Evolutionary History of Alpha Satellite DNA Repeats Dispersed within Human Genome Euchromatin

Isidoro Feliciello¹,²,*, Željka Pezer¹, Dušan Kordić³, Branka Bruvo Mađarić¹, and Đurđica Ugarković¹,*

¹Department of Molecular Biology, Ruđer Bošković Institute, Zagreb, Croatia
²Dipartimento di Medicina Clinica e Chirurgia, Università degli Studi di Napoli Federico II, Italy
³Department of Molecular and Biomedical Sciences, Jožef Stefan Institute, Ljubljana, Slovenia

*Corresponding authors: E-mails: ifelici@unina.it; ugarkov@irb.hr.

Accepted: 14 October 2020

Abstract

Major human alpha satellite DNA repeats are preferentially assembled within (peri)centromeric regions but are also dispersed within euchromatin in the form of clustered or short single repeat arrays. To study the evolutionary history of single euchromatic human alpha satellite repeats (ARs), we analyzed their orthologous loci across the primate genomes. The continuous insertion of euchromatic ARs throughout the evolutionary history of primates starting with the ancestors of Simiformes (45–60 Ma) and continuing up to the ancestors of Homo is revealed. Once inserted, the euchromatic ARs were stably transmitted to the descendant species, some exhibiting copy number variation, whereas their sequence divergence followed the species phylogeny. Many euchromatic ARs have sequence characteristics of (peri)centromeric alpha repeats suggesting heterochromatin as a source of dispersed euchromatic ARs. The majority of euchromatic ARs are inserted in the vicinity of other repetitive elements such as L1, Alu, and ERV or are embedded within them. Irrespective of the insertion context, each AR insertion seems to be unique and once inserted, ARs do not seem to be subsequently spread to new genomic locations. In spite of association with (retro)transposable elements, there is no indication that such elements play a role in ARs proliferation. The presence of short duplications at most of ARs insertion sites suggests site-directed recombination between homologous motifs in ARs and in the target genomic sequence, probably mediated by extrachromosomal circular DNA, as a mechanism of spreading within euchromatin.

Key words: satellite DNA, euchromatin, heterochromatin, evolution, proliferation, primates.

Significance

Satellite DNAs are major constituents of pericentromeric and centromeric regions in many eukaryotes and their role in centromere and kinetochore assembly and heterochromatin formation has been extensively investigated. However, the role of satellite repeats found dispersed in euchromatin, outside of centromere/pericentromere regions, remains largely unexplored. Here, we analyze the dynamics of dispersion of human alpha repeats throughout euchromatin during the evolutionary history of primates and the mechanism of their proliferation. The results contribute to our understanding of the possible evolutionary and functional significance of satellite DNA repeats spread throughout euchromatin.

Introduction

Satellite DNAs are tandemly repeated sequences assembled in large arrays within constitutive heterochromatin in (peri)centromeric and/or telomeric regions of eukaryotic chromosomes. Within euchromatin, longer arrays of tandem satellite repeats are generally rare, probably due to the instability caused by intrastrand homologous recombination, although blocks of euchromatic tandem repeats have been found in several species (Kuhn et al. 2012; Pavlek et al. 2015; Pita et al. 2017; Vlahovic et al. 2017). Bioinformatic
analyses of sequenced genomes however reveal many single repeats or short arrays of satellite DNAs dispersed in the vicinity of genes within euchromatin in diverse species such as mouse (Bulut-Karsioglu et al. 2012) or insects (Brajković et al. 2012, 2018; Kuhn et al. 2012; Ruiz-Ruano et al. 2016). The pattern of dispersion of satellite DNA repeats within euchromatin is very dynamic and differs among related species or even among strains of the same species, as shown for Drosophila and Tribolium castaneum satellite DNAs, respectively, suggesting that similar to transposable elements, euchromatic satellite repeats are subjected to cycles of proliferation (Feliciello, Akrap, Brajković, et al. 2015; Feliciello, Akrap, Ugarković 2015; Sproul et al. 2020). Some euchromatic satellite repeats such as those of a major satellite DNA in the beetle Tribolium castaneum modulate the local chromatin environment upon heat stress, affecting the expression of neighboring genes (Feliciello, Akrap, Ugarković 2015). Moreover, euchromatic repeats seem to be in spatial contact with heterochromatin, suggesting that the interplay between euchromatic and heterochromatic repeats could play a role in gene expression modulation (Feliciello, Akrap, Ugarković 2015; Lee et al. 2020). In the mosquito Aedes aegypti, satellite repeats located at a single euchromatic locus promote sequence-specific gene silencing via the expression of abundant PWI-interacting RNAs (piRNAs; Halbach et al. 2020). Euchromatic satellite repeats also facilitate X chromosome recognition/dosage compensation in Drosophila (Menon et al. 2014; Joshi and Meller 2017).

Alpha satellite DNA makes up to 10% of the human genome, it is located in the centromeric and pericentromeric regions of all chromosomes, contributing to essential chromosomal functions such as centromere and kinetochore assembly and heterochromatin formation (McNulty and Sullivan 2018). Based on its wide presence among primates (Willard 1991; Alexandrov et al. 2001; Cacheux et al. 2016) and according to primate phylogeny (Finstermeier et al. 2013; Pozzi et al. 2014), the age of alpha satellite DNA could be estimated to ~65–70 Myr. The fundamental unit of human alpha satellite DNA is based on diverged 171-bp monomers estimated to ~100–200-bp flanking sequences at 5′ and 3′ site were used for detection of orthologous loci by UCSC Genome Browser and syntenicity of flanking sequences for each AR was examined to confirm orthologous loci across primate species.

Annotation of human ARs into suprachromosomal families and monomer classes was retrieved using: https://genome.ucsc.edu/cgi-bin/hgTracks?db=hg38&htg.customText=https://raw.githubusercontent.com/enigene/AS-tracks/master/GRCh38-GCA_000001405.15/human-GRCh38-M1Sfsv2.2.bed.gz.

Analysis of Junction Regions
Analysis of the junction regions on both sides of ARs annotated by RepeatMasker, with 100–200-bp flanking sequences at 5′ and 3′ site were used for detection of orthologous loci by UCSC Genome Browser and syntenicity of flanking sequences for each AR was examined to confirm orthologous loci across primate species.
et al. 2015) to computationally detect motifs that are present at the 5’ and 3’ junction regions of ARs.

Statistical Analyses

We used bedtools (Quinlan and Hall 2010) and custom scripts to analyze content of repetitive element classes. Tables ncbiRefSeqCurated and rmsk corresponding to genes and repetitive elements were downloaded from UCSC Genome Browser. Statistical analyses were performed in R. Random introns were chosen such that their length was between 5 and 300 kb, matching the range of ARs-containing introns. For analysis of repetitive classes surrounding the intergenic ARs, we extended coordinates of ARs upstream and downstream by 20 kb and intersected the intervals with rmsk data.

