Tandem repeats modify the structure of human genes hosted in segmental duplications
Anna De Grassi and Francesca D Ciccarelli

Address: Department of Experimental Oncology, European Institute of Oncology, IFOM-IEO Campus, Via Adamello, 20139 Milan, Italy.

Correspondence: Francesca D Ciccarelli. Email: francesca.ciccarelli@ifom-ieo-campus.it

© 2009 De Grassi and Ciccarelli; licensee BioMed Central Ltd.
This is an open access article distributed under the terms of the Creative Commons Attribution License (http://creativecommons.org/licenses/by/2.0), which permits unrestricted use, distribution, and reproduction in any medium, provided the original work is properly cited.

Abstract

Background: Recently duplicated genes are often subject to genomic rearrangements that can lead to the development of novel gene structures. Here we specifically investigated the effect of variations in internal tandem repeats (ITRs) on the gene structure of human paralogs located in segmental duplications.

Results: We found that around 7% of the primate-specific genes located within duplicated regions of the genome contain variable tandem repeats. These genes are members of large groups of recently duplicated paralogs that are often polymorphic in the human population. Half of the identified ITRs occur within coding exons and may be either kept or spliced out from the mature transcript. When ITRs reside within exons, they encode variable amino acid repeats. When located at exon-intron boundaries, ITRs can generate alternative splicing patterns through the formation of novel introns.

Conclusions: Our study shows that variation in the number of ITRs impacts on recently duplicated genes by modifying their coding sequence, splicing pattern, and tissue expression. The resulting effect is the production of a variety of primate-specific proteins, which mostly differ in number and sequence of amino acid repeats.

Background
The completion of the human genome and recent advances in sequencing technologies have revealed the presence of recently duplicated genomic segments with high degrees of sequence identity. Some of these regions have reached fixation during primate evolution and are known as segmental duplications (SDs) [1]. Other segments are still polymorphic and represent copy number variants (CNVs) within the human population [2-4]. Duplicated blocks are usually enriched in genes, thus providing raw material for the evolution of novel gene families [5-7].

Newly duplicated paralogs undergo rearrangements that usually cause their non-functionalization [8]. Sporadically, these modifications lead to advantageous events, such as the development of a novel function (neo-functionalization) or the repartition of the original function between paralogs (sub-functionalization). Under these circumstances, the new genes are rapidly preserved and fixed into the population [9-14]. Rapid divergence of paralogs immediately after gene duplication is a consequence of the relaxed evolutionary pressure that favors the retention and the propagation of the mutated
alleles [8,15]. In this particular context, errors in DNA replication act as a major source of evolutionary innovation.

One of the most frequent replication errors involves internal tandem repeats (ITRs), which are short genomic regions that undergo homologous unequal crossing-over and replication slippage. ITRs are very frequent in eukaryotic genomes [16] and show a positive correlation with genome size in metazoans [17,18]. The biological role of ITRs has been a matter of long-standing debate. In a gene-centric view of genome evolution, these regions have been often tagged as junk DNA, particularly when they localize in intergenic segments and within introns. However, growing evidence has shown that ITRs are important for the evolution of eukaryotic genomes because they act as potential source of genetic variation owing to their 'mutator properties' [19]. Several examples supporting this role have been accumulating over the years, including cell adhesion in yeast [20], morphological modifications in dogs [21], social behaviors in voles [22], and differences in sexual behavior between primates [23]. In coding exons, repetitions mostly involve trinucleotides due to selection against frameshift [24]. In this context, ITRs, and particularly short repeats or microsatellites (< 10 bp), are highly polymorphic within the human population. Polymorphic trinucleotides are often associated with human genetic diseases, one of the best-known examples being the expansion of polyglutamine traits in Huntington disease and various other spinocerebellar ataxias (for recent reviews, see [25,26]). The association between trinucleotide polymorphisms and genetic diseases might lead to the conclusion that repeat variations are always evolutionarily deleterious. However, this is not true: the CAG repeat of the SCA2 gene is under positive selection within the CEU population, although the biological reasons for this selection are still unknown [27].

We have recently shown other possible evolutionary outcomes deriving from ITR variations. Namely, we described the structural modifications occurring in PRDM7, a primate-specific member of the PRDM gene family, where the repeat contributes to the acquisition of a complex pattern of splicing variants and tissue-specific expression [14]. In the present study, we extend the analysis to all variable ITRs in human paralogs lying in SDs with the aim of verifying whether ITR-driven modifications represent a widespread mechanism for the evolution of novel genes.

