

# A Comparative Approach Shows Differences in Patterns of Numt Insertion During Hominoid Evolution

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**Abstract** Nuclear integrations of mitochondrial DNA (numts) are widespread among eukaryotes, although their prevalence differs greatly among taxa. Most knowledge of numt evolution comes from analyses of whole-genome sequences of single species or, more recently, from genomic comparisons across vast phylogenetic distances. Here we employ a comparative approach using human and chimpanzee genome sequence data to infer differences in the patterns and processes underlying numt integrations. We identified 66 numts that have integrated into the chimpanzee nuclear genome since the human–chimp divergence, which is significantly greater than the 37 numts observed in humans. By comparing these closely related species, we accurately reconstructed the preintegration target site sequence and deduced nucleotide changes associated with numt integration. From >100 species-specific numts, we quantified the frequency of small insertions, deletions, duplications, and instances of microhomology. Most human

and chimpanzee numt integrations were accompanied by microhomology and short indels of the kind typically observed in the nonhomologous end-joining pathway of DNA double-strand break repair. Human-specific numts have integrated into regions with a significant deficit of transposable elements; however, the same was not seen in chimpanzees. From a separate data set, we also found evidence for an apparent increase in the rate of numt insertions in the last common ancestor of humans and the great apes using a polymerase chain reaction–based screen. Last, phylogenetic analyses indicate that mitochondrial-numt alignments must be at least 500 bp, and preferably >1 kb in length, to accurately reconstruct hominoid phylogeny and recover the correct point of numt insertion.

**Keywords** Numt · Hominid · Hominoid · Evolution · Mitochondria · Phylogenetic · Ape

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## Introduction

Nuclear integrations of mitochondrial DNA (numts) are fragments of the mitochondrial genome that have incorporated into germline nuclear DNA (nDNA), and they have been reported in animals, plants, and fungi (Thorsness and Fox 1990; Zischler et al. 1995a; Blanchard and Schmidt 1996; Zhang and Hewitt 1996; Bensasson et al. 2001; Leister 2005). They are usually referred to as “pseudogenes.” However, there are some instances in which they may have been exonized in a few species (Noutsos et al. 2007), including a human-specific numt that inserted into a 3'-UTR (Ricchetti et al. 2004). Although they are non-functional, numts offer a model for early eukaryotic evolution in which hundreds of genes are believed to have retained function after their migration from the

protomitochondrial endosymbiont to the nuclear genome (Margulis 1970; Andersson et al. 2003; Lang et al. 1999; Blanchard and Lynch 2000). Numts are commonly believed to “fossilize” after their integration into the nuclear genome, i.e., the nuclear translocated mitochondrial copy is more likely to resemble the ancestral mitochondrial haplotype at the time of its insertion than its modern mitochondrial counterpart because there is a much lower mutation rate in the nuclear genome (Zischler et al. 1995a). As such, numts offer interesting opportunities to study mitochondrial DNA evolution. Numts can also be problematic if mistaken for authentic mitochondrial DNA (mtDNA), potentially confounding interpretations in wildlife genetics, forensics, ancient DNA, or medical studies (van der Kuyl et al. 1995; Zischler et al. 1995b; Wallace et al. 1997; Bensasson et al. 2001; Jensen-Seaman et al. 2004; Anthony et al. 2007).

Although they are taxonomically widespread, the prevalence of numts varies tremendously among species, suggesting that the processes of numt integration may change with time and/or differ between taxa. Analyses of complete genome sequences have shown numts to be present in all mammals examined thus far, with large numbers reported in humans, chimpanzees, and cats but fewer found in mice and rats (Mourier et al. 2001; Tourmen et al. 2002; Woschnik and Moraes 2002; Richly and Leister 2004; Antunes et al. 2007). Honeybees display an unusually large number of numts (Behura 2007; Pamilo et al. 2007), whereas there are few to none in *Drosophila* and *Anopheles* (Richly and Leister 2004). There has been some controversy over the presence of numts in any fish species; however, for now it appears that at least the Fugu genome is devoid of them (Antunes and Ramos 2005; Venkatesh et al. 2006).

Despite this increasing awareness of the prevalence of numts in many taxa, comparisons of numt distributions among relatively closely related species are rare (however, see Krampis et al. 2006; Hazkani-Covo and Covo 2008). One approach to identifying differences in insertion rates between species, especially in locating temporal fluctuations in numt insertion rates, has been to infer the point of insertion using a phylogeny-based approach. In primates, this has led to the suggestion that a burst of insertions occurred near the time of the divergence between Old World and New World monkeys (Bensasson et al. 2003; Hazkani-Covo et al. 2003; Gherman et al. 2007). Another study suggested that humans may be experiencing a recent increase in numt integration (Ricchetti et al. 2004), although that conclusion was based on the assumption that neutral alleles in the human populations are expected to have a coalescence time within the last 100,000 years. Within great apes, it has been suggested that the frequency of numts is increased in gorillas, although these

observations are limited to the mitochondrial D-loop (Jensen-Seaman et al. 2004; Thalmann et al. 2004; Anthony et al. 2007). Whether or not this claim can be substantiated awaits a more detailed comparison of insertion rates between taxa and across the entire mitochondrial genome.