Phylogenetic Analysis

Alignments of ARs were performed online with MAFFT version 7, using “Auto” or “E-INS-i” strategy (Katoh et al. 2019; https://mafft.cbrc.jp/alignment/server/index.html). Junction regions were removed prior to alignment. Uncorrected p-distances were calculated using MEGA 7.0.25 (Kumar et al., 2016). Neighbor-Joining (NJ) trees based on the p-distance model were calculated in MEGA 7.0.25 (Kumar et al. 2016), and the robustness of the cladess was assessed through 1,000 bootstrap replicates. Maximum-likelihood (ML) trees were constructed in PhyML 3.0 web-server (Guindon et al. 2016), and the robustness of the cladest was calculated through 1,000 bootstrap replicates. Maximum-likelihood (ML) trees were constructed in PhyML 3.0 web-server (Guindon et al. 2010), with automatic model selection by SMS (determined through AIC selection criterion) (Lefort et al. 2017) and aLRT SH-like support (Anisimova and Gascuel 2006). Resulting trees were edited in FigTree v.1.4.3. (http://tree.bio.ed.ac.uk/software/figtree/). To check for higher order repeat organization, we produced self-dot plots of sequences by using Flexidot (Seibt et al. 2018) and setting the word size to 25 bp.

Synteny Analysis

To analyze the order of repetitive elements in the vicinity of ARs and compare it between primate species, we first extended the genomic coordinates of each AR in the human assembly: for intergenic AR elements coordinates were extended 2-kb up- and downstream and for intronic elements the coordinates of the whole intron were considered. We used these extended coordinates to find orthologous regions with the LiftOver program (Hinrichs et al. 2006). LiftOver was run on 16 existing primate chain files for assemblies other than human, that were downloaded from http://hgdownload.soe.ucsc.edu/goldenPath/hg38/liftOver: PanTro6, PanPan2, GorGor5, ChiSaB2, PonAbe3, NomLeu3, MacFas5, RHEMAC10, PapAnu4, NasLar1, RhRox1, CalJac3, SaiBo1, TarSyr2, MicMUr2, and OttoGar3. The resulting coordinates were intersected with coordinates of annotated repetitive elements (rmsk tables) of the respective species downloaded from the UCSC ftp site (ftp://hgdownload.soe.ucsc.edu/goldenPath/). The resulting annotated repetitive elements that overlapped given coordinates were plotted with genoPlotR package (Guy et al. 2010).

Gene Ontology Analysis

For gene ontology analysis, the following tools were used: GOIrilla (http://cbl-gorilla.cs.technion.ac.il/; Eden et al. 2009) and Panther (http://www.pantherdb.org/; Mi et al. 2019).

Results

Dispersed ARs in Human Genome Euchromatin

ARs annotated in the human genome assembly hg19 were extracted from the rmsk table from the UCSC Table Browser. In this way, 1,287 ARs were identified in the assembled chromosomes, ranging in size from 14 to 160,602 bp. Because it is possible that some short ARs represent false positives, only hits with at least 50% of 171-bp monomer sequence length were considered for further analyses. Most of the ARs are located within centromeric and pericentromeric regions and 1,071 are organized within clusters which are composed of 2–52 ARs positioned within a distance <1 kb. In order to study the evolutionary history and mode of dispersion of ARs within euchromatin, we focused on ARs located outside of (peri)centromeric regions, preferentially on those not organized within clusters but present as single repeats (supplementary table 1, Supplementary Material online). For such ARs, it is possible to detect orthologous regions within other primate genomes in order to date their insertion and to follow the dynamics throughout the evolutionary history of primates. The presence and organization of euchromatic ARs was additionally checked in the human genome assembly GRCh38.hg38, revealing them on almost all human chromosomes.

There are 32 euchromatic ARs which overlap with genes, all located in introns of 18 protein coding genes, two ncRNAs, and one pseudogene. Most intronic ARs are short, between 0.5 and 4 mer size, except three ARs of several kb size within the ANKRD30BL gene, and are preferentially organized as single repeats except in three cases where clusters of two, three, and seven repeats are found within genes LINC01580, ANKRD30BL, and SACS, respectively (table 1). In addition to intronic ARs, we analyzed the evolutionary history of 36 intergenic ARs, 29 of them organized as single repeats, whereas the others form clusters composed of a few adjacent repeats (table 1 and supplementary table 1, Supplementary Material online). By examination of clustered ARs, we expected to reveal if the ARs within the same cluster share a common evolutionary history, meaning that they were dispersed simultaneously, possibly together within a single insertion event, or separately during different evolutionary periods. The genes containing intronic ARs as well as genes most proximal to intergenic alpha repeats and their distances
### Table 1

