutation of the \( LHβ \) gene. The Type II \( CGβ \) genes, \( hCGβ1 \) and \( hCGβ2 \), are unique to African Great Apes and evolved through duplication and minor mutation of the \( LHβ \) gene.

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