### Results

**ITRs modify around 7% of human genes hosted in segmental duplications**

SDs are regions of the human genome longer than 1 kb, with at least 90% sequence identity, and that underwent duplications during the last 25 million years of primate evolution [28]. We grouped all human genes hosted in SDs into 2,948 discrete gene loci, each composed of all overlapping mRNAs lying on the same DNA strand (Figure 1a). This definition of gene loci allowed the identification of human genes hosted within SDs by using directly the genome annotation rather than pre-compiled collections of genes. We performed an all-against-all alignment between all exons in these gene loci to extract the 2,008 loci that are associated through nearly identical exons (Figure 1b). We only considered for further analysis the exon alignments with a variable number of ITRs between the aligned exons (Figure 1c). By looking at the exon-intron structure of the corresponding genes, we identified 102 alignments with variable ITRs retained within both exons and 162 alignments with ITRs occurring at exon-intron boundaries (Figure 1c). After manual inspection to eliminate false positives (see Materials and methods), we identified 53 exon and 27 intron modifications (Figure 1d). We carefully analyzed the transcription evidence supporting each of these modifications to exclude that they were artifacts of RNAs with multiple matches on highly identical genomic sequences, such as SDs. Strikingly, almost all loci with variable ITRs (> 96%) are supported by RNAs with unique or best matches, and only less than 4% are associated with ambiguous transcription evidence (Table 1). This result confirms that ITR-driven gene modifications are real events and not artifacts of genome mapping.

Due to the multiple rounds of duplications during primate evolution, the same modification could be found in several gene loci. Overall, the 80 modifications were detectable in 210 gene loci, lying within 496 SDs (Table 1; Additional file 1).

### Table 1

| Modifications | Associated gene loci | Associated SDs | Unique match | Best match | Multiple match |
|---------------|----------------------|----------------|--------------|------------|----------------|
| Exon (53)     | 180                  | 474            | 48           | 5          | 2              |
| Intron (27)   | 106                  | 370            | 21           | 4          | 1              |
| Both (18)     | 76                   | 154            | -            | -          | -              |
| Total (80)    | 210                  | 496            | 68           | 9          | 3              |

In 18 cases, the same variable ITR can result in both exon and intron modifications. RNAs supporting the modification can have a unique or best match in that locus, or multiple matches in the genome (see Materials and methods).
Variable ITRs therefore affect 7% (210 out of 2,498) of the human genes hosted in primate-specific duplications.

Variable ITRs occur in large groups of recently duplicated paralogs
To better characterize the genes that undergo ITR-driven modifications, we compared the paralogs of the 210 loci with variable ITRs with those of the remaining 1,798 nearly identical loci. We counted the number of paralogs of the 210 loci that were associated through any nearly identical exon, and through only ITR-containing exons. Both comparisons showed that the 210 loci with variable ITRs are significantly enriched in larger groups of paralogs (P-value < 10^-3, Wilcoxon test; Figure 2a, b). The same trend is detectable also when exon and intron modifications are analyzed separately (P-value < 10^-3; Figure S1 in Additional file 2). These data suggest that ITR-driven modifications have occurred in genes that underwent several rounds of duplications. This is not surprising as these genes had higher chances to undergo modifications and likely experienced periods of relaxed evolutionary pressure due to functional redundancy [8].

Interestingly, SDs with variable ITRs tend to occur within the same chromosome more frequently than all other SDs (70.4% and 39.3%, respectively, P-value < 10^-3, chi-squared test). Since intrachromosomal SDs are known to be recent duplications [29], this enrichment may suggest that ITR-driven gene modifications occurred recently during primate evolution. To
be able to date the appearance of the loci with variable ITRs
during primate evolution, we relied on the percentage of iden-
tity between pairs of SDs, which returns an indication of when
the duplication occurred in time [30,31]. When compared to
the rest, the 496 SDs hosting the loci with variable ITRs are
enriched in recent SDs (Figure 2c). In particular, 161 of them
(32.5%) share more than 98% sequence identity and hence
underwent duplication during or after the speciation between
human and chimpanzee [30]. This percentage is significantly
higher compared to all SDs (11.4%) and increases to 46.4%
when, for each of the 80 modifications, only the 168 SDs with
the longest version of each ITR (168). The last two are enriched in highly identical SDs. Genes with variable ITRs, and
especially those with the longest ITR version, have formed
through recent duplications.

Genes with variable ITRs lie in polymorphic regions
of the human genome
The results reported above may suggest that the 496 SDs
bearing variable ITRs continue to undergo further rearrange-
ments and fixation in the human population. We therefore
measured the co-occurrence of SDs with variable ITRs and
human CNVs, which are large polymorphic regions (> 1 kb) of
the human genome accounting for a large portion of human
variation [3,32]. We observed the expected general trend
[30,33,34] in which recent SDs tend to undergo variation
within the human population (Figure 2d). However, when
only SDs with variable ITRs are considered, they are significa-
cantly more represented within human CNVs, independent of
the age of the SD. Also in this case, the signal is still detectable
when only SDs with the longest version of the repeat are con-
sidered (Figure 2d). This observation suggests that ITR-driven modifications preferentially occur in evolutionarily dynamic regions of the genome that are still undergoing modification within the human population.

The fact that SDs with the longest version of the ITRs are particularly enriched in recent SDs as well as in human CNVs may indicate that the direction of ITR modifications within the primate lineage is towards expansion more than contraction, possibly through replication slippage or unequal crossover. To further verify this, we counted the number of ITRs in the orthologous exons of two outgroup species, mouse and dog. For 11 out of 80 modifications we could detect no orthologous sequence (Additional file 1), suggesting that the exon itself originated in primates. For the remaining 69 ITR modifications, at least one ortholog was recovered in mouse or dog. In all cases but two (ZNF100 and FOXD4L) the number of ITRs was higher in human than in the other species. This result confirms that variable ITRs in SDs mostly expanded in the primate lineage, resulting in exon and intron elongations.