Numts do not appear to show a preference for specific target sequences, as has been found for retrotransposable elements (Cost et al. 2002), although whether they integrate truly randomly remains an open question. Large-scale analyses of human numts, made possible by the complete human genome sequence, suggested that the immediate flanking regions (approximately 15 bp) contained fewer transposable elements (TEs) than expected by chance, whereas slightly more distal regions (15 to 150 bp) contained more TEs than expected (Mishmar et al. 2004). The opposite conclusion was reached by Gherman et al. (2007), who found a decrease in the TE content of the first 150-bp sequence flanking human numts, with this decreased TE content extending across at least 1 kb away from the numt. With respect to any sequence composition preference for human numt insertions, most studies have not found a strong pattern. Exceptions include the suggestion that numts are preferentially found in regions with a different GC-content than that of the surrounding chromosomal G-band (Mishmar et al. 2004) and another being the recent observation that human numts prefer low GC-content isochores (Lascaro et al. 2008). Finally, it has been reported that older numts tend to be found outside genic regions, whereas more recent human-specific numts preferentially insert within introns (Ricchetti et al. 2004).

Numts are believed to integrate predominantly, or perhaps exclusively, by way of DNA-mediated transfer during the use of nonhomologous end-joining (NHEJ) repair of double-stranded breaks (Blanchard and Schmidt 1996; Ricchetti et al. 1999; Hazkani-Covo and Covo 2008). Compared with repair by way of homologous recombination, classical NHEJ does not seek out truly homologous sequences as templates for repair but typically uses short (1–4 bp) stretches of sequence identity, or “microhomology”, to facilitate end-joining. NHEJ is inherently error prone, commonly involving short insertions and deletions at the repair site, with some of these insertions deriving from the fill-in of staggered double-strand breaks (Roth et al. 1985). The inference of microhomology, insertions, and deletions requires knowledge of both the target sequence before numt integration as well as the postintegration sequence. For this reason, only a few naturally occurring numt junctions had been examined (Zischler et al. 1995a; Ricchetti et al. 2004) before the recent work by Hazkani-Covo and Covo (2008), who demonstrated the preponderance of NHEJ repair with microhomology, although with a decreased frequency of deletions relative to experimental systems, indicating that numts may help

decrease the deleterious effects of deletions during double-stranded break repair.

Here we employ a comparative genomic approach to address several questions related to numt integrations in primates. A complete genome assembly exists for humans and chimpanzees, whose nuclear DNA differs by approximately 1% (CSAC 2005), making these species ideal for the comparative study of numts in closely related taxa. This study is divided into four main objectives. Firstly, we use closely related species as proxies for the preintegration site to accurately determine the extent of the numt insertion and to quantitatively infer the presence of microhomology and indels in >100 species-specific numts. Secondly, we use these comparative data to test for an overabundance or deficit of TEs near numt insertions. Third, we use a polymerase chain reaction (PCR)-based assay to determine the likely time of insertion of a range of numts across the entire ape phylogeny to test whether numt insertions are uniform through time. Finally, we assess how well these insertion times can be recovered using computationally based phylogenetic methods.

## Materials and Methods

### Identification of Human and Chimpanzee Numts

The human mitochondrial genome sequence (NC\_001807; Ingman et al. 2000) was aligned to the human nuclear genome (March 2006 assembly; National Center for Biotechnology Information build 36.1) using the *blastn* program of locally installed *BLAST* (Altschul et al. 1990), with an Expect (e) value of 10. Similarly, the chimpanzee mtDNA sequence (NC\_001643; Horai et al. 1995) was *BLAST*ed to the chimpanzee genome (March 2006 assembly; build 2, v. 1). Hits to “chromosome unknown” or “random” chromosome contigs were ignored. Because *BLAST* tends to fragment contiguous matches interrupted by more diverged sequences (Jareborg et al. 1999), hits within 1 kb were automatically grouped into a single hit. No attempt was made to exhaustively identify all, particularly highly divergent, numts. For each hit, the candidate human numt, the corresponding portion of the human mtDNA, and 500 bp of the left and right flanking human nuclear sequence were compared with the chimpanzee genome using *BLAT* (Kent 2002). This was followed by visual inspection to determine whether the numt was human specific or present in both species. The same process was carried out for putative chimpanzee numts using *BLAT* to compare them with the human genome to identify chimpanzee-specific numts (Supplementary Fig. 1). Comparing the putative numt region with that of a closely related species to confirm that an insertion indeed took place

permits use of a low stringency *BLAST* search, increasing the chance of identifying short and/or more diverged numts while eliminating false-positive *BLAST* hits. Differences in the number of species-specific numts between humans and chimpanzees were tested using a G-test (Sokal and Rolf 1995). For species-specific numts, the homologous nuclear genomic regions, including the numt and 50 bp on either side from both species, along with the homologous mitochondrial region plus 50 bp on either side, were aligned locally with *MUSCLE* (Edgar 2004) and examined by eye to determine the extent of microhomology and to identify small insertions, deletions, and duplications that apparently occurred concomitantly with the integration of mtDNA.

### Transposable Element Content of Preintegration Sites

We used the chimpanzee genomic sequence as the proxy for the preintegration site for human-specific numts. We specified the point of numt insertion in the homologous chimpanzee sequence as (1) the base pair at which the two homologous human flanking sequences meet in the absence of the numt or (2) the midpoint of the short intervening gap if the human homologous flanking sequences did not quite meet when aligned to the chimpanzee genome with *BLAT* (see Supplementary Fig. 1). The same criteria were applied to the delimitation of chimpanzee-specific numts using *BLAT* to determine their presence or absence in the human genome (Supplementary Fig. 1). From this point, 10 nonoverlapping windows of 100 bp at increasing distances (0–1 kb) from either side of each numt insertion point were extracted from the preintegration cross-species proxy sequence and analyzed for repetitive element content using locally installed *RepeatMasker* with the *-e* WU-*BLAST* option (Smit et al. <http://www.repeatmasker.org>). To test for significance, we generated a distribution of similar data taken from throughout the genome. For this, we randomly selected a number of genomic locations equivalent to the number of species-specific numts (i.e., 37 random locations for human and 66 random locations for chimpanzee) and again extracted 100-bp windows for analysis with *RepeatMasker* as described previously. This random selection was repeated 10,000 times using a custom *perl* script to create a distribution with which to compare our observed values.

