Extensive Gene Conversion Drives the Concerted Evolution of Paralogous Copies of the SRY Gene in European Rabbits

Armando Geraldes,*†1,2,3 Teri Rambo,4 Rod A. Wing,4 Nuno Ferrand,1,2 and Michael W. Nachman3

1CIBIO, Centro de Investigação em Biodiversidade e Recursos Genéticos, Universidade do Porto, Campus Agrário de Vairão, Vairão, Portugal
2Departamento de Biologia, Faculdade de Ciências da Universidade do Porto, Porto, Portugal
3Department of Ecology and Evolutionary Biology, University of Arizona
4Arizona Genomics Institute, Department of Plant Sciences, University of Arizona

†Present address: Department of Botany, University of British Columbia, Vancouver BC, Canada

*Corresponding author: E-mail: geraldes@interchange.ubc.ca.
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Abstract

The human Y chromosome consists of ampliconic genes, which are located in palindromes and undergo frequent gene conversion, and single-copy genes including the primary sex-determining locus, SRY. Here, we demonstrate that SRY is duplicated in a large palindrome in the European rabbit (Oryctolagus cuniculus). Furthermore, we show through comparative sequencing that orthologous palindrome arms have diverged 0.40% between rabbit subspecies over at least 2 My, but paralogous palindrome arms have remained nearly identical. This provides clear evidence of gene conversion on the rabbit Y chromosome. Together with previous observations in humans, these results suggest that gene conversion is a general feature of the evolution of the mammalian Y chromosome.

Key words: Y chromosome, SRY, gene conversion, Oryctolagus cuniculus.

The completion of the sequence of the euchromatic portion of the male-specific region of the human Y chromosome (MSY) yielded many novel insights into the structure and evolutionary dynamics of this chromosome (Rozen et al. 2003; Skaletsky et al. 2003). Skaletsky et al. (2003) suggested that the human MSY is composed of three sequence classes (ampliconic, X-degenerate, and X-transposed) and that these three classes have distinct properties. According to this view, ampliconic genes occur in multiple copies, are expressed mostly in the testes, and are located in palindrome arms that undergo extensive gene conversion. X-degenerate genes are typically single copy and are generally widely expressed. X-transposed genes derive from a recent transposition from the X to the Y in the human lineage. Other authors have emphasized that some Y-linked genes do not fit easily into these categories, suggesting that Y-linked genes form an evolutionary continuum of degradation from a proto-XY ancestor (Graves 2006).

The ampliconic genes on the MSY are noteworthy because they have been shown to undergo frequent gene conversion in humans (Rozen et al. 2003). Rozen et al. (2003) suggested that such gene conversion might prevent the decay of these genes, although they noted that this would require a directional bias favoring the restoration of the ancestral state.

The human SRY (sex-determining region on the Y) is unusual in that it is the only X-degenerate gene with expression mostly restricted to the testis (primordial gonad), a pattern of expression that is otherwise associated with ampliconic genes (Skaletsky et al. 2003). SRY triggers the development of the undifferentiated gonadal primordia in to testis, and its presence or absence determines whether an individual is born as a male or a female (Gubbay et al. 1990; Sinclair et al. 1990; Koopman et al. 1991). Although in humans SRY is a single-copy gene, it is present in multiple copies in several rodents (Bianchi et al. 1993; Nagamine 1994; Lundrigan and Tucker 1997; Bullejos et al. 1999; Turner et al. 2007) and lagomorphs (Geraldes and Ferrand 2006; Putze et al. 2007). In the only case where SRY gene conversion has been explored, the nucleotide differences among copies suggest that extensive gene conversion is not occurring (Turner et al. 2007).