**ITR-driven modifications are due to expansion of minisatellites**

Variable ITRs responsible for gene modifications are composed, on average, of 30-bp units that are repeated 4 times for a total length of 160 bp (Table S1 in Additional file 2). When compared to all ITRs within exonic and non-exonic regions hosted in SDs as well as in the whole human genome, variable ITRs affecting the gene structure are significantly longer (Figure 3a) as a consequence of longer units (Figure 3b) rather than of higher numbers of repetitions (Figure 3c). Therefore, ITR-driven modifications of genes hosted in SDs are preferentially mediated by minisatellites. This result can be explained by different and concomitant reasons. First, it partly reflects the fact that we focused on almost identical regions, thus favoring the detection of longer repeat units. As a general trend, ITRs lying in SDs have, on average, repeat units significantly longer than ITRs dispersed in the rest of human genome (Figure 3b). Second, long repeats are more variable than short repeats probably because they enlarge the target sequence for slippage or unequal crossover [35]. Finally, the absence of variable ITRs with repeat units shorter than 9 bp (Figure 3b) suggests a preferential retention of repeats that can significantly diversify the sequence of the encoded proteins.

**Fifty percent of variable ITRs modify protein sequences**

In agreement with an active role in modifying protein sequences, we found that 50% of the detected ITRs occur in the coding sequence of genes lying in SDs (Table 2). This is different, for example, from smaller ITRs in housekeeping genes, which preferentially occur within untranslated regions [36]. We manually analyzed all these 40 modifications in order to verify the effect of the repeats on the resulting proteins. In the majority of cases, the reading frame of the original protein is preserved and variable ITRs cause the elongation of low complexity regions in between globular domains as well as of amino acid repeats, such as zinc fingers and protein-specific repeats (Table S2 in Additional file 2). Often these modifications occur in polymorphic human proteins, such as the keratin-associated proteins, the VCX/Y proteins, the nuclear pore interacting proteins, and the prostate-ovary-testis-endometrium proteins. In this latter case, the formation of an amino acid repeat is involved in the modification of the protein’s cellular localization [37].

In seven cases variable ITRs introduce frame shifts with the formation of novel amino acid sequences (Table S2 in Addi-
### Table 2

| Modification | Non-coding RNAs | UTRs | ITR unit (bp) | In-frame | Out of frame |
|--------------|----------------|------|---------------|----------|--------------|
| Exon (53)    | 10             | 13   | < 40 (9-39)   | 14       | 3            |
|              |                |      | > 60 (63-228) | 9        | 4            |
| Intron (27)  | 9              | 8    | < 40 (9-30)   | 3        | 0            |
|              |                |      | > 60 (62-126) | 6        | 1            |

The number of ITR modifications localized in non-coding mRNAs, untranslated regions (UTRs), and coding sequences is reported. For ITRs lying within coding sequences, the length of the ITR unit in base pairs and the effect on the reading frame of the encoded proteins are also shown.

### ITRs are involved in the diversification of Morpheus paralogs

One of the most complex cases of ITR-driven modification that we identified involves the paralogs of morpheus, a primate-specific gene under strong positive selection in hominoids [11]. To investigate the role of variable ITRs in the diversification of potentially important genes in human evolution, we manually analyzed and reconstructed the structure of the ITR-containing exons in this family. Overall, we identified 26 paralogous loci, most of which have uniquely or best mapping transcripts that encode nuclear pore interacting proteins (NPIPs; Table 3). All these genes but one are hosted in a region of human chromosome 16 that underwent several rounds of duplications during primate evolution [39-41]. The paralogous exons can host two different ITRs, namely type 1 and type 2 repeats (Figure 4a). Type 1 repeats are associated with two different units of 57 bp and 69 bp, respectively, which in turn can be translated into two different frames. As a result, type 1 repeats can produce four distinct amino acid sequences. Type 2 ITRs are much simpler repeats with a single 87-bp unit and a unique reading frame. Depending on the percentage of identity between exons, morpheus paralogs can be assembled into two groups (G29 and G40; Additional file 1), which also reflect a different degree of expansion of the ITR units. Members of G29 host a maximum of four repeats of type 1 and two repeats of type 2, while members of G40 show a large repeat expansion, with up to 39 copies of type 1, and 4 repeats of type 2 (Table 3). Not all variable ITRs present at the genomic level are also found in the mature transcripts and often the same locus is associated with transcripts that differ in the number of ITRs (Figure 4b). However, rather than being due to a complex pattern of alternative splicing, this variability seems the result of the high structural polymorphism of these regions in the human population. All morpheus loci but two overlap with human CNVs, and in at least three cases the polymorphic region corresponds to the repeats (Table 3). Interestingly, there are only two paralogs with less than three ITRs of type 1 and in none are the ITRs in coding exons, suggesting that at least three units of type 1 repeat are required for protein function.