### Determining the Point of Insertion of Hominoid Numts Using Cross-Species PCR

To empirically determine the point of numt insertions more broadly throughout hominoid evolution, we *BLAT*ed all putative human numts identified in the original mtDNA-to-nDNA *BLAST* search and their flanking sequences to the rhesus macaque genome sequence assembly (January 2006 assembly) to identify those numts not present in the

macaque genome (i.e., numts that integrated since the cercopithecoïd–hominoid split). We also excluded numts shown to be human-specific because we already determined their time of insertion computationally (see previous text). From the human–macaque sequence alignment, PCR primers were then designed from conserved regions flanking each numt. We also excluded very large numts or those flanked by too many repetitive elements to be able to design reliable primers. PCR amplification was attempted for 50 of these loci using genomic DNA samples from human (*Homo sapiens*), chimpanzee (*Pan troglodytes*), gorilla (*Gorilla gorilla*), orangutan (*Pongo pygmaeus*), gibbon (*Hylobates lar*), and macaque (*Macaca mulatta*). PCR primer sequences and reaction conditions are available on request. Presence or absence of each numt for each species was scored based on the expected size of the amplification product with and without the numt. Of the 34 numts successfully placed (of 50 attempted) in this manner, 8 PCR products were sequenced in whichever of the above species is most closely related to human, but found to lack that particular numt, to confirm its absence. The observed phylogenetic distribution of nuclear integrations was compared with an expected distribution assuming equal rates of numt insertion across all branches leading to humans, and evaluated using  $\chi^2$  test. For this we used hominoid divergence dates from Raaum et al. (2005) and Stauffer et al. (2001).

### Phylogenetic Analysis

We assessed the ability of tree-based methods to correctly infer the point of insertion of 40 numts in the hominoid phylogeny by conducting phylogenetic analysis on each of the 34 loci that had been placed empirically by cross-species PCR (see previous text), along with 6 more determined to be human specific through comparison to the chimp genome (see previous text). To do this, mtDNA sequences from human, chimp, gorilla, orangutan, gibbon, and macaque were first aligned without their corresponding nuclear copy using *Clustal X* (Larkin et al. 2007). Maximum parsimony (MP) and maximum likelihood (ML) phylogenetic analysis were then carried out to identify those mitochondrial data sets (without the numt) with sufficient signal to recover the accepted primate phylogeny (Goodman et al. 1998). MP and ML analyses were carried out in PAUP\* v4.4 (Swofford 2002) using starting trees obtained from the stepwise addition and neighbor-joining method options, respectively. Heuristic searches were conducted with the tree-bisection-reconnection method, and branch support was obtained from 100 bootstrap replicates of the data. Starting evolutionary parameters for ML analyses were obtained from the Akaike Information Criterion option in *ModelTest* (Posada and Crandall 1998). The 8 mtDNA data sets that correctly recovered the recognized

hominoid phylogeny were then also subjected to Bayesian phylogenetic analysis to check for consistency across methods. Bayesian analysis was carried out using *BEAUTi/BEAST* v1.4.6 (Drummond and Rambaut 2007) using tree priors based on the accepted tree topology and approximate divergence times between taxa using an uncorrelated, log-normal clock to allow for among lineage rate variation and with starting evolutionary parameters from *ModelTest*. A Monte Carlo Markov Chain (MCMC) of 50 million steps in length was run with a sampling interval every 1000 steps. The appropriate burn-in period (10%) was determined from visual inspection of output in *Tracer* v1.4 (Drummond and Rambaut 2007). Using these search options, all parameter values could be estimated from effective sample sizes  $\geq 100$ . For those 8 cases for which mtDNA sequences were able to recover the accepted primate phylogeny, the point of numt insertion in the phylogeny was then tested in a phylogenetic framework by imposing a backbone constraint on the underlying primate phylogeny and using ML to find the most likely placement of the numt insertion.

## Results

### Chimpanzees Have More Recent Numts Than Humans

*BLAST* searches of the human genome assembly using the human mitochondrial genome sequence as query and grouping hits <1 kb apart yielded a total of 519 putative loci. Similarly, *BLAST* searches using the chimpanzee mtDNA sequence against the chimpanzee genome assembly yielded 579 hits. We compared each putative numt, left and right flanking regions, and the homologous mtDNA domain from the human to the chimpanzee nuclear genome, or vice versa, using *BLAT* followed by manual inspection of the *BLAT* visualization to score each numt as either species-specific or shared between human and chimpanzee (Supplementary Fig. 1). This approach identified 37 human-specific numts and 66 chimpanzee-specific numts (Supplementary Tables 1, 2). The greater number of chimpanzee numts is significant ( $G_{\text{adj}} = 8.237$ ;  $p < 0.005$ ,  $df = 1$ ). The mean length of the chimpanzee-specific numts is larger than that of the human numts (chimp mean = 554 bp and human mean = 321 bp), although this difference is not significant ( $p = 0.37$ , Student's *t* test with Welch's correction for unequal variances).