In order to investigate whether SRY in the European rabbit (Oryctolagus cuniculus) is located in a palindrome and thus potentially subject to gene conversion, we sequenced 92,657 bp of a bacterial artificial chromosome (BAC) clone previously shown to contain SRY. This clone was mapped to Yp12 by fluorescent in situ hybridization (Hayes et al. 2002) and then sequenced and assembled following standard protocols (The Rice Chromosome 10 Sequencing Consortium 2003) (GenBank accession numbers for all sequences in this manuscript: HM230423–HM230433). In this clone, we detected 23,086 bp of an inverted repeat (Y Palindrome Arms 1 and 2—YP1A1 and YP1A2, fig. 1a) separated by 18,911 bp of unique sequence (Y Palindrome 1 Spacer—YP1S, fig. 1a). The SRY open reading frame was located in the arms of a palindrome, and transcription was oriented outwards (fig. 1a). This organization is similar to the palindromic organization of ampliconic genes on the human Y chromosome. We found 11 single nucleotide polymorphisms (SNPs) between palindrome arms (99.94% homology)
and 5 indel polymorphisms located in regions of repetitive sequence. We scanned this clone for interspersed repeats with Repeat Masker and found that repeats make up 37.8% of the total sequence. In the human MSY, repeats make up a similar fraction of ampliconic regions (35.5%), a value much lower than the value for the entire euchromatic MSY (47.4%) or the X-degenerate regions (56.8%) (Skaletsky et al. 2003). After masking all interspersed repeats, we used GeneScan to identify other genes in this BAC clone. Two putative genes were identified, but these were discarded because they had significant Blast hits to retroviral elements present throughout the rabbit genome. Our results demonstrate that there are at least two copies of \textit{SRY} in the rabbit genome and that these copies are located in the arms of an inverted repeat.

Next, we sought to determine the age of the \textit{SRY} duplication by asking whether the duplication boundaries are shared among closely related species. We designed primers (all primers used in this manuscript are available upon request) to polymerase chain reaction (PCR) amplify the two boundaries between palindrome arms and the internal single-copy spacer in the rabbit subspecies (\textit{O. c. algirus} and \textit{O. c. cuniculus}), one hare (\textit{Lepus} spp.), and one cottontail (\textit{Sylvilagus} spp.). Amplification of the boundary between \textit{YP1A1} and \textit{YP1S} was successful in all samples, but the boundary between \textit{YP1S} and \textit{YP1A2} was only successfully amplified in \textit{O. c. algirus} and \textit{O. c. cuniculus}. The two rabbit subspecies are distributed parapatrically in Iberia and diverged approximately 2 Ma (Biju duval et al. 1991; Branco et al. 2000; Geraldes et al. 2006; Carneiro et al. 2009), whereas \textit{Oryctolagus} diverged from \textit{Lepus} and \textit{Sylvilagus} approximately 11.8 Ma (Matthee et al. 2004). This indicates that the \textit{SRY} gene is located in the same palindromic structure in both rabbit subspecies and that the duplication occurred at least 2 Ma. It is also possible that the palindrome predates the divergence between \textit{Oryctolagus} and \textit{Lepus}. Although we could only amplify one of the boundaries in \textit{Lepus}, previous work showed that \textit{SRY} is not a single-copy gene in this genus (Putze et al. 2007). The finding that in rabbits \textit{SRY} has the characteristics of an ampliconic gene, although in humans it is a single-copy X-degenerate gene, reinforces the idea that these three classes of genes are somewhat fluid and that ampliconic genes can arise from X-degenerate genes or from genes acquired from other genomic regions (Graves 2006).

**Fig. 1.** \textit{SRY} palindromic structure in rabbits and levels of divergence between orthologous and paralogous regions. (a) Shown are the results of the sequencing and assembly of a MSY BAC clone from a domestic rabbit genomic library. The two arms of the palindrome are shown as white boxes. \textit{YP1A1} extends from the beginning of the BAC clone to 23,086 bp, and \textit{YP1A2} begins at 41,999 bp and extends at least until 65,085 bp. The hatched box represents 27,572 bp of sequence that is single copy in the sequenced clone but that possibly is still part of \textit{YP1A2}. The black box indicates the 18,911 bp \textit{YP1S} between the two palindrome arms. Ten base pairs of sequence at the boundary between \textit{YP1A1} and \textit{YP1S} and between \textit{YP1S} and \textit{YP1A2} are shown, as well as 10 bp of the \textit{YP1S} sequence in the same regions. Note that these \textit{YP1A1} and \textit{YP1A2} sequences are complementary. The \textit{SRY} gene is shown as a gray box, and the direction of transcription of each copy is indicated by an arrow. Above the scale (in kilobases), each line indicates an interspersed repeat identified with Repeat Masker. (b) Divergence between orthologous sequences of \textit{YP1S} (in black) of \textit{Oryctolagus cuniculus cuniculus} and \textit{O. c. algirus} in the right. Divergence between paralogous \textit{YP1A1} and \textit{YP1A2} in \textit{O. c. cuniculus} at the top, and in \textit{O. c. algirus} at the bottom (all in white). Divergence between orthologous palindrome arms at the center.
The discovery that SRY is located in a palindrome provided the opportunity to ask whether there is evidence for gene conversion between palindrome arms in rabbits, as seen in the human MSY (Rozen et al. 2003). If there is ongoing gene conversion between palindrome arms, divergence between paralogous arms of palindromes should be reduced relative to divergence between orthologous arms. Thus, we predict that YP1A1 and YP1A2 within each rabbit subspecies should be more similar to each other than either is to its ortholog in the other subspecies. To test this hypothesis, we PCR amplified and sequenced ~1.2 kb from the palindrome arms of both rabbit subspecies. We were able to amplify fragments corresponding to each arm of the palindrome by placing one PCR primer within the YP1S region and one primer within the palindrome arm for each fragment. We found that paralogous copies (YP1A1 and YP1A2) within each rabbit subspecies were identical, yet divergence between orthologous copies was 0.40% (fig. 1b and table 1), providing clear evidence for gene conversion. Furthermore, previous work provided some indirect evidence for gene conversion at SRY: a population survey of the distribution of a 7 bp insertion in the 3' region of SRY in rabbit subspecies in Iberia (Geraldes and Ferrand 1992–1999) showed that although this insertion was absent in O. c. cuniculus, in O. c. algirus some individuals had the insertion in the two SRY copies, some had the insertion in just one SRY copy, and some, just like O. c. cuniculus, completely lacked the insertion. The authors suggested that either 1) an identical insertion arose independently in each SRY copy or 2) the insertion arose in just one of the SRY copies and was then converted in some but not all individuals onto the other SRY copy.