The reason for the high variability in the number of ITRs in the morpheus family is currently unknown. Together with positive selection occurring at the upstream exons 2 and 4 of morpheus [11], it could be a sign of the fixation process that the entire family is currently undergoing in hominoids. Interestingly, the protein encoded by morpheus localizes at the nuclear membrane, where it interacts with the nuclear pore complex [11]. According to several secondary structure predictors [42-46], morpheus hosts a transmembrane segment in its amino-terminal part, followed by a helical portion before the repeats (Figure 4c). Variation in amino acid repeats is a known mechanism to vary the surface of interaction with different targets [47]. This could be the case also for the different paralogs of morpheus, which in this way could adapt and fine-tune their binding to different interactors.

### Discussion

Tandem repetitions of short sequences represent an effective case of ‘dynamic mutations’ in which a secondary event can easily occur after the primary duplication [48]. As a consequence of such high dynamism, it is not surprising that ITRs highly differ between lineages [49,50], species [51], and even individuals [52]. In this study we show that long and variable ITRs are involved in the modification of 7% of human genes hosted in primate-specific SDs. These genes are very recent and therefore likely to be still in the process of formation and fixation. Confirming their evolutionary dynamism, genes with variable ITRs are enriched in human variant regions, in spite of the overall paucity of CNVs that overlap with RefSeq genes [53]. At least 50% of variable ITRs contribute to the modification of coding sequences, mostly leading to the elongation of amino acid repeats in the encoded proteins. When
variable ITRs occur at the exon-intron boundary, they may cause the formation of novel introns (Table S3 in Additional file 2). The majority of such intron modifications (66% of the total; Table 1) also show support for the alternative transcript, in which the repeat is retained within the exon. Alternative transcripts occur less frequently in exon modifications (34%, \(P\)-value = 0.01, chi-squared test), suggesting that the formation of novel introns is a more complex event that requires further rearrangements to generate novel splice sites [54]. For some of the reported cases, variable ITRs cause the activation of cryptic splice sites and the formation of novel introns [55]. This model of intron formation has so far been invoked only very seldom [14,56,57], likely because the fast divergence of intronic sequences makes the identification of

| Group ID | Genomic locus of exon 8 | Exon length (bp) | Type1 | Type2 | CNVs | Associated proteins or transcripts | Exon length (aa) | Type1 | Type2 |
|----------|------------------------|------------------|-------|-------|------|-----------------------------------|------------------|-------|-------|
| G29      | chr.16:68567795-68568136 | 342              | 1     | 0     | 1    | NR_003610, PDXC2 * (U)           | -                | 1     | 0     |
|          | chr.18:11192873-11929237 | 504              | 2     | 2     | 0    | -                                 | -                | -     | -     |
|          | chr.16:14952973-14953380 | 408              | 3     | 0     | 4    | BC023572 * (M)                    | -                | 2     | 0     |
|          | chr.16:15365020-15365427 | 408              | 3     | 0     | 2    | -                                 | -                | -     | -     |
|          | chr.16:1609511-1609918  | 408              | 3     | 1     | 0    | -                                 | -                | -     | -     |
|          | chr.16:28261432-28262010 | 579              | 3     | 2     | 1    | -                                 | -                | -     | -     |
|          | chr.15:105678-15106116  | 439              | 4     | 0     | 1    | -                                 | -                | -     | -     |
|          | chr.16:16351450-16351914 | 465              | 4     | 0     | 3    | BAC85871 (U)                      | 155              | 4     | 0     |
|          | chr.16:16394814-16395312 | 465              | 4     | 0     | 3    | NP_848636 (M)                     | 155              | 4     | 0     |
|          | chr.16:18319329-18319793 | 465              | 4     | 0     | 5    | NP_848636 (M)                     | 155              | 4     | 0     |
|          | chr.16:18359482-18359946 | 465              | 4     | 0     | 5    | NP_848636 (M)                     | 155              | 4     | 0     |
|          | chr.16:28735249-28735848 | 600              | 4     | 2     | 2    | -                                 | -                | -     | -     |
|          | chr.16:28690975-28691574 | 600              | 4     | 2     | 4    | -                                 | -                | -     | -     |
|          | chr.16:28576850-28577449 | 600              | 4     | 2     | 5    | -                                 | -                | -     | -     |
|          | chr.16:72982790-72983476 | 687              | 4     | 2     | 5    | AA660029, NPIPL2 (U)             | 229              | 4     | 2     |
|          | chr.16:28970894-28971553 | 660              | 4     | 2     | 1    | -                                 | -                | -     | -     |
| G40      | chr.16:22452448-22455204 | 2757             | 35    | 4     | 12   | NP_001129337 (U)                 | 919              | 35    | 4     |
|          | chr.16:29300271-29303111 | 2841             | 37    | 3     | 2‡   | AAH94882 (B)                      | 544              | 18    | 4     |
|          | chr.16:30141785-30144625 | 2841             | 37    | 4     | 2‡   | BAC87606 (B)                      | 794              | 29    | 4     |
|          | chr.16:21321092-21324115 | 3024             | 39    | 4     | 5‡   | BAG65049 (B)                      | 970              | 39    | 3     |
|          | chr.16:21753543-21756566 | 3024             | 39    | 4     | 10‡  | BAA13210 (B)                      | 840              | 31    | 4     |