### Use of Outgroup to Define Junctions and Infer Preintegration State

The use of *BLAST* identifies numts by sequence similarity to the mitochondrial genome, but it does not necessarily accurately delineate the exact boundaries of the numt. Here

```

H:2:149      CTTTAGTCCTTACCTCTAAATCATCGTGGTGATT...GATGTGAGCCCGTCTAAACAGTTTCCACCTGTGTC
P:2b:153    CTTTAGTCCTTACCTC-----...-----AGGTTTCCACCTGTGTC
H:M:513     TGCTTGATGCTTGTCCTTTTGTATCGTGGTGATT...GATGTGAGCCCGTCTAAACATTTTTCAGTGTATTGCT

H:3:68.8    ATGGCTCTTGGGCTAGAGCCCAAGCATGTTCGTTACATGGTCCATCATAGTCCTCTGGTAAATAGGTCTTTT
P:3_70.5    ATGGCTCTTGGGCTAG-----TTCTTAGTCCTCTGGTAAATAGGTCTTTT
H:M:12513   TCTCCATAATATTCATCCCTGTAGCATGTTCGTTACATGGTCCATCATAGAATTCTCACTGTGATATATAAA

H:13:55.4   TGGGTCTGAGAGTAAAGGGAATAGTAGGCCCTCCTAGG...GAGTAATAGAAATGCGGTAATACAAAGCAGAAT
P:13_55.9   TGGGTCTGAGAGTAAA-----...-----GGTAATACAAAGCAGAAT
H:M:5009    GGGCAAAAAGCCGGTTAGCGGGGGCAGCCCTCCTAGG...GAGTAGTAGGAATGCGGTAGTAGTAGGATAAT

H:14:32.0   TCCCTGCAAGGGATAGGTGTTGGTATAGA...CAGTCCTTAGCTGCAAATGAGTCCTTAGCAAGGCACTCATT
P:14_31.5   TCCCTGCAAGGGA-----...-----CATGAACTCATT
H:M:5484    CGAAAAATCAGAAATAGGTGTTGGTATAGA...CAGTCCTTAGCTGTTGCAGAAATTAAGTATTGCAACTTACT

```

**Fig. 1** Examples of the use of the chimpanzee genomic sequence to accurately define the boundaries of the inserted numt. *Shaded nucleotides* represent BLAST-defined homology between human mtDNA and the human numt region. *Boxed nucleotides* are the actual nucleotides inserted, as inferred from comparison with the chimpanzee sequence, which served as proxy for the preintegration target site.

we define the numt as the stretch of nucleotides that was inserted into the genome compared with the inferred preintegration site, which in many cases is slightly different from the stretch of nucleotides that share significant similarity with mtDNA in a *BLAST* search for two main reasons. First, several additional bases are commonly added during the insertion, often from small duplications and direct flanking repeats (Fig. 1). Second, short stretches of similarity are frequently found between the mtDNA and the preintegration sequence, termed “microhomology” (Fig. 1). These two features cause underestimation and overestimation, respectively, of the length of the numt. Of the 37 human-specific numts we identified, 23 possess at least some nucleotides inserted without similarity to mtDNA; 18 show deletions of nuclear DNA on insertion; and nearly all show some microhomology at one or both flanks (see Supplementary Fig. 2 for alignments of all human-specific numts). Because nucleotides appear to be commonly deleted from the nuclear target site during the process of integration, as inferred from human–chimpanzee alignments (Fig. 1), the flanking sequences, even when properly defined, do not offer the best estimation of the preintegration sequence (i.e., “target site”). We were unable to accurately align the human and chimpanzee nuclear sequences at the point of insertion for two chimpanzee-specific numts caused by complex rearrangements, leaving 64 chimpanzee numts for the integration site analyses below (see Supplementary Fig. 3 for alignments of all chimpanzee-specific numts).

#### Microhomology and Indels at Integration Sites

Comparing the chimpanzee sequence (representing the preintegration state) with the human mtDNA sequence

*Bold nucleotides* indicate insertions in human or deletions in chimpanzee. Sequences are labeled by species (*H* *Homo*, *P* *Pan*), followed by chromosome number (or “*M*” for the mitochondrial genome), followed by beginning position rounded to the nearest Mb for nuclear DNA or nearest bp for mitochondrial DNA

immediately flanking the numt insertion point shows that most (35 of 37) of the numt insertions occur in the presence of microhomology at one or both of the numt–nuclear junctions. (Fig. 2a, c; Supplementary Fig. 2). Similarly, 45 of the 64 chimpanzee-specific numts possess microhomology at one or more of the junctions (Fig. 2; Supplementary Fig. 3). The distribution of lengths of observed microhomology for both species is similar to that reported for other types of DNA integration into eukaryotic DNA, with most microhomology being limited to 1–4 bp, although one of the human-specific numts did contain a 10-bp stretch of identical nucleotides (Fig. 2d). These estimates are conservative in that only perfect uninterrupted matches were counted. We are operationally defining microhomology as exact matches regardless of length to compare with other data sets (Fig. 2d). However, this does not necessarily imply that the matches were used in the numt integration process and may have occurred simply by chance.