List of Euchromatic Human ARs Used in This Study

| AR No. | Size | Monomer Type | Position (Associated Gene), Distance (bp) | Insertion Site Characteristics | Dating of Insertion | TS-Duplicated Sequence |
|--------|------|--------------|------------------------------------------|-------------------------------|---------------------|------------------------|
| sat_1  | 1.3  | Intron (AGDR121), –240,820 | Adjacent to L1PA10 | Simiformes | CTT |
| sat_83 | 2.4  | M1+, X | Intron (PLA2G12B) | Btw L1MB8 and L2c | Simiformes | AGGAGT |
| sat_368| 0.8  | Intron (MYCBP2), –73,091 | Btw AluSx6 and L1ME1 | Simiformes | — |
| sat_380| 1    | R1 | Intron (LINC01580), ncRNA | Adjacent to MIR (SINE) | Simiformes | GAA |
| sat_381| 0.9  | R1 | Intron (LINC01580), ncRNA | Adjacent to MIR (SINE) | Simiformes | GAA |
| sat_496| 1.4  | Intron (AVH1) | Btw AluSx1 and MLT1C (ERV1) | Simiformes | AAG |
| sat_623| 0.5  | | Intron (PLCB4) | No adjacent repeats | Simiformes | — |
| sat_703| 1    | X, M, X | Intron (ACTL6A), 29,969 | L1MC1 (5900) | Simiformes | TGA |
| sat_704| 1.4  | Intron (RPL392), 15,390 | No adjacent repeats | Simiformes | AGTG |
| sat_685| 1.9  | Intron (FILIP11) | Adjacent to L1PA11 | Simiformes | AAA |
| sat_730| 0.5  | Um | Intron (MIR297), –630,154 | Btw L1PA8A and simple repeat | Simiformes | ATGAAAAAAA |
| sat_721| 0.9  | Intron (RPL21P44), 37,365 | Btw TAn and ATG6n | Simiformes | ATAT |
| sat_722| 1    | | Intron (LOC105372247), 129,191 | Btw (ATAT)n and FordPrefect hAT | Simiformes | GCTA |
| sat_825| 0.7  | Intron (GABRB2), 125,243 | Btw L1MB2 and AluSx4 | Simiformes | GAAA |
| sat_826| 0.8  | Intron (C6orf106), 7,067 | No adjacent repeats | Simiformes | — |
| sat_827| 0.7  | M1+ | Intron (FAM83B), –46,144 | Tigger3 (DNA; 205) | Simiformes | GCT |
| sat_828| 1.4  | | Intron (PRIM2) | Adjacent to AluR | Simiformes | GAAAAAG |
| sat_1122| 0.5  | | Intron (TG), 2,283 | No adjacent repeats | Simiformes | GTGA |
| sat_1147| 1.5 | M1+ | Intron (MAP7) | L1ME3 (5618) | Simiformes | ATC |
| sat_60 | 0.5  | | Intron (STAM) | MER778 (ERV1; 311) | Catarrhini | AT |
| sat_85 | 0.6  | | Intron (LRRC4C) | Adjacent to MIR (SINE) | Catarrhini | — |
| sat_87 | 0.5  | | Intron (LRRC4C) | Adjacent to MERSA hAT-Charlie | Catarrhini | GAG |
| sat_497| 0.6  | M1+ | Intron (ZNF675) | AluSc (73) | Catarrhini | — |
| sat_621| 1.8  | M1+ | Intron (SLC40A1), –33,821 | No adjacent repeats | Catarrhini | — |
| sat_605| 0.7  | | Intron (SLC30A6) | AluSx3 (169) | Catarrhini | AGA |
| sat_606| 0.5  | | Intron (LINCO0486), ncRNA | No adjacent repeats | Catarrhini | CTT |
| sat_612| 0.7  | | Intron (TMEM131) | L1M2c (2147) | Catarrhini | GCT |
| sat_705| 1.4  | | Intron (AFAP1) | Adjacent to L1PA10 | Catarrhini | — |
| sat_373| 0.9  | | Intron (ERO1A) | No adjacent repeats | Catarrhini | GTTTT |
| sat_706| 3.4  | M1+ | Intron (ACOX), 23,643 | Btw LTR188 and L2a | Catarrhini | TTA |
| sat_58 | 0.5  | | Intron (SUSD4), 12,805 | HERV771A-int (ERV1; 5439) | Hominoidea | AAC |
| sat_86 | 0.3  | | Intron (LRRC4C) | Adjacent to MERSA | Hominoidea | — |
| sat_358| 2.7  | M1+ | Intron (CPNE8), 59,985 | MLT1D (ERV1-MaLR LTR; 216) | Hominoidea | TCAC |
| sat_380| 1.5  | | Intron (SACS) | Adjacent to AluYk | Hominoidea | AGA |
| sat_361| 0.9  | | Intron (SACS) | Adjacent to AluYk | Hominoidea | AGA |
| sat_362| 3.7  | | Intron (SACS) | Adjacent to AluYk | Hominoidea | TCA |
| sat_363| 1    | | Intron (SACS) | Adjacent to AluYk | Hominoidea | TTGT |
| sat_364| 1.2  | | Intron (SACS) | Adjacent to AluSp | Hominoidea | ACT |
| sat_365| 0.8  | | Intron (SACS) | Adjacent to AluSp | Hominoidea | TAT |
| sat_366| 0.8  | | Intron (SACS) | Adjacent to AluSp | Hominoidea | AGCT |
| sat_367| 1.5  | | Intron (PARP4), –9,495 | No adjacent repeats | Hominoidea | TGT |
| sat_729| 0.7  | M1+ | Intron (CXCL13), 22,197 | No adjacent repeats | Hominoidea | TGT |
| sat_378| 1.8  | | Intron (SNRPN), 69,307 | | Hominoidea | AAG |
| sat_410| 0.5  | | Intron (LINCO1566), 36,721 | Btw L1M3 and AluY | Hominoidea | GAT |
| sat_652| 5    | M1+ | Intron (PRAMENP), –90,051 | Btw AluSx2 and AluSc | Hominoidea | AC |
| sat_653| 27   | M1+, Um | Intron (PRAMENP), –91,217 | Btw AluSc and (TG)n | Hominoidea | AC |
| sat_723-25 | 32 | M1+ | Intron (TECRL), 949,009 | Btw MSTA (ERV1) and L1PA4 | Hominoidea | TTG |
| sat_726 | 31 M1+ | Intron (TECRL), 933,498 | Btw L1PA4 and MER11C (ERV1) | | Hominoidea | — |
| sat_727 | 15 M1+ | Intron (TECRL), 929,843 | Adjacent to MER11C (ERV1) | | Hominoidea | TTG |
| sat_823 | 0.7 | | Intron (LINCO2159), 101,431 | Btw 2 L1PA7 | Hominoidea | TAA |
| sat_732 | 11 M1+ | Intron (C5orf17), 64,041 | Adjacent to AluY | Hominoidea | AAACCTG |

(continued)
Table 1 Continued

| AR No. | Size | Monomer Type | Position (Associated Gene), Distance (bp) | Insertion Site Characteristics | Dating of Insertion | TS-Duplicated Sequence |
|--------|------|--------------|------------------------------------------|-------------------------------|---------------------|-----------------------|
| sat_733 | 26   | M1+          | Intergenic (C5ORF17), 59,303             | Btw AluY and L1PA7           | Hominidae           | TG                    |
| sat_864 | 2    | M1+          | Intergenic (MAF), 11,153                 | Btw MER21C and MLT2B3 (ERV)  | Hominidae           | TTGG                  |
| sat_59  | 0.5  | M1+          | Intergenic (DIP2C), –18,185             | Btw 2 MSTD-int ERV           | Hominidae           | CTA                   |
| sat_257 | 0.7  | –            | Intron (DLG2)                            | Adjacent to L1PA8            | Hominidae           | CAT                   |
| sat_372 | 5.9  | M1+          | Intergenic (LINC01296), –19,382         | No adjacent repeats          | Hominidae           | ACAT                  |
| sat_379 | 1.2  | Um           | Intron (MYO1E)                           | No adjacent repeats          | Hominidae           | ACT                   |
| sat_607 | 0.9  | R2, X        | Intergenic (TGFA), 15,683               | Btw L1PA2 and Charlie8       | Hominidae           | AAC                   |
| sat_731 | 0.5  | M1+          | Intergenic (LIN00613), 290,201          | Adjacent to AluX1            | Hominidae           | CTCCAA                |
| sat_822 | 1.1  | M1+          | Intron (NRCIC)                          | Btw AluSc and Alu            | Hominidae           | CTT                   |
| sat_1123 | 4.1 | D2, M1       | Intron, LOC101928195-pseudogene        | Btw MamGypLTR3a and L1PA3   | Hominidae           | AACAG                 |
| sat_1145 | 4.1 | D2, M1       | Intergenic (LOC101928381), ncrRNA, 36,431 | Btw MamGypLTR3a and L1PA3 | Hominidae           | AACAG                 |
| sat_1146 | 0.9  | –            | Intergenic (PPPR23B), 15,781            | Adjacent to AluO             | Hominidae           | TGA                   |
| sat_84  | 1    | J1A          | Intergenic (OR6A2), 14,531              | L1PA16 (3051)                | Hominini            | GTA                   |
| sat_613 | 0.9  | R1           | Intergenic (DPPI), 58,435               | LTR43 (ERV; 506)            | Hominini            | CAT                   |
| sat_614 | 67   | M1, R2, X    | Intron (ANKRD30BL)                      | Adjacent to SVA, D-E        | Homo                | ACACTG                |
| sat_615 | 49   | R1–2, M1, Um | Intron (ANKRD30BL)                      | Btw SVA, D-E and L1PA3      | Homo                | GAA                   |
| sat_616–618 | 77  | D1–2, R, M  | Intron (ANKRD30BL)                      | Adjacent to L1PA3           | Homo                | ATAA                  |

Notes.—Size is expressed as number of monomers, monomer types are inferred according to Shevelev et al. [2015], and positions relative to genes are shown; negative distances mean 5’ position of a gene to the AR. The association with other repetitive elements at the insertion site and dating of the insertion is listed as well as target site (TS)-duplicated sequences at the insertion sites. For ARs inserted in other repetitive elements, position of insertion within the particular element is shown in parentheses. ARs embedded in other repeats are indicated in bold and clustered ARs in italic.