The paralogous sequences of morpheus exon 8 were detected and characterized for the presence and number of type 1 and 2 ITRs. Group IDs refer to Additional file 1. Exon length is reported in base pairs and in amino acids (aa) only when supported by coding mRNAs. Unique (U), best (B) or multiple (M) transcript support is reported for each locus. CNVs that overlap the genomic locus of exon 8 are indicated. *ITRs residing in non-coding RNAs. †CNVs overlapping exclusively with ITRs, and corresponding to identifiers ‡30774, §30757 and 30761 of the Database of Genomic Variants.
intron gains very challenging and often questionable [58-61]. Because our analysis focuses on recent events, it enables the capture of signs of intron formation before they disappear as a result of sequence divergence. None of the putative intron gains reported in our study had been previously identified, likely because our approach does not limit the search for repeated regions to intron boundaries [62] but extends it to the entire ITRs contained within exons. This approach also reduces the chance of false positives due to RT-PCR artifacts [63]. When no canonical splice sites can be identified (Table S3 in Additional file 2), the discrepancy between the number of putative ITRs between genomic and transcribed sequences can be better explained by structural polymorphisms more than by intron gains. This is the case of *morpheus* paralogs, where both ITRs associated with these genes undergo copy number variation within the human population (Table 3).

While microsatellites can be involved in CNV formation [31,53], we found that the repetition of minisatellites seems to play a role especially in the diversification of recently acquired paralogs. How do these modifications affect the function of these genes? The general paucity of functional information on primate-specific genes prevents us from fully addressing this issue. Some hints can be derived, however, from the functional enrichment and the tissue expression of genes with variable ITRs. As expected for recent paralogs,

**Figure 4**
Effect of variable ITRs on the coding sequence of nuclear pore interacting proteins. (a) Amino acid sequences encoded by the ITR units of the human *morpheus* paralog BAG65049 were taken as representatives. Type 1 repeats are associated with two different units of 57 bp and 69 bp, respectively, and are translated into two different frames. This results in four distinct amino acid repeats (1a, 1b, 1c and 1d). Type 2 repeats are much simpler and only produce one sequence. Amino acids are highlighted according to the Clustal color scheme [71]. (b) Representation of the protein sequences encoded by the human paralogs of exon 8 of *morpheus*. Only proteins associated with transcripts shown in Table 3 are reported. (c) Cartoon of the possible three-dimensional structural organization of *morpheus*, based on secondary structure predictions (see text). These predictions were confirmed also for the other RefSeq transcripts. The representation is not to scale.
Recent modifications allow the observation of events that are probably involved in the regulation of transcription (G11 and G12; Additional file 1). Furthermore, variable ITRs can influence the tissue expression [14] and localization of the encoded protein [37], thus confirming that their presence and/or between unrelated SDs were discarded. From the resulting set, alignments bearing a different number of ITRs were extracted. ITRs were recovered from the UCSC singleRepeat annotation track generated using Tandem Repeat Finder (TRF) [66]. As reported in the UCSC website, TRF was run under the following parameters: 2,7,7 = weights for match, mismatch and indels used in the Smith-Waterman alignment; 80, 10 = matching and indel probability; 50 = minimum alignment score; 2000 = maximum size of the repeat unit. Applied to the entire sequence of the human genome, TRF is able to detect ITRs with a total length of 20 to 100,000 bp. Alignments were divided into two groups, according to the position of the variable ITRs in respect to the gene structure. ITR variation could occur either within exons, resulting in exon modification, or at exon-intron boundaries, leading to intron modification. For each modification, representative alignments were manually checked to eliminate false positives due to errors of TRF and/or incorrect alignment between repeats. The 524 RNAs associated with ITR modifications were carefully analyzed to verify the robustness of the transcription support for the new gene structure arrangements. Transcripts were classified as: unequivocally associated with the locus, if their only genomic match in terms of sequence identity and coverage corresponded to the locus with variable ITR; best associated with the locus, if the best genomic match corresponded to the locus with variable ITR; or multiply associated if they had multiple best matches on the genome.

Groups of paralogs, age of SDs and overlap with human CNVs
For each of the 2,008 gene loci, the number of paralogs was defined as the number of loci associated by at least one nearly identical exon (> 95% coverage, > 90% identity). The resulting distributions of paralogs between the 210 loci with variable ITRs and the remaining 1,798 loci were compared, using the non-parametric Wilcoxon test. The percentage of identity between pairs of SDs was recovered directly from UCSC. A non-redundant set of 25,914 pair-wise alignments between SDs was derived, 496 of which involve the 210 loci with variable ITRs. The genomic coordinates of 21,178 human CNVs grouped in 6,558 non-overlapping CNV loci were recovered from the Database of Genomic Variants [67,68] (version 7, March 2009).