The majority (23 of 37) of human-specific numts contain insertions of nucleotides that appear neither to have been derived from mtDNA nor present before insertion—again using the chimpanzee nuclear sequence as an estimate of the preintegration site. These insertions are between 1 and 37 bp (mean 7.2; median 5), examples of which are shown in Fig. 3a–f. Almost all of these insertions can be explained as being derived from 1 of 3 sources: flanking direct repeats, tandem direct repeats, and tandem inverted repeats. Considering only insertions of at least 4 bp, 8 of 37 human-specific numts contain flanking direct repeats of 4–14 bp, which are always found precisely at the junction between the numt and flanking DNA (Fig. 3a, b). Four cases of tandem direct repeats of 5–22 bp were found, only 1 of which was truly tandem in that the duplicated nucleotides immediately follow the source nucleotides (Fig. 3c),



whereas the other 3 included between 2 and 9 nucleotides spacing the duplication (e.g., Fig. 3d). Interestingly, and most certainly anecdotally, the 6 nucleotides at the left flank of 1 particular numt insertion that includes a tandem duplication are a perfect complement to the mitochondrial sequence (Fig. 3c, underlined text). Finally, 5 of the 37 human-specific numt insertions were accompanied by inverted repeat sequences of between 8 and 12 nucleotides, spaced by 1–6 bp (e.g., Fig. 3e, f). Eighteen of the 37 human-specific numts show evidence for deletions of 1–157 bp in the preintegration sequence using the chimpanzee sequence as proxy. As with humans, the majority (40 of 64) of chimpanzee-specific numts contain insertions of 1–60 bp

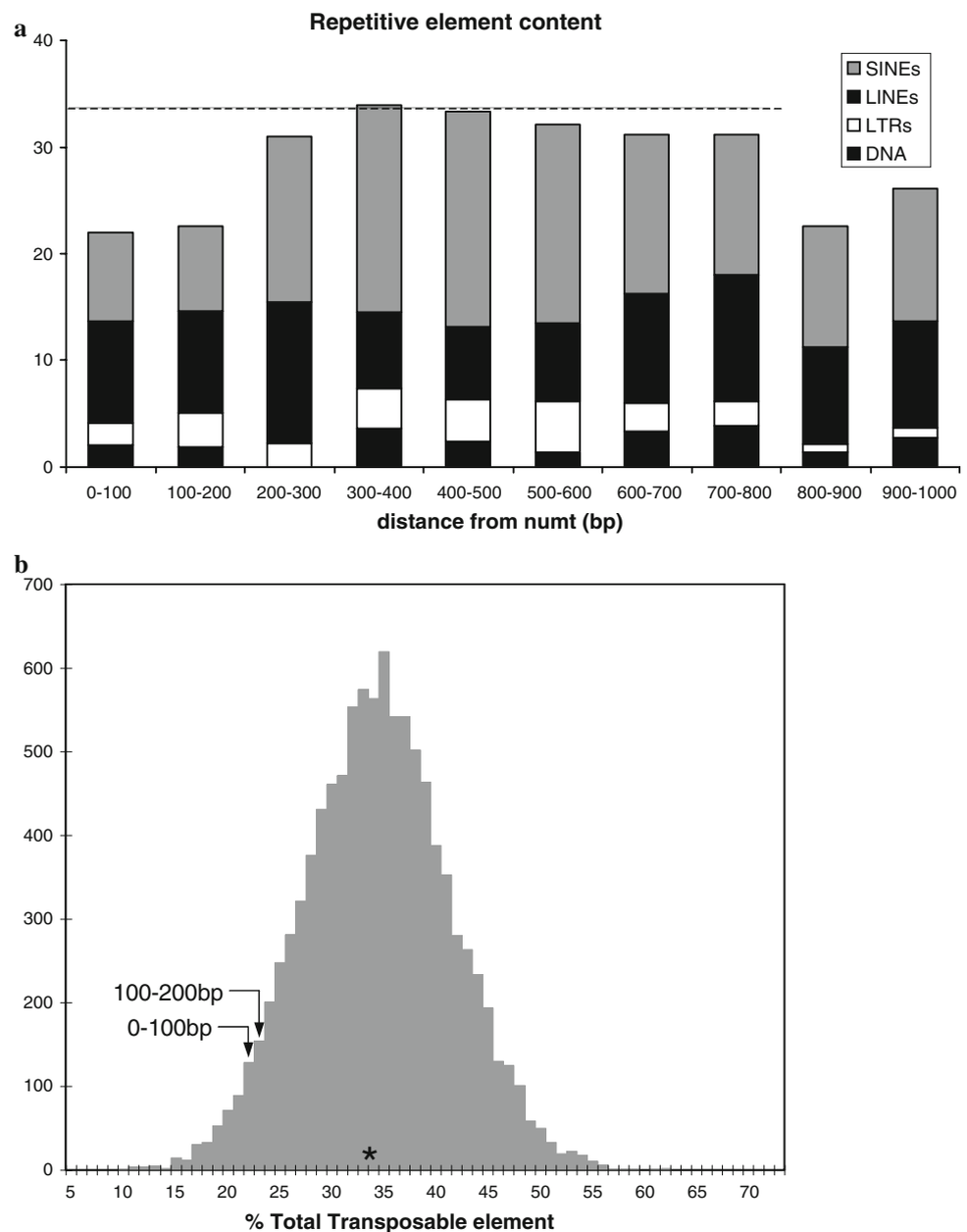
(mean 5.4; median 1.5), including 8 flanking direct repeats (4–14 bp), 9 tandem direct repeats (4–12 bp), and 4 tandem inverted repeats (7–15 bp). Alignments of all human-specific and chimpanzee-specific numts with their preintegration sequences are shown in Supplementary Figs. 2 and 3.