Orthologous Dispersed ARs in Primate Species—Dating of Insertions

We searched for sequence orthologs to all 32 intronic and 36 intergenic dispersed human ARs across the primate genomes in order to trace when their dispersion and insertion in the euchromatin occurred during the evolutionary history. The oldest insertions were traced back 45–60 Ma in the ancestors of Simiformes, which include Platyrhini (New World monkeys) and Catarrhini (Old World monkeys and hominoids) (fig. 1a), and are characteristic for 11 single intergenic ARs as well as for six single and two clustered intronic ARs (table 1). The ARs inserted in the ancestors of Simiformes can be detected in almost all available genomes of the descendant species belonging to the branches of Platyrhini and Catarrhini, revealing that after insertion, ARs are transmitted and preserved in the descendant species. Some ARs exhibit copy number variation (CNV) such as sat_83 which is in the form of 2.3–2.4 mer in Homininae, gibbon, rhesus, and new world monkeys, whereas in baboon, green monkey, proboscis, and golden snub-nosed monkey, it is found as 3.4 mer and in orangutan as 1.4 mer (fig. 1a). The CNV of sat_83 can be explained by a single intrasstrand recombination event occurring between two homologous regions within AR monomers [supplementary fig. 1a, Supplementary Material online, and fig. 1b]. There is no correlation between the size of sat_83 and species phylogeny suggesting that CNV results from the process of recombination which occurred several times independently during the evolutionary history, but at the same specific site of the sequence. According to species phylogeny, a recombination event which resulted in CNV probably occurred in the ancestor of Platyrrhini (3.4–2.4 mer), in the ancestor of Hominoidea (3.4–2.4 mer), after separation of orangutan from Homininae (2.4–1.4 mer) and in the ancestor of the rhesus–macaque group of old world monkeys (3.4–2.4 mer) (fig. 1a).

The next round of AR insertions can be traced 30–45 Ma in the ancestors of Catarrrhini and includes nine intronic and two intergenic ARs (table 1). It is interesting that the intergenic sat_706 is detected as a 1.2 mer in old world monkeys (Cercopithecidae), whereas in Hominioidea it is 2.2 mer in gibbon, 4.4 mer in orangutan, and 3.4 mer in gorilla, bonobo, chimpanzee, and human (fig. 1b). Alignment of sat_706 sequences in different primates indicates that CNV could result from intrasstrand recombination facilitated by homologous sequence motifs in multimers [supplementary fig. 1b, Supplementary Material online, and fig. 1b]. The origin of sat_706 in Hominioidea can be explained by a single intrasstrand recombination occurring between two homologous regions within monomers, whereas the genesis of a 1.2 mer in old world monkeys is proposed to be due to an additional recombination event (fig. 1b). According to the phylogeny, recombination occurred several times through evolutionary history: in the common ancestor of old world monkeys (4.4–1.2 mer), in the ancestor of Hylobatidae (4.4–2.2 mer), and in the ancestor of Homininae (4.4–3.4 mer) (fig. 1a).
Another round of insertion is dated 20–30 Ma in the ancestors of Hominoidea (Hominidae, Hylobatidae) and is characteristic for seven clustered ARs within an intron of the SACS gene as well as for a single intronic and four intergenic ARs (Table 1 and fig. 1a). The AR sat_378 exhibits CNV, in gibbon (Hylobatidae) it is a 0.9 mer, whereas in Hominidae it is a 1.8 mer (orangutan, human, and chimp) except in gorilla and bonobo where a larger region encompassing sat_378 is deleted. The excision of the monomer in gibbon can be explained by a single internal recombination occurring between two homologous regions.

Insertions in the ancestors of Hominidae (orangutan, gorilla, pan, and human) 18–20 Ma are characteristic for four intergenic ARs (Table 1). The intergenic sat_372 shows CNV: it is 5.9 mer in human, 3.9 mer in bonobo and chimp, 4.9 mer in gorilla, and 10.9 mer in orangutan and can be explained by a single intrastand recombination event which occurred independently several times, in the lineages of gorilla (10.9–4.9

**Table 1**

| AR     | Homo sapiens | Pan troglodytes | Pan paniscus | Gorilla gorilla gorilla | Pongo pygmaeus abelii | Nomascus leucogenys | Macaca mulatta | Macaca fascicularis | Papio anubis | Chlorocebus sabaeus | Rhinopithecus roxellana | Nasalis larvatus | Callithrix jacchus | Saimiri boliviensis |
|--------|--------------|-----------------|--------------|------------------------|-----------------------|---------------------|------------------|---------------------|--------------|---------------------|------------------------|-----------------|----------------|-------------------|
| sat_83 | 2.4 mer      | 3.4 mer         | 2.3 mer      | 2.3 mer                | 3.4 mer               | 3.4 mer             | 1.4 mer          | 4.4 mer             | 2.4 mer      | 2.2 mer             | 1.2 mer                | 2.4 mer         | 2.4 mer         | 2.2 mer           |
| sat_706| 3.9 mer      | 3.4 mer         | 3.4 mer      | 4.9 mer                | 4.9 mer               | 4.9 mer             | 4.9 mer          | 10.9 mer            | 2.2 mer      | 1.2 mer             | 1.2 mer                | 1.2 mer         | 1.2 mer         | 1.2 mer           |
| sat_372| 5.9 mer      | 3.9 mer         | 3.9 mer      | 10.9 mer               | 10.9 mer              | 10.9 mer            | 10.9 mer         | 10.9 mer            | 3.9 mer      | 3.9 mer             | 3.9 mer                | 3.9 mer         | 3.9 mer         | 3.9 mer           |

**Figure 1**

(a) Number of single human ARs inserted within euchromatin during evolutionary history is indicated in red on the phylogenetic tree of simians (Simiformes). Species for which genome sequence is available are indicated: human (*Homo sapiens*), chimp (*Pan troglodytes*), bonobo (*Pan paniscus*), gorilla (*Gorilla gorilla gorilla*), orangutan (*Pongo pygmaeus abelii*), gibbon (*Nomascus leucogenys*), rhesus (*Macaca mulatta*), crab-eating macaque (*Macaca fascicularis*), baboon (*Papio anubis*), green monkey (*Chlorocebus sabaeus*), golden snub-nosed monkey (*Rhinopithecus roxellana*), proboscis monkey (*Nasalis larvatus*), marmoset (*Callithrix jacchus*), and squirrel monkey (*Saimiri boliviensis*). For alpha repeats sat_83, sat_706 and sat_372 which show CNV, size of repeats in each species is indicated. (b) Schematic representation of intrastrand recombination responsible for CNV of alpha repeats sat_83 and sat_706. Homologous regions in AR sequences at which recombination occurred are indicated by different signs. In sat_83, there is a single recombination site, whereas in sat_706 there are two sites.
mer), chimp/bonobo (10.9–3.9 mer), and human (10.9–5.9 mer) (fig. 1a). Three intergenic clusters examined, composed of two and three ARs respectively, were also inserted in the ancestors of Hominidae (table 1) and remained preserved in all species except chimpanzee where clustered sat_726–sat_727 was removed as a part of larger deletion.

Insertions occurring 10–18 Ma in the ancestors of Homininae (gorilla, pan, and human) include four intronic as well as six intergenic ARs (table 1). ARs sat_1145 and sat_1123 show CNV: 4.1 mer in human and 2.9 mer in gorilla, whereas in chimp the large region encompassing these ARs is deleted (table 1). The insertions occurring in the ancestors of Hominini (pan, human) ~7–10 Ma are two intergenic ARs, whereas the most recent insertion is of three clustered ARs within intron of the ANKRD30BL gene occurring in the ancestor of Homo (table 1 and fig. 1a).