**Genomes**

**Materials and methods**

**Identification of gene loci and detection of ITR-driven modifications**

The genomic coordinates of 51,809 alignments between 30,833 primate-specific SDs [28] were recovered from the assembly of the human genome (hg18, March 2006) at UCSC [64]. Starting from 11,118 GenBank human RNAs lying within SDs for at least 95% of their sequence (UCSC mRNA track, frozen at May 2007), 2,948 discrete gene loci were derived by merging all RNAs mapping in the same locus and on the same strand. All exons within SDs were aligned using all-against-all BLAT [65]. Only pairwise alignments with at least 90% identity and covering at least 95% of the shortest exon were retained. Alignments between isoforms of the same exon and/or between unrelated SDs were discarded. From the resulting set, alignments bearing a different number of ITRs were extracted. ITRs were recovered from the UCSC singleRepeat annotation track generated using Tandem Repeat Finder (TRF) [66]. As reported in the UCSC website, TRF was run under the following parameters: 2,7,7 = weights for match, mismatch and indels used in the Smith-Waterman alignment; 80, 10 = matching and indel probability; 50 = minimum alignment score; 2000 = maximum size of the repeat unit. Applied to the entire sequence of the human genome, TRF is able to detect ITRs with a total length of 20 to 100,000 bp. Alignments were divided into two groups, according to the position of the variable ITRs in respect to the gene structure. ITR variation could occur either within exons, resulting in exon modification, or at exon-intron boundaries, leading to intron modification. For each modification, representative alignments were manually checked to eliminate false positives due to errors of TRF and/or incorrect alignment between repeats. The 524 RNAs associated with ITR modifications were carefully analyzed to verify the robustness of the transcription support for the new gene structure arrangements. Transcripts were classified as: unequivocally associated with the locus, if their only genomic match in terms of sequence identity and coverage corresponded to the locus with variable ITR; best associated with the locus, if the best genomic match corresponded to the locus with variable ITR; or multiply associated if they had multiple best matches on the genome.

**Groups of paralogs, age of SDs and overlap with human CNVs**

For each of the 2,008 gene loci, the number of paralogs was defined as the number of loci associated by at least one nearly identical exon (> 95% coverage, > 90% identity). The resulting distributions of paralogs between the 210 loci with variable ITRs and the remaining 1,798 loci were compared, using the non-parametric Wilcoxon test. The percentage of identity between pairs of SDs was recovered directly from UCSC. A non-redundant set of 25,914 pair-wise alignments between SDs was derived, 496 of which involve the 210 loci with variable ITRs. The genomic coordinates of 21,178 human CNVs grouped in 6,558 non-overlapping CNV loci were recovered from the Database of Genomic Variants [67,68] (version 7, March 2009).
Orthology assignment, tissue expression and functional enrichment

Orthologous regions corresponding to the human ITR-containing exons were extracted from the pair-wise BlastZ alignments between human and mouse (mm9) and human and dog (canFam2) using Galaxy [69]. The human/dog alignments were screened only in case the alignment between human and mouse was not available. The portions of the alignments corresponding to variable ITRs were manually checked. For 377 out of the 524 mRNAs with variable ITRs and for the 5,256 out of 8,638 mRNAs in 2,008 loci it was possible to extract information on the tissue type directly from GenBank. Tissues that were represented by at least 15 mRNAs with variable ITRs (> 3%) were selected for chi-squared comparison between transcripts with variable ITRs and other transcripts in SDs. Forty-nine percent of genes with variable ITRs can be associated with functional categories according to the Gene Ontology [70]. The functional enrichment was measured in comparison to other genes hosted in SDs. The functional terms in common between the two groups at levels 3 to 9 of the Gene Ontology hierarchy were compared using the chi-squared test and P-values were adjusted using the Bonferroni correction for multiple testing.

Abbreviations

CNV: copy number variant; ITR: internal tandem repeat; SD: segmental duplication; TRF: Tandem Repeat Finder.

Authors’ contributions

FDC conceived and designed the study; ADG performed the experiments; ADG and FDC analyzed the data and wrote the paper.

Additional files

The following additional data are available with the online version of this paper: an Excel file containing genomic coordinates and transcription evidence supporting ITR-driven gene modifications (Additional file 1); a Word file containing properties of variable ITRs, their effect on coding sequences and groups of paralogs associated with exon and intron modifications (Additional file 2); an Excel file providing functional analysis of genes with variable ITRs (Additional file 3).

Acknowledgements

This work was supported by AIRC Start-Up grant and by the Italian Ministry of Health to FDC.

References

1. Bailey JA, Gu Z, Clark RA, Reinert K, Samonte RV, Schwartz S, Adams MD, Myers EW, Li PW, Eichler EE. Recent segmental duplications in the human genome. Science 2002, 297:1003-1007.

2. Korbel JO, Urban AE, Affourtit JP, Godwin B, Grubert F, Simons JF, Kim PM, Palejev D, Carriero NJ, Du L, Taillon BE, Chen Z, Tanzer A, Saunders AC, Chi J, Yang F, Carter NP, Hurles ME, Weissman SM, Harkins TT, Gerstein MB, Egholm M, Snyder M. Paired-end mapping reveals extensive structural variation in the human genome. Science 2007, 318:420-426.