#### Deficit of TEs in Human Numt Flanks

Taking the 37 human-specific numts, we used the homologous insertion point in the chimpanzee genome to represent the preintegration state. There is decreased density of TEs in the first 2 100-bp windows on either side of the insertion point (Fig. 4a). This includes a decrease in all 4

**Fig. 4** **a** Transposable element content in 100-bp windows flanking human-specific numts. Each column shows the major classes of TEs estimated from 7400 bp (37 numts  $\times$  2 flanking regions  $\times$  100 bp). *Dashed line* indicates the average (33.8%) of the total transposable element content found in 10,000 randomly generated data sets (each data set consisted of 37 regions  $\times$  2 flanking regions  $\times$  100 bp). **b** Distribution of the total transposable element content of the 10,000 randomly generated data sets along with the values from the first 100 bp and the second 100 bp from the flanking regions of the 37 human-specific numts.

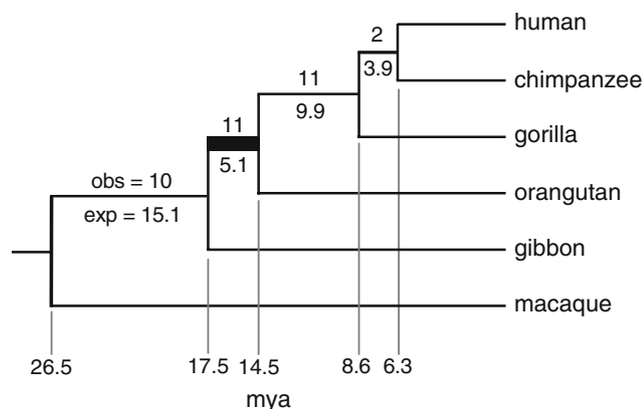
\* Average of the distribution



categories of TEs compared with the genome-wide average (1–100 bp: SINEs 8.39%, LINEs 9.49%, LTRs 2.09%, DNA-based mobile elements 2.03%; 101–200 bp: SINEs 7.97%, LINEs 9.55%, LTRs 3.25%, DNA-based mobile elements 1.84%). The total proportion of TEs in the 100-bp windows immediately flanking the numt insertions (22.00%) falls within the most extreme 5% of a distribution made from 10,000 randomly generated data sets (4.46 percentile; Fig. 4b), whereas the next 100 bp (100–200 bp away from the insertion point) nearly does (5.32 percentile). When chimpanzee-specific numts are examined in analogous fashion (using the human genome as the outgroup sequence to define the preintegration state), no significant decrease in TE content is seen at or near the point of numt insertion.

#### Determination of Numt Insertion Time

We determined the time of insertion of 34 numts using cross-species PCR. Most branches of the hominoid phylogenetic tree have approximately the same or fewer insertions than the expected number based on estimated divergence times between internal nodes, with the exception of the stem hominid (great ape and human) lineage that follows the split with gibbons (Fig. 5). This distribution of numts across the primate phylogeny differs significantly than expected ( $\chi^2 = 9.78$ ;  $p = 0.021$ ;  $df = 3$ ), driven almost entirely by the excess of numt insertions in the stem hominid, using the Bayesian estimates of divergence dates as reported by Raaum et al. (2005). Our result of a significant excess of numt insertions in the hominid ancestor is highly



**Fig. 5** Observed (above branch) and expected (below branch) distribution of hominoid numt insertions, as determined with cross-species PCR, shown on the universally accepted phylogeny. Bayesian posterior probability estimates of divergence times are given below the tree, taken from Raaum et al. (2005), and used to calculate the expected number of numts on each branch. A significant excess of numts have inserted into the common ancestor of humans and the great apes after their divergence with gibbons ( $p < 0.05$ ; indicated by thick branch)

dependent on the estimated divergence dates, which vary widely depending on methodology, data sets, and choice of fossil calibration. Indeed, the observed uneven phylogenetic distribution is not significantly different than the expected uniform distribution when using dates estimated with ML on the same mitochondrial genome data (Raaum et al. 2005) ( $\chi^2 = 6.88$ ;  $p = 0.076$ ) or using dates derived from nuclear data (Stauffer et al. 2001).

Taken together, MP and ML methods recovered the accepted hominoid phylogeny in a total of 8 of 40 data sets. Of these, MP only recovered the topology correctly in 4 cases, whereas ML recovered the topology successfully in 7 cases. Of these 8 instances, Bayesian analyses correctly recovered the same topology in 7 cases, only 1 of which was not recovered in ML. The size of the alignments for these 8 loci ranged from 149 to 2457 bp in length (mean 1055.3; median 676.5). In contrast, alignments that failed to recover the accepted hominoid topology ranged from 50 to 487 bp in size (mean 149.5; median 135.0), with a significant difference in the 2 medians of these groups (Mann–Whitney  $U$  test  $p < 0.001$ ). ML analysis with the backbone constraint imposed reliably assigned the numt to the correct position in only 5 of the 8 cases in which the underlying primate topology was recovered using one or more phylogenetic methods.

#### Discussion

##### Nonrandom Numt Insertion Through Time

It has been well established that some species contain a greater number of numts than others (Bensasson et al. 2001). From available genome sequence data, it appears that *Homo*, *Arabidopsis*, and *Apis* have substantial numbers of numts, whereas *Drosophila*, *Anopheles*, *Fugu*, and *Galus* do not (Richly and Leister 2004; Pamilo et al. 2007; Venkatesh et al. 2006; Pereira and Baker 2004). Most of the general conclusions that can be made from these comparisons, however, are across large phylogenetic distances and are seemingly explained by differences between broad taxonomic categories (e.g., fish vs. mammals, plants vs. animals, dipterans vs. hymenopterans).