The analysis of sequences orthologous to dispersed euchromatic human ARs across the primate genomes revealed their continuous insertion throughout evolutionary history of primates starting in the ancestors of Simiformes (45–60 Ma) and continuing up to the ancestors of Homo. Once inserted, the ARs were preserved and transferred to the descendant species, whereas some of them exhibited CNV due to recombination occurring independently in different evolutionary periods.

Phylogenetic Analysis and Sequence Evolution of Dispersed ARs

Phylogenetic analysis divides human monomeric ARs into different age groups, reflecting the evolution of centromeric alpha satellite DNA sequences in primates (Schueler and Sullivan 2006) which proceeds through proximal expansion of central active centromeric regions and moving of the previous centromeric DNA distally onto each arm (Schueler et al. 2005). According to that model, human alpha repeats proximal to the euchromatin chromosome arms are remnants of the ancestral primate centromere. Based on sequence features, human ARs are classified into 12 distinct monomer types which belong to five suprachromosomal families, SF1-5, and their annotation to classes and suprachromosomal families is available (Shepelev et al. 2015).

The analysis of human ARs dispersed in euchromatin shows that half of them are annotated as specific monomer types characteristic for particular suprachromosomal families (table 1). The annotated euchromatic ARs preferentially belong to SF4+ (monomer type M1+) and SF5 (monomer types R1 and R2) suprachromosomal families which are evolutionary old and constitute pericentromeric regions. Only a few ARs contain monomers of evolutionary new suprachromosomal families which are characteristic of active centromeres, such as sat_84 which belongs to the SF1 (monomer type J1A), whereas sat_1145 and sat_1123 as well as few kb size AR in intron of ANKRD30BL (sat_616–618) contain monomers D1 and D2 of the SF2. The ARs belonging to the new suprachromosomal families were inserted within the last 18 Myr (table 1).

In order to see if there is any sequence clustering of single dispersed human ARs that could indicate homogenization at the level of chromosome, neighboring or tandemly arranged repeats, sequence alignment and phylogenetic analyses were performed. Because almost half of the ARs are partial monomers, the alignment was performed on those repeats that mutually overlap in the sequences, whereas the others were excluded from the analysis. Finally, 77 partial or full sized monomers which derive from 57 dispersed human ARs were aligned (supplementary file 1a, Supplementary Material online). Phylogenetic analysis performed by the NJ and ML methods resulted in a phylogenetic tree with generally very weak resolution (supplementary fig. 2, Supplementary Material online). Only a few clusters composed of two ARs such as neighboring sat_380 and sat_381 as well as sat_1145 and sat_1123 are well supported (bootstrap values >0.7). Tandemly arranged monomers within sat_372 (5.9 mer) and sat_706 (3.4 mer) as well as clustered sat_360–366 are partially grouped in the phylogenetic tree, although the groups are not significantly supported. The average sequence divergence (p-distance) of human dispersed alpha satellite monomers is 0.3. We additionally checked the phylogeny of alpha monomers within three long clustered ARs (sat_614–618) in the intron of the ANKRD30BL gene, which were inserted relatively recently, in the ancestor of Homo. All three phylogenetic trees are characterized by weak resolution with very few supported groups of 2–3 monomers (supplementary fig. 3a and file 1b–d, Supplementary Material online). Although multiple monomers within sat_616–618 belong to the SF2 subfamily which is characteristic of active centromeres, the phylogenetic tree gave no indication for centromere-specific higher order organization of monomers and the self-dot plot confirmed this (supplementary fig. 3b, Supplementary Material online). The average sequence divergence of monomers within long clustered ARs in the ANKRD30BL gene is 0.20.

The sequence evolution of single ARs dispersed within euchromatin was followed in different primate species. Six ARs in introns (sat_828, sat_1147, sat_705, sat_685, and two adjacent ARs sat_380–381) and five in intergenic regions (sat_621, sat_704, sat_1, sat_703, and sat_722) were examined, all inserted in the ancestors of Simiformes and are therefore widely spread among extant species. In addition, the examined ARs are longer than a 1 mer and do not exhibit CNV among primate species. Average sequence divergences (uncorrected p-distances) between respective ARs of different primate species are generally low and range from 0.0468 in sat_1147 to 0.0835 in sat_685. NJ and ML phylogenetic trees reveal evolution of ARs’ sequences which generally supports the primate species phylogeny, in particular, major clades of new and old world monkeys as well of hominoids are
supported by high confidence, in addition to most of the
nodes within clades (fig. 2 and supplementary fig. 4 and file
1e–n, Supplementary Material online). The results reveal that
dispersed ARs’ sequences are not subjected to abrupt changes
and rearrangements except CNV as previously described but
evolve gradually, reflecting the evolution of primate species.

Association of ARs with Other Repetitive Elements and
Synteny Analysis

Over 80% of single ARs within human euchromatin have in
the vicinity, within 100-bp distance, other repetitive DNA ele-
ments (table 1). To see if there is any preference for specific
repetitive families to be nearby ARs, we analyzed association
of all 1,287 annotated ARs with annotated transposable ele-
ments (TEs). Out of 618 TEs found immediately next to ARs,
over half (329) belong to the L1 repeat family and 32% (197)
are Alu elements, whereas the majority of the rest are ERV
elements (~15%). Only 25 ARs actually overlap other TEs, the
majority intersecting ERV elements. To test if ARs are inserted
in regions enriched with specific repetitive elements what
might also suggest that such elements could mediate ARs
dispersion, we analyzed repeat composition of 22 introns
containing ARs and compared it with the composition of
100 randomly chosen introns without ARs (fig. 3a and b).
There is 3.6x higher variance of simple repeats proportions
in random introns (P value = 0.005; F test) compared with
ARs-containing introns and the only repetitive class “DNA”
(DNA transposons) is significantly underrepresented in introns
with ARs (P value = 0.0038; Welch Two Sample t-test), com-
prising on average about 8% of repetitive elements compared
with about 13% in randomly chosen introns. Overall, the
proportion of repetitive element classes seems to be less var-
iable within the 22 introns compared with arbitrarily chosen
introns. We also analyzed the proportion of repetitive

![ML trees based on human dispersed ARs, intronic sat_828 and sat_380-381 as well as intergenic sat_704 and sat_703 and their orthologous sequences in different primate assemblies: hg38-human, panTro6-chimp, panPan2-bonobo, gorGor5-gorilla, panAbe3-orangutan, nomLeu3-gibbon, chlSab2-green monkey, macFas5-crab-eating macaque, rheMac-rhesus, papAnu4-baboon, rhiRox1-golden snub-nosed monkey, nasLar1-proboscis monkey, calJac3-marmoset, and saIb01-squirrel monkey. Numbers on the nodes depict ML aLRT/NJ bootstrap support values.](image)
elements 20 kb around intergenic ARs listed in table 1 (fig. 3c) and found a substantially smaller proportion of SINE elements (26%) compared with ARs-containing introns (40%; P value = 0.0011; Welch Two Sample t-test). The average proportion of repeats in introns with ARs, randomly chosen introns or in intergenic regions around ARs is not significantly different from the average proportion of repetitive elements in the human genome overall (fig. 3d). The results indicate that preferential location of ARs near SINE, LINE, and ERV elements is more likely due to the general abundance of such repeat elements in introns and intergenic regions rather than to the specific association of ARs with these elements.