3. Redon R, Ishikawa S, Fitch KR, Feuk L, Perry GH, Andrews TD, Fiegler H, Shapoero MH, Carson AR, Chen W, Cho EK, Dallaire S, Freeman JL, Gonzalez JR, Gratasco M, Huang J, Kalaitzopolous D, Komura D, MacDonald JR, Marshall CR, Mei R, Montgomery L, Nishimura K, Okamura K, Shen D, Somerville MJ, Tchinda J, Valsea S, Woodward C, Yang F, et al.: Global variation in copy number in the human genome. Nature 2006, 444:444-454.

4. Wong KK, deLeeuw RJ, Dosanjh NS, Kimm LR, Cheng Z, Horsman DE, MacAsayu C, Ng RT, Brown CJ, Eichler EE, Lann WL: A comprehensive analysis of common copy-number variations in the human genome. Am J Hum Genet 2007, 80:91-104.

5. Eichler EE. Recent duplication, domain accretion and the dynamic mutation of the human genome. Trends Genet 2001, 17:461-469.

6. Zhang L, Lu HHS, Chung W-y, Yang J, Li W-H: Patterns of segmental duplication in the human genome. Mol Biol Evol 2005, 22:135-141.

7. Bailey JA, Eichler EE: Primate segmental duplications: crucibles of evolution, diversity and disease. Nat Rev Genet 2006, 7:552-564.

8. Lynch M, Conery JS: The evolutionary fate and consequences of duplicate genes. Science 2000, 290:1151-1155.

9. Long M, Betran E, Thornton K, Wang W: The origin of new genes: glimpses from the young and old. Nat Rev Genet 2003, 4:865-875.

10. Ciccarelli FD, von Mering C, Suyama M, Harrington ED, Izaurrelde E, Bork P: Complex genomic rearrangements lead to novel primate gene function. Genome Res 2005, 15:343-351.

11. Johnson ME, Viggiano L, Bailey JA, Abdul-Rauf M, Goodwin G, Rocchi M, Eichler EE: Positive selection of a gene family during the emergence of humans and African apes. Nature 2001, 413:514-519.

12. Birtle Z, Goodstadt L, Ponting C: Duplication and positive selection among hominin-specific PRAME genes. BMC Genomics 2005, 6:120.

13. Semple C, Rolfe M, Dorin J: Duplication and selection in the evolution of primate beta-defensin genes. Genome Biol 2003, 4:R31.

14. Fantoni I, Meini N, Rambaldi D, Scaffegata E, Alcayde M, Ciccarelli FD: Family expansion and gene rearrangements contributed to the functional specialization of PRDM genes in vertebrates. BMC Evol Biol 2007, 7:187.

15. Taylor JS, Raes J: Duplication and divergence: the evolution of new genes and old ideas. Annu Rev Genet 2004, 38:615-643.

16. Ellegren H: Microsatellites: simple sequences with complex evolution. Nat Rev Genet 2004, 5:435-445.

17. Katti MV, Ranjekar PK, Gupta VS: Differential distribution of simple sequence repeats in euchromatic genome sequences. Mol Biol Evol 2001, 18:161-167.

18. Toth G, Gaspari Z, Jurka J: Microsatellites in different eukaryotic genomes: survey and analysis. Genome Res 2000, 10:967-981.

19. Kashi Y, King DG: Simple sequence repeats as advantageous mutators in evolution. Trends Genet 2006, 22:253-259.

20. Verstrepen KJ, Jansen A, Lewitter F, Fink GR: Intrageneric tandem repeats generate functional variability. Nat Genet 2005, 37:986-990.

21. Fondon JW, Garner HR: Molecular origins of rapid and continuous morphological evolution. Proc Natl Acad Sci USA 2004, 101:18058-18063.

22. Hammock EA, Young Lj: Microsatellite instability generates diversity in brain and sociobehavioral traits. Science 2005, 308:1630-1634.

23. Jensen-Seaman Ml, Li WH: Evolution of the hominoid semenogelin genes, the major proteins of ejaculated semen. J Mol Evol 2003, 57:261-270.

24. Metzgar D, Bycroft J, Wills C: Selection against frameshift mutations limits microsatellite expansion in coding DNA. Genome Res 2000, 10:72-80.

25. Gatche JR, Zoghbi HY: Diseases of unstable repeat expansion: mechanisms and common principles. Nat Rev Genet 2005, 6:743-755.

26. Usdin K: The biological effects of simple tandem repeats: lessons from the repeat expansion diseases. Genome Res 2008, 18:1011-1019.

27. Yu F, Sabeti PC, Hardenbol P, Fu Q, Fry B, Lu X, Ghose S, Vega R,
45. Tusnady GE, Simon I: The HMMTOP transmembrane topology prediction server. Bioinformatics 2001, 17:849-850.

46. Bryson K, McGufflin LJ, Marsden RL, Ward JH, Siddhi JS, Jones DT: Protein structure prediction servers at University College London. Nucleic Acids Res 2005, 33:W349-W351.

47. Andrade MA, Perez-eraso C, Ponting CP: Protein repeats: structures, functions, and evolution. J Struct Biol 2001, 134:117-131.

48. Richards RI, Sutherland GR: Dynamic mutations: a new class of mutations causing human disease. Cell 1992, 70:709-712.