We also sequenced ~1.8 kb in the YP1S region and found that divergence between rabbit subspecies was 0.55% (fig. 1b and table 1). This value is slightly higher than the divergence between orthologous palindrome arms, though both estimates are based on only a few substitutions and are therefore approximate. This is in contrast to the divergence between Oryctolagus and Lepus, where divergence at YP1S (3.03%) is lower than divergence at YP1A (5.51%). Assuming a divergence time between Oryctolagus and Lepus of 11.8 Ma (Matthee et al. 2004), we estimate that rabbit subspecies diverged 0.87 Ma based on palindrome arms and 2.14 Ma based on the palindrome spacer. Although approximate, the value of 2.14 Ma is similar to previous reports based on mitochondrial DNA (Biju-duval et al. 1991; Branco et al. 2000), X-linked (Geraldes et al. 2006), and autosomal (Carneiro et al. 2009) data.

Here, we provide the first evidence of gene conversion within MSY palindromes outside of primates, suggesting that this might be a general feature of mammalian Y chromosomes. This, coupled with previous evidence for gene conversion on the Z chromosome of birds (Backstrom et al. 2005), suggests that gene conversion might be a common feature of sex chromosomes that lack homologous recombination. Rozen et al. (2003) suggested that gene conversion of Y ampliconic genes might prevent gene decay by preferentially restoring ancestral alleles. Consistent with this hypothesis, they observed lower levels of sequence divergence between humans and chimpanzees for palindromes than for single-copy regions. In contrast, we observed slightly higher levels of divergence between Oryctolagus and Lepus in the palindrome than in the single-copy spacer (table 1), providing no evidence that gene conversion is reducing the rate of evolution in these taxa. Although multiple copies of SRY have been described in other mammalian species, the genomic context was not characterized and therefore it was not possible to determine if they were located in a palindrome or not. It remains to be seen whether SRY is an ampliconic gene in other species of mammals and whether primates are the exception to the rule by having just one copy of SRY.

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References


Table 1. Nucleotide Divergence between Orthologous MSY Sequences in Two Rabbit Subspecies and Hares.

<table>
<thead>
<tr>
<th>Region</th>
<th>Length (bp)</th>
<th>Dxy (%)</th>
<th>Divergence Time (Ma)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Oryctolagus cuniculus algirus to O. c. cuniculus</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>YP1A</td>
<td>1,239</td>
<td>0.40</td>
<td>5.51</td>
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<tr>
<td>YP1S</td>
<td>1,831</td>
<td>0.55</td>
<td>3.03</td>
</tr>
<tr>
<td>O. c. algirus to O. c. cuniculus</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

a Average pairwise distance.
b Divergence time between rabbit subspecies was calculated assuming a molecular clock and divergence time between rabbits and hares of 11.8 Ma (Matthee et al. 2004).


Carneiro M, Ferrand N, Nachman MW. 2009. Recombination and speciation: loci near centromeres are more differentiated than loci near telomeres between subspecies of the European rabbit (Oryctolagus cuniculus). Genetics 181:593–606.