To study how the organization of repetitive elements within regions proximal to dispersed ARs evolves among primates, we analyzed the distribution of repetitive families within 20 kb of intergenic ARs, as well as within ARs-containing introns. The analysis showed a significant conservation of synteny (order of repeats) within intergenic and intronic regions around many dispersed ARs between human and other primates despite periods of divergent evolution of up to 45–60 Myr and the high abundance of different repetitive families (fig. 4 and supplementary fig. 5, Supplementary Material online). The regions proximal to most of dispersed ARs are not prone to the rearrangements and the observed stable transmission of ARs among species is in accordance with the preserved synteny of their neighboring repeats.

Mechanism of ARs Dispersion within Euchromatin

Analysis of junctions on both sides of single dispersed euchromatic alpha repeats reveals that short segments of sequence duplication of 2–10 bp size occur at most of the insertion sites (table 1). The sequence duplication is detected irrespective of the alpha repeat insertion within different context: within other repeats, adjacent to repeats, or in the unique regions without repeats within at least 100 bp (table 1). We propose that regions of short homology of 2–10 bp between an alpha repeat and its target site facilitated insertion of ARs by homologous recombination (fig. 5). The junction sequences which are duplicated are mutually different and also reflect a difference in the segments or parts of alpha satellite monomers which are inserted at the particular site (table 1). ARs embedded within other repetitive elements are not inserted within the same position in the repeats of the same type: for example, sat_605 and sat_497 are inserted at positions 169 and 73 of Alu repeat, respectively, whereas sat_84, sat_612, sat_1147, and sat_703 are inserted at positions 3051, 2147, 5618, and 5900 of L1 repeat, respectively (table 1).
**FIG. 4.**—Organization of repetitive families in the vicinity of intergenic ARs sat_827 and sat_1122 as well as intronic sat_605, in different primate species indicated in the phylogenetic tree. ARs are shown in red and other repetitive families are marked with different colors (see legend). Sat_827 and sat_605 are embedded in Tigger DNA transposon and Alu (SINE), respectively, whereas sat_1122 is inserted within unique region.
Therefore, each AR insertion seems to be unique, characterized by a specific part of the AR monomer(s) inserted at a specific genomic region.

Because some of ARs are embedded in other repetitive elements, it is possible that these elements are responsible for the dispersion of ARs. To check if ARs and the associated repeats are transferred together to the particular genomic location, we analyzed the orthologous regions in those primates separated before the insertion of the particular AR occurred (Table 2). For the sat_605 and sat_497 embedded within the Alu sequence as well as for sat_60 inserted within ERV and sat_612 in L1, respectively, in the ancestor of Catarrhini, we analyzed the orthologous regions in the species of Platyrhini: marmoset and squirrel monkey. The analyses revealed the presence of Alu sequences at sat_605 and sat_497 orthologous loci in new world monkeys but without inserted ARs (Table 2). The sat_60 and sat_612 orthologous loci also contained ERV and L1, respectively, in new world monkeys but without ARs insertion. For sat_827 inserted into the Tigger element as well as sat_703 and sat_1147 in L1, respectively, in the ancestors of Simiformes, we examined orthologous loci in tarsier (Tarsiidae) and in mouse lemur (Lemuridae). At the sat_827 orthologous locus, only the Tigger element was present without sat_827 insertion in both species. At the locus orthologous to sat_703 in the mouse lemur, L1 element without sat_703 is present, whereas in tarsier the large region encompassing orthologous sat_703 locus is absent. The regions encompassing sat_1147 orthologous loci are also absent in both species (Table 2). For sat_58 and sat_358 embedded in ERV in the ancestor of Hominioidea, examination in the old world monkeys (rhesus, baboon) revealed an intact orthologous ERV without an AR insert. Finally, for sat_84 and sat_613 embedded into L1 and ERV in the ancestors of Hominini, examination of orthologous locus in gorilla and orangutan revealed L1 and ERV element without inserted AR. All these observations show that ARs were subsequently inserted within other repetitive elements, probably by the same mechanism used for ARs insertion in unique regions without repeats, or between different repeats.

Once ARs were inserted within other repeats, it is possible that such hybrid elements are further spread along the genome by (retro)transposition or by recombination facilitated by larger segments of homology between abundant dispersed repeats such as Alu, L1, or ERVs. Most transposons in the human genome have been inactive for the last 500 Myr (Lander et al. 2001), but some retrotransposable elements including the prevalent LINE and SINE repeats which dominate the human genome are still active (Mills et al. 2007). However, our search of the human genome assembly for elements homologous to hybrid AR-(retro)transposons gave negative results, suggesting the absence of their spreading to new loci. The same is true for ARs inserted near other repeat elements. Each of the AR insertion is flanked by a specific combination of other repeats or repeat subfamilies and is present at a single genomic locus, as revealed by BLAT search. The only exceptions are sat_1145 and sat_1123, which are highly similar 4.1 mers flanked by MamGypLTR3a and L1PA3,
and are present at two different genomic loci as a result of duplication of a long chromosomal region of ∼190 kb within chromosome 9 and not of the specific spreading of the particular AR. Using UCSC browser, we searched if segmental duplications annotated in assembled human genome intersect other dispersed ARs. The results reveal that apart from sat_1145 and sat_1123, intergenic sat_372 is associated with the duplication of ∼280 kb long segment on chromosome 14. Therefore, the results indicate that most AR insertions are unique and present at a single genomic locus, whereas no evidence of subsequent spreading of inserted ARs across the genome exists.

Considering the dispersion of clustered ARs within euchromatin, sat_614-618 clustered within intron of ANKRD30BL gene corresponds to the site of a relic, previously active centromere (Miga 2017). Examination of other clustered ARs within introns of the genes SACS and LINC01580 and of three intergenic clusters revealed that all ARs within a cluster seem to be inserted within the same period, suggesting that the cluster could result from a single insertion event. Within clusters, ARs are interspersed with other repetitive elements such as Alu in the case of SACS intronic cluster or with L1 and ERV-LTRs in intergenic sat_723-727 clusters, respectively (supplementary fig. 6, Supplementary Material online). Regions composed of different types of mutually interspersed repeats are characteristic for pericentromeric heterochromatin and it is possible that they could be transferred from heterochromatin to euchromatin as larger blocks in the same way as shorter single AR repeats, probably by a recombination event relying on short sequence homology. The existence of extrachromosomal circular DNAs (eccDNA) in human cells composed of ARs ranging in size from <2 kb to over 20 kb was previously shown (Cohen et al. 2010), revealing that tandemly arranged alpha repeats are prone to generate eccDNA. We suggest the possible involvement of eccDNA in the dispersion of ARs within euchromatin and propose that their insertions within the genome were facilitated by short sequence homology between ARs and their target sequences (fig. 5).

Discussion

Our analysis of ARs annotated in the human genome reveals that majority of them, over 80%, are organized within clusters composed of ARs located within a distance <1 kb, and such clusters are preferentially located in the pericentromeric and centromeric regions. We focused on ARs not organized within clusters but present as single repeats within euchromatin and followed their evolutionary history among primates. Continuous insertion of ARs from the ancestors of Simiformes (45–60 Ma) up to the lineage of Homo is detected. In the available genomes of prosimians, tarsier (Tarsius syrichta), mouse lemur (Microcebus murinus), and bushbaby (Otolemur garnetti), we did not find alpha repeats within regions orthologous to those containing dispersed human ARs, suggesting that spreading preferentially started in the ancestors of simians. Alternatively, chromosomal rearrangements may have resulted in deletion of such regions. It is important to mention that the quality of assembly of all primate genomes is not the same and that some of them could contain the unassembled stretches of ARs which can affect our analysis. Long-read sequencing technology recently used for great ape genome assembly (Kronenberg et al. 2018) is expected to improve future studies of satellite DNA evolution.