Here we take advantage of the complete genome sequence of two closely related species to compare the rates of numt integration and infer preintegration target sequences. Chimpanzees have a significantly greater number of recent species-specific numts than humans. Numts can increase in frequency in two ways: greater mitochondria-to-nucleus transfer or increased intranuclear postinsertion duplications (Hazkani-Covo et al. 2003). None of the chimpanzee-specific numts appear to be postinsertion duplicates of each other; therefore, they likely reflect an increased rate of transfer from the mitochondria.

It is difficult to explain why chimpanzees should have more numts, and we offer no definitive answers here. If numts are considered slightly deleterious mutations, as TEs usually are, and with all else being equal, selection should more efficiently remove them from a larger population. However, chimpanzees have historically larger effective population sizes than humans (Jensen-Seaman et al. 2001; Yu et al. 2003; Burgess and Yang 2008). Therefore, in the absence of other factors we might expect to see more numts in humans, which is not the case. It is also conceivable that chimpanzees and humans differ slightly in their double-strand break repair mechanisms, including overall efficiency or preference for one mechanism over another, which may be related to the likelihood of mtDNA being incorporated during repair. Nonhuman primates have substantially lower rates of cancer than humans, which may be in part caused by genetic differences, including DNA repair enzymes (Puente et al. 2006). Alternatively, differences may exist between humans and chimpanzees in the cellular availability of degraded mtDNA in germline cells (see Richly and Leister 2004; Willett-Brozick et al. 2001). Chimpanzee sperm cells, and therefore potentially zygotes, may contain more mtDNA than their human counterparts because the volume of the sperm midpiece, which is packed with mitochondria, is substantially greater in chimpanzees compared with humans (Anderson and Dixon 2002).

It is important to consider the possibility that the greater number of observed numts in chimpanzees may be an artifact of methodology or data quality. We believe that all of the chimp-specific numts identified here are true numt insertions, not spurious *BLAST* matches nor deletions in humans because their identification was not only based on similarity to mtDNA but also the absence of precisely that sequence in the human genome at the homologous location (and vice versa regarding human-specific numts). Furthermore, we do not believe that we are overcounting chimpanzee numts by including 1 single numt insertion split into 2 numts by the insertion of a TE; no 2 chimp numts are within 1 Mb of each other. Concerning methodology, we note that our approach to identifying human numts did find a nearly identical set of numts as that recently described by others. Specifically, all 34 human-specific numts previously identified using a different approach (Hazkani-Covo and Graur 2007) were identified herein, along with 3 additional numts. We used a newer version of the human genome assembly, which may account for the additional 3 numts discovered. Similarly, all 27 human-specific numts described by Ricchetti et al. (2004) were found with our approach. The lower quality of the chimpanzee genome assembly might account for a greater number of identified numts, although it is not clear how. If anything, we might expect a smaller number to be found in a more fragmented, error-

prone genome assembly. Indeed, Hazkani-Covo and Covo (2008) found substantially more numts in the build 2 version of the chimpanzee genome than Hazkani-Covo and Graur (2007) found in the build 1 version using similar methods, suggesting that increasing assembly quality will only add to the number of recovered numts. We therefore believe that the greater number of observed numts in chimpanzees relative to humans reflects a true biologic difference. Nonetheless, we do heed the admonition by Venkatesh et al. (2006) that all numts from shotgun-sequenced genomes must be empirically verified and anticipate that future research will do so.

Using cross-species PCR on a subset of human numts, we observed an excess of numt insertions in the stem hominid branch leading to the human and great ape clade, with 11 of 34 hominid-specific numts inserting along this short branch. The approximate time of these insertions, i.e., the mid-Miocene, saw an impressive adaptive radiation of apes with greater species diversity than seen previously or since. However, it should be emphasized that the increases in numt insertions in the stem hominid are strongly dependent on the accuracy of our estimated divergence dates. Although recent years have seen an explosion of sequence data used to date these events with the molecular clock, the fossil record is still woefully inadequate for accurate calibration (Jensen-Seaman and Hooper-Boyd 2008). We also note that this increase could be due in part to an ascertainment bias in that some numt loci failed to amplify in all species and were excluded from analyses. We cannot envision, however, how this could have led to an increase the number of observed numt insertions at this point in the tree. If anything, we might expect a bias toward more recent events.

Several previous studies have suggested a nonuniform rate of numt insertions into the human genome, particularly in identifying a burst of insertions near the time of the split between Old World (catarrhine) and New World (platyrrhine) anthropoids, especially along the branch leading to all extant catarrhines (Hazkani-Covo et al. 2003; Bensasson et al. 2003) or even earlier (Gherman et al. 2007). It is, however, difficult to accurately determine the time of insertion of any numt with a purely computational phylogenetic approach for several reasons. First, it requires including mitochondrial and nuclear sequences in the same tree and as such requires the evolution of these sequences to be modeled with the same parameters. It is widely known that nuclear and mitochondrial sequences have different mutation rates, different transition-transversion biases, and different patterns of among-site rate heterogeneity—all critical variables in modeling sequence evolution. The difficulties with respect to accurately placing numts using phylogenetic methods were explored in depth by Bensasson et al. (2003), who demonstrated not only a

strong dependency on which model of sequence evolution was used but also a consistent decrease in the nonuniformity of numt insertions as increasingly realistic models were used. In this light, the more simplistic models used by Gherman et al. (2007) may explain, at least in part, their observation of an extremely strong temporal burst of numt insertions.