49. Gibbs RA, Weinstock GM, Metzker ML, Muzny DM, Sodergren EJ, Scherer S, Scott G, Steffen D, Worley KC, Burch PE, Okwuonu G, Hines L, Lewis S, DeRamo C, Delgado O, Dugan-Rocha S, Miner G, Morgan M, Hawes A, Gill R, Celera , Holt RA, Adams MD, Amanatides PG, Baden-Tillson H, Barnstead M, Chin S, Evans CA, Ferreria S, Fosler C, et al.: Genome sequence of the Brown Norway rat yields insights into mammalian evolution. Nature 2004, 428:493-512.

50. Waterston RH, Lindblad-Toh K, Birney E, Rogers J, Abril JF, Agarwal P, Ainscough R, Alexandersson M, An P, Antonarakis SE, Astwood J, Baertsch R, Bailey J, Barlow K, Beck S, Berry E, Birren B, Bloom T, Bork P, Boucherby M, Bray N, Brent MR, Brown DG, Brown SD, Built C, Burton J, Campbell RD, Carninci P, et al.: Initial sequencing and comparative analysis of the mouse genome. Nature 2004, 429:520-562.

51. Webster MT, Smith NG, Ellegren H: Microsatellite evolution inferred from human-chimpanzee genomic sequence alignments. Proc Natl Acad Sci USA 2002, 99:8748-8753.

52. Ellegren H: Heterogenous mutation processes in human microsatellites DNA sequence from human chromosome 16p and 16q. Nat Genet 1999, 24:400-402.

53. Conrad DF, Pinto D, Redon R, Feuk L, Gokcumen O, Zhang Y, Aerts M, Andrews TD, Barnes C, Campbell P, Fitzgerald T, Hu M, Ihm CH, Kristiansson K, MacArthur DG, MacDonald JR, Onyiah I, Pang AWC, Robinson S, Stirrup S, Troyanska S, Walther K, White C, Smith CR, Byers DR, Lee H, Scherer SW, Hurles ME: Origins and functional impact of copy number variation in the human genome. Nature 2009 in press.

54. Catania F, Lynch M: Where do introns come from? PLoS Biol 2008, 6:e282.

55. Rogers JH: How were introns inserted into nuclear genes? Trends Genet 1989, 5:213-216.

56. Venekatesh B, Ying Y, Brenner S: Late changes in spliceosomal introns define clades in vertebrate evolution. Proc Natl Acad Sci USA 1999, 96:10257-10261.

57. Knowles DG, Mclysaght A: High rate of recent intron gain and loss in simultaneously duplicated Arabidopsis genes. Mol Biol Evol 2006, 23:1548-1557.

58. Roy SW, Gilbert W: The evolution of spliceosomal introns: patterns, puzzles and progress. Nat Rev Genet 2006, 7:27-36.

59. Knowles DG, Mclysaght A: High rates of recent intron gain and loss in simultaneously duplicated Arabidopsis genes. Mol Biol Evol 2006, 23:1548-1557.

60. Roy SW, Penny D: Smoke without fire: most reported cases of intron gain in nematodes instead reflect intron losses. Mol Biol Evol 2006, 23:2259-2262.

61. Roy SW, Fedorov A, Gilbert W: Large-scale comparison of intron positions in mammalian genomes shows intron loss but no gain. Proc Natl Acad Sci USA 2003, 100:7158-7162.

62. Coulombe-Huntington J, Majewski J: Characterization of intron loss events in mammals. Genome Res 2007, 17:23-32.

63. Zhou D, Madden R, Etela SA, Chabot B: Modern origin of numerous alternatively spliced human introns from tandem arrays. Proc Natl Acad Sci USA 2007, 104:882-886.

64. Roy SW, Irimia M: When good transcripts go bad: artificial RT-PCR ‘splicing’ and genome analysis. Bioessays 2008, 30:601-605.

65. Uscs Genome Bioinformatics [http://genome.ucsc.edu/]

66. Kent WJ: BLAT - the BLAST-like alignment tool. Genome Res 2002, 12:656-664.

67. Benson G: Tandem repeats finder: a program to analyze DNA sequences. Nucleic Acids Res 1999, 27:573-580.

68. Database of Geneic Variants [http://projects.tcg.ca/variation/]

69. Iwafune J, Feuk L, Rivera MN, Listewnik ML, Donahoe PK, Qi Y, Scherer SW, Lee C: Detection of large-scale variation in the human genome. Nat Genet 2004, 36:949-951.

70. Galaxy [http://main.b2.bio.psu.edu/]

71. Ashburner M, Ball CA, Blake JA, Botstein D, Butler H, Cherry JM, Davis AP, Dolinski K, Dwight SS, Eppig JT, Harris MA, Hill DP, Issel-Tarver L, Kasarskis A, Lewis S, Matese JC, Richardson JE, Ringwald M, Rubin GM, Sherlock G: Gene ontology: tool for the unification of biology. The Gene Ontology Consortium. Nat Genet 2000, 25:25-29.
71. Larkin MA, Blackshields G, Brown NP, Chenna R, McGettigan PA, McWilliam H, Valentin F, Wallace IM, Wilm A, Lopez R, Thompson JD, Gibson TJ, Higgins DG: Clustal W and Clustal X version 2.0. Bioinformatics 2007, 23:2947-2948.