Based on sequence features, alpha satellite monomers are classified into distinct groups and families (Shepelev et al. 2015). According to such classification, many dispersed alpha repeats belong to evolutionary old alpha satellite families which constitute pericentromeric regions, indicating dispersion of repeats from heterochromatin to euchromatin. The high sequence divergence of dispersed ARs also suggests their origin from the (per)centromeric alpha repeats which form distinct subfamilies characterized by specific, mutually divergent monomers (Willard 1985; Vissel and Choo 1992). Many extant human single dispersed ARs are up to the monomer size, whereas longer, multimeric ARs often exhibit CNV between primate species which can be explained by intrastrand recombination between homologous sequence motifs in the monomers. The recombination process seems to occur randomly and independently in different lineages during different evolutionary periods. Some single ARs composed of multimers such as sat_358 (2.7 mer), sat_685 (1.9 mer), and sat_621 (1.8 mer) have remained stable since their insertion in the ancestor of Hominoidea, Simiformes, and Catarhini, respectively. However, it is possible that these ARs show CNV in some other primate species which were not examined in our study due to the lack of genome assembly data. Excluding CNV, single ARs inserted within euchromatin are stably transmitted to descendant species showing gradual sequence evolution which generally follows species evolution. In addition, the organization of repeats in the regions proximal to dispersed alpha repeats seems to be conserved among simian primates. When compared with X chromosome euchromatic satellites in Drosophila species which exhibit a high rate of rearrangements and reorganization resulting in change of abundance, location, and composition (Sproul et al. 2020), euchromatic ARs in primate species show greater stability and lower evolutionary dynamics.

Single repeats of a major alpha satellite DNA dispersed within euchromatin are often found adjacent to other abundant repetitive elements such as Alu, L1, or ERV or are embedded within them. Some DNA transposons such as those belonging to the Helitron superfamily have a propensity to capture and mobilize flanking DNA sequences (Thomas et al. 2014) and based on such characteristics it was proposed that satellite arrays flanked by Helitron transposons can be spread throughout the genome by the process of transposition (Brajković et al. 2012; Satović et al. 2016). In this study, we did not find evidence for (retro)transposable elements to...
be specifically associated with ARs. In addition, there is no indication that (retro)transposons play a role in spreading of adjacent or embedded alpha repeats throughout the euchromatin, either by recombination with other members of the same family or by retrotransposition. We also show that segmental duplications within human genome can be associated with dispersion of only a few ARs, whereas ANKRD30BL intronic alpha repeat array corresponds to the previously mapped relic chimpanzee centromere (Miga 2017). The most probable mechanism of alpha repeats spreading that we propose is based on extrachromosomal circles of alpha satellite DNA which can be created by intrachromatid recombination of alpha repeats within heterochromatin (Feliciello et al. 2006; Feliciello, Akrap, Brajković, et al. 2015). The presence of short duplications at most AR insertion sites supports this model of satellite repeats proliferation. Extrachromosomal circular satellite DNA are common across diverse eukaryotic organisms including insects, plants, and mammals (Cohen et al. 2006; Navratilova et al. 2008; Cohen and Segal 2009; Paulsen et al. 2018, Sproul et al. 2020), including human cell lines (Cohen et al. 2010). Extrachromosomal satellite DNA circles are proposed to be amplified by rolling circle replication and reintegrated within the genome by a random process of site-specific recombination which occurs between short sequence motifs within circularized satellite repeats and homologous motifs at different chromosomal sites, either within euchromatin or heterochromatin.

In conclusion, our research reveals spreading of alpha repeats in the human genome euchromatin which occurred gradually throughout evolutionary history of primates and discloses the most probable mechanism of alpha repeats proliferation. Considering the influence of dispersed satellite repeats on local chromatin structure and gene expression in different organisms (Feliciello, Akrap, Ugarković 2015; Joshi and Meller 2017; Halbach et al. 2020; Feliciello et al. 2020), studies of their evolutionary dynamics and mechanisms of proliferation within euchromatin could contribute to the explanation of their potential role in the evolution of gene regulatory networks.

**Supplementary Material**

Supplementary data are available at Genome Biology and Evolution online.

**Acknowledgments**

This work was supported by the Croatian Science Foundation (grant IP-2019-04-6915), the Slovenian-Croatian Bilateral Cooperation, the Adris Foundation, the Italian Ministry of Education, University and Research (MIUR) and fund for Investments on Basic Research (FIRB) and the International Staff Mobility Program of University of Naples Federico II.

**Data Availability**

The data underlying this article are available in the article and in its Supplementary Material online.

**Literature Cited**

Alexandrov I, Kazakov A, Tumeneva I, Shepelev V, Yurov Y. 2001. Alpha-satellite DNA of primates: old and new families. Chromosoma 110(4):253–266.

Anisimova M, Gascuel O. 2006. Approximate likelihood ratio test for branches: a fast, accurate and powerful alternative. Syst Biol. 55(4):539–552.

Bailey TL, Johnson J, Grant CE, Noble WS. 2015. The MEME suite. Nucleic Acids Res. 43(W1):W39–W49.

Brajković J, Feliciello I, Bravo-Madarić B, Ugarković D. 2012. Satellite DNA-like elements associated with genes within euchromatin of the beetle Tribolium castaneum. G3 (Bethesda) 2:931–941.

Brajković J, et al. 2018. Dispersion profiles and gene associations of repetitive DNAs in the euchromatin of the beetle Tribolium castaneum. G3 (Bethesda) 8: 875–886.

Bulut-Karslioglu A, et al. 2012. A transcription factor-based mechanism for mouse heterochromatin formation. Nat Struct Mol Biol. 19(10):1023–1030.

Cachex L, Ponger L, Gerbault-Seureau M, Richard FA, Escudé C. 2016. Diversity and distribution of alpha satellite DNA in the genome of an Old World monkey: Cercopithecus solatus. BMC Genomics. 17(1):916.

Cohen S, Agmon N, Sobol O, Segal D. 2010. Extrachromosomal circles of satellite repeats and SS ribosomal DNA in human cells. Mob DNA 1(1):11.

Cohen S, Segal D. 2009. Extrachromosomal circular DNA in eukaryotes: possible involvement in the plasticity of tandem repeats. Cytogenet Genome Res. 124(3–4):327–338.

Cohen Z, Bacharach E, Lavi S. 2006. Mouse major satellite DNA is prone to eccDNA formation via DNA ligase IV-dependent pathway. Oncogene 25(33):4515–4524.

Eden E, Navon R, Steinfeld I, Lipson D, Yakhini Z. 2009. GOrilla: a tool for discovery and visualization of enriched GO terms in ranked gene lists. BMC Bioinf. 10(1):48.

Feliciello I, Akrap J, Brajković J, Zlatar I, Ugarković D. 2015. Satellite DNA as a driver of population divergence in the red flour beetle Tribolium castaneum. Genome Biol Evol. 7(1):228–239.

Feliciello I, Akrap I, Ugarković D. 2015. Satellite DNA modulates gene expression in the beetle Tribolium castaneum after heat stress. PLoS Genet. 11(8):e1005466.