In trying to improve on the ability to phylogenetically place human numt insertions, we applied alternative approaches (MP, ML, and Bayesian) to assess which mitochondrial data had sufficient phylogenetic information to recover the accepted primate tree. Only 8 of 40 numts (20%) contained sufficient phylogenetic signal, indicating that a previous selection of data sets is necessary before attempting to infer numt insertion points. We doubt the reliability of any method to accurately place numts shorter than 500 bp, reinforcing the need to complement phylogenetic analyses with wet laboratory techniques or comparative genomic studies when candidate numt loci are  $\leq 500$  bp.

#### Nonrandom Numt Insertion in Genomic Space

Human-specific numts preferentially integrate into regions of low TE density. More than 200 bp away from the insertion point, this no longer holds true, largely because of the abundant SINEs several hundred nucleotides away. Previous studies examining the flanking regions of all human numts have presented conflicting results. Mishmar et al. (2004) reported a striking deficit of TEs within 15 bp of the numt boundary, and an excess of TEs between 15 and 150 bp from the boundary, for 247 human numts. In contrast, Gherman et al. (2007) described a decreased proportion of TEs across the entire first 500 bp flanking 266 human numts, with a monotonically increasing TE frequency moving away from the point of insertion for at least 1000 bp. Our data differ in being based on a smaller number of numts, but we have the advantage of using a closely related species to accurately define the numt insertion point in the genome. In addition, because we are only examining recent numts, the TE content of the chimpanzee genomic sequence likely represents the state at the time of insertion because the majority of human TEs inserted before the human–chimpanzee divergence. In contrast, examining much older numts likely also includes counting TEs that inserted after the numt. To summarize, our observation of a decreased proportion of TEs near the numt insertion point is broadly in agreement with Mishmar et al. (2004) and Gherman et al. (2007). However, we were unable to replicate the results of Gherman et al. (2007) in finding a continuously increasing proportion of TEs with increasing distance from the numt, even when using a data set composed of the flanking sequences of all identifiable human numts ( $n = 403$ ; data not shown).

Although the pattern of numts inserting into low TE regions is clear, a mechanistic explanation is less forthcoming. TEs may induce conformational changes in DNA, which may make them less susceptible to double-strand breaks or more likely to lead to correctly repairing such breaks when they occur without incorporating extranuclear DNA. TEs themselves integrate nonrandomly into the primate genome, with LINEs more often found in GC-poor areas of the genome, whereas SINEs show a preference for GC-rich genic regions (International Human Genome Sequencing Consortium 2001; Gasior et al. 2007). As such, it may be likely that the negative association between numts and TEs is caused by a co-correlation with some other unknown factor. Finally, it may be that numts that have inserted into TEs may be more frequently removed from the genome by way of nonhomologous recombination with another copy of that TE or other mechanisms to excise TEs.

Although previous studies have identified microhomology between the target site and the mtDNA, these have been limited to only a handful of occurrences (Zischler et al. 1995a; Blanchard and Schmidt 1996; Ricchetti et al. 1999) before the recent study by Hazkani-Covo and Covo (2008). By the use of a closely related outgroup, we were able to accurately define the extent of microhomology present in  $>100$  recent human- and chimp-specific numt insertions. Nearly all numts contain some degree of microhomology, at least at one end. Our quantification of microhomology is conservative in that for human numts we are comparing modern human mtDNA with chimpanzee nDNA as proxies for the molecules present at the time of integration (and vice versa for chimp-specific numts). As such, any mutations that have occurred since that time may be obscuring more substantial matches. This is especially relevant considering the high mutation rate of mtDNA. In addition, short stretches of microhomology that occur in positions not immediately adjacent to the boundary of numt insertion were not considered in the quantitative analysis. These microhomologies, along with the common occurrence of small deletions and insertions of flanking repeats, tandem repeats, and inverted repeats, are the hallmarks of the NHEJ pathway of double-strand break repair (Varga and Aplan 2005). The occasional presence of longer stretches of microhomology ( $\geq 7$  bp), and recessed microhomology, may indicate the use of an alternative end-joining mechanism (Daley and Wilson 2005; Corneo et al. 2007; Decottignies 2007; Yan et al. 2007), but the data presented here suggest that this is rare, at least for mammalian numts. Close examination of the numt junctions, along with the outgroup for comparison, provides some hints as to the inexactness of this repair pathway and confirms its description as “dirty” (Odersky et al. 2002). The reliance on short microhomology to complete the double-strand break repair with NHEJ further indicates a nonrandom component to numt insertion. Although this mechanism does

not specifically target genomic locations based on long stretches of homology, it does suggest that there must be some portions of the genome where no sufficient microhomology could be found, leading to a failure to repair DNA damage and thus to cell death. As such, we can only observe numts that originated in germline cells that were able to successfully complete the repair process.

Although the insertion of numts into the genome is often considered random, we show here that primate numts do not insert uniformly through time nor randomly with respect to genomic location. Although mechanistic explanations for these patterns remain elusive, the impending availability of several more hominoid genome sequences may likely provide clues. The use of comparative data to accurately determine the preintegration site is shown here to be essential and provides evidence that microhomology and small indels are frequently associated with integration, reinforcing the potential role of NHEJ in numt insertion.

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