Feliciello I, Picariello O, Chiniá G. 2006. Intra-specific variability and unusual organization of the repetitive units in a satellite DNA from Rana dalmatina: molecular evidence of a new mechanism of DNA repair acting on satellite DNA. Gene. 383:81–92.

Feliciello I, Sermek A, Pozer Ž, Matulić M, Ugarković D. 2020. Heat stress affects H3K9me3 level at human alpha satellite DNA repeats. Genes 11(6):663.

Finstermeier K, et al. 2013. A mitogenomic phylogeny of living primates. PLoS One 8(7):e69504.

Guindon S, et al. 2010. New algorithms and methods to estimate maximum-likelihood phylogenies: assessing the performance of PhyML 3.0. Syst Biol. 59(3):307–321.

Guy L, Kultima JR, Andersson SGE. 2010. genoPlotR: comparative gene and genome visualization in R. Bioinformatics 26(18):2334–2335.

Halbach R, et al. 2020. A satellite repeat-derived piRNA controls embryonic development of Aedes. Nature 580(7802):274–277.

Hinrichs AS, et al. 2006. The UCSC Genome Browser Database: update 2006. Nucleic Acids Res. 34(90001):D590–D598.
Joshi SS, Meller VH. 2017. Satellite repeats identify X chromosome for dosage compensation in Drosophila melanogaster males. Curr Biol. 27(10):1393–1402.

Katoh K, Rozewicki J, Yamada KD. 2019. MAFFT online service: multiple sequence alignment, interactive sequence choice and visualization. Briefings Bioinf. 20(4):1160–1166.

Kent WJ, et al. 2002. The human genome browser at UCSC. Genome Res. 12(6):996–1006.

Kronenberg ZN, et al. 2018. High-resolution comparative analysis of great ape genomes. Science 360(6393):eaar6343.

Kuhn GC, Küttler H, Moreira-Filho O, Heslop-Harrison JS. 2012. The 1.688 repetitive DNA of Drosophila: concerted evolution at different genomic scales and association with genes. Mol Biol Evol. 29(1):7–11.

Kumar S, Stecher G, Tamura K. 2016. MEGA7: molecular evolutionary genetics analysis version 7.0 for bigger datasets. Mol Biol Evol. 33(7):1870–1874.

Lander ES, et al. 2001. Initial sequencing and analysis of the human genome. Nature 409(6822):860–921.

Lee C, Wevrick R, Fisher RB, Ferguson-Smith MA, Lin CC. 1997. Human centromeric DNAs. Hum Genet. 100(3–4):291–304.

Lee YCG, et al. 2020. Pericentromeric heterochromatin is hierarchically organized and spatially contacts H3K9me2 islands in euchromatin. PLoS Genet. 16(3):e1008673.

Lefort V, Longueville JE, Gascuel O. 2017. SMS: smart model selection in PhyML. Mol Biol Evol. 34(9):2422–2424.

McNulty SM, Sullivan BA. 2018. Alpha satellite DNA biology: finding function in the recesses of the genome. Chromosome Res. 26(3):115–138.

Menon DU, Coarfa C, Xiao W, Gunaratne PH, Meller VH. 2014. siRNAs from an X-linked satellite repeat promote X-chromosome recognition in Drosophila melanogaster. Proc Natl Acad Sci U S A. 111(46):16460–16465.

Mi H, Muruganujan A, Ebert D, Huang X, Thomas PD. 2019. PANTHER version 14: more genomes, a new PANTHER GO-slim and improvements in enrichment analysis tools. Nucleic Acids Res. 47(D1):D419–D426.

Miga KH. 2017. Chromosome-specific centromere sequences provide an estimate of the ancestral chromosome 2 fusion event in hominin genomes. J Hered. 108(1):45–52.

Mills RE, Bennett EA, Iskow RC, Devine SE. 2007. Which transposable elements are active in the human genome? Trends Genet. 23(4):183–191.

Navratilova A, Koblikova A, Macas J. 2008. Survey of extrachromosomal circular DNA derived from plant satellite repeats. BMC Plant Biol. 8(1):90.

Paulsen T, Kumar P, Koseoglu MM, Dutta A. 2018. Discoveries of extrachromosomal circles of DNA in normal and tumor cells. Trends Genet. 34(4):270–278.

Pavlek M, Gelfand Y, Plohl M, Melstrovic N. 2015. Genome-wide analysis of tandem repeats in Tribolium castaneum genome reveals abundant and highly dynamic tandem repeat families with satellite DNA features in euchromatic chromosomal arms. DNA Res. 22(6):387–401.

Pita S, et al. 2017. Comparative repeatome analysis on Triatoma infestans Andean and non-Andean lineages, main vector of Chagas disease. PLoS One 12(7):e0181635.

Pozi I, et al. 2014. Primate phylogenetic relationships and divergence dates inferred from complete mitochondrial genomes. Mol Phylogenetics Evol. 75:165–183.

Quinlan AR, Hall IM. 2010. BEDTools: a flexible suite of utilities for comparing genomic features. Bioinformatics 26(8):841–842.

Rudd MK, Willard HF. 2004. Analysis of the centromeric regions of the human genome assembly. Trends Genet. 20(11):529–533.

Ruz-Ruano FJ, López-León MD, Cabrero J, Camacho JP. 2016. High-throughput analysis of the satellitome illuminates satellite DNA evolution. Sci Rep. 6(1):28333.

Šatović E, Vojvoda Zeljko T, Luchetti A, Mantovani B, Plohl M. 2016. Adjacent sequences disclose potential for intra-genomic dispersal of satellite DNA repeats and suggest a complex network with transposable elements. BMC Genomics. 17(1):997.

Schueler MG, Sullivan BA. 2006. Structural and functional dynamics of human centromeric chromatin. Annu Rev Genomics Hum Genet. 7(1):301–313.

Schueler MG, et al. 2005. Progressive proximal expansion of the primate X chromosome centromere. Proc Natl Acad Sci U S A. 102(30):10563–10568.

Seibt KM, Schmidt T, Heitkam T. 2018. FlexiDot: highly customizable, ambiguity-aware dotplots for visual sequence analyses. Bioinformatics 34(20):3575–3577.

Shepelev VA, et al. 2015. Annotation of suprachromosomal families reveals uncommon types of alpha satellite organization in pericentromeric regions of hg38 human genome assembly. Genomics Data 5:139–146.

Sproul JS, et al. 2020. Dynamic evolution of euchromatic satellites on the X chromosome in Drosophila melanogaster and the simulans clade. Mol Biol Evol 37(8):2241–2256.

Thomas J, Phillips CD, Baker RJ, Pritcham EJ. 2014. Rolling-circle transposons catalyze genomic innovation in a mammalian lineage. Genome Biol. 6(10):2595–2610.

Vissel B, Choo KHA. 1992. Evolutionary relationships of multiple alpha satellite subfamilies in the centromeres of human chromosomes 13, 14, and 21. J Mol Evol. 35(2):137–146.

Vlahovic I, Gluncic M, Rosandic M, Ugarkovic D, Paar V. 2017. Regular higher order repeat structures in beetle Tribolium castaneum genome. Genome Biol Evol. 9(10):2668–2680.

Willard HF. 1985. Chromosome-specific organization of human alpha satellite DNA. Am J Hum Genet. 37(3):524–532.

Willard HF. 1991. Evolution of alpha satellite. Curr Opin Genet Dev. 1(4):509–514.

Associate editor: Ellen Pritham