Research

Coevolution of retroelements and tandem zinc finger genes

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Vertebrate genomes encode large and highly variable numbers of tandem C2H2 zinc finger (tandem ZF) transcription factor proteins. In mammals, most tandem ZF genes also encode a KRAB domain (KZNF proteins). Very little is known about what forces have driven the number and diversity of tandem ZF genes. Recent studies suggest that one role of KZNF proteins is to bind and repress transcription of exogenous retroviruses and their endogenous counterpart LTR retroelements. We report a striking correlation across vertebrate genomes between the number of LTR retroelements and the number of host tandem ZF genes. This correlation is specific to LTR retroelements and ZF genes and was not explained by covariation in other genomic features. We further show that recently active LTR retroelements are correlated with recent tandem ZF gene duplicates across vertebrates. On branches of the primate phylogeny, we find that the appearance of new families of endogenous retroviruses is strongly predictive of the appearance of new duplicate KZNF genes. We hypothesize that retroviral and LTR retroelement burden drives evolution of host tandem ZF genes. This hypothesis is consistent with previously described molecular evolutionary patterns in duplicate ZF genes throughout vertebrates. To further explore these patterns, we investigated 34 duplicate human KZNF gene pairs, all of which underwent an early burst of divergence in the major nucleotide contact residues of their ZF domains, followed by purifying selection in both duplicates. Our results support a host-pathogen model for tandem ZF genes.

[Supplemental material is available for this article.]

Vertebrate genomes contain large and highly variable numbers of tandem C2H2 zinc finger (tandem ZF) transcription factor genes. Outside of mammals, almost nothing is known about the function of these genes. Within mammals, tandem ZF genes are dominated by those with a KRAB domain, which expanded from a KRABtandem ZF fusion gene near the root of tetrapod vertebrates (Birtle and Ponting 2006). Most KRAB zinc finger (KZNF) proteins consist of an N-terminal KRAB domain followed by multiple tandem ZF domains (Bellefroid et al. 1991; Huntley et al. 2006). Some KZNF proteins have an additional SCAN protein-interaction domain N-terminal to their KRAB domain (Edelstein and Collins 2005). The KRAB domain represses transcription by binding TRIM28 (also called KAP1), which is part of a large protein complex that modifies histones to promote closed chromatin (e.g., Nielsen et al. 1999; Ryan et al. 1999; Lechner et al. 2000; Schultz et al. 2002; Sripathy et al. 2006). The tandem ZF domains confer DNA-binding specificity in a modular manner, with a turn-helix segment of each ZF domain binding to three nucleotides in target DNA sites (Pavletich and Pabo 1991; Kim and Berg 1996).

Tandem ZF genes have been gained by an ongoing process of lineage-specific duplication and divergence (e.g., Shannon et al. 2003; Emerson and Thomas 2009; Nowick et al. 2010). Throughout vertebrates, tandem ZF gene expansions are characterized by strong positive selection that has changed the number and DNAbinding specificity of zinc fingers, while retaining a conserved KRAB domain (Schmidt and Durrett 2004; Emerson and Thomas 2009). Though most tandem ZF genes outside of mammals lack a KRAB domain, the structure and evolution of their zinc fingers is strikingly similar to that of KZNF genes in mammals (Emerson and

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Thomas 2009). These evolutionary patterns, combined with remarkably little functional information, have given rise to a set of long-standing puzzles. What are the organismal functions of tandem ZF genes? Why are there so many genes, and why do the gene numbers and their repertoire of DNA-binding sites change so quickly?

Recent work suggests a plausible functional explanation for KRAB tandem ZF gene evolution. First, a series of papers showed that restriction of murine leukemia virus (MLV) in mouse embryonic stem cells results from transcriptional repression by the mouse-specific KZNF gene Zfp809 acting via the TRIM28 complex (Wolf and Goff 2007, 2009). Zfp809 binds integrated MLV DNA at its primer binding site (PBS), which MLV requires to prime reverse transcription via a complementary host tRNA (Wolf and Goff 2009). Shortly thereafter, two papers showed that deletion of TRIM28 (or the TRIM28 effector SETDB1) in mouse embryonic cells causes massive transcriptional derepression of several mouse endogenous retroviruses (ERVs) (Matsui et al. 2010; Rowe et al. 2010). Since TRIM28 and SETDB1 are thought to be shared effectors of all KZNF proteins, this result suggests that a suite of KZNF genes repress transcription of diverse ERVs (for review, see Rowe and Trono 2011). These findings have the potential to explain the large number of evolutionarily volatile KZNF genes based on a host-pathogen interaction. ERVs are the genomic footprints of historical retroviral infections that resulted in viral insertions in the germ line (for review, see de Parseval and Heidmann 2005; Blikstad et al. 2008). When an ERV first appears in a genome, it typically appears as a burst of multiple elements, due either to recurrent viral insertions or to transposition (Belshaw et al. 2004, 2005). This process results in a genomic signature that reflects a sampling of the history of retroviral infections, or at least the subset of infections that successfully established a germ-line copy. Based on the diversity and ages of ERVs present today, both the mouse and human lineages have episodically suffered a large

Coevolution of retroelements and zinc finger genes

number of infections by diverse retroviruses over the past 80 million years (e.g., de Parseval and Heidmann 2005; Stocking and Kozak 2008). Less extensive analysis suggests that a similar process occurs throughout mammals (e.g., Mouse Genome Sequencing Consortium 2002; Mikkelsen et al. 2007).

Presumably, both new retroviral infections and divergence of endogenous LTR retroelements will drive selection for a host response. One possible host response is the generation of new transcriptional repressors that evolve to target the DNA of the new retrovirus or retrotransposon. The simple modular biochemistry of KZNF transcriptional repression makes KZNF genes particularly suitable for such a role. The existing TRIM28 complex should require only recruitment by a new DNA-binding specificity to result in a repressed chromatin state. This repressed state can spread many kilobases from the DNA-binding site (Groner et al. 2010), so, in principle, binding anywhere in a retroelement could provide effective repression.

Here, we present data that supports the hypothesis that most vertebrate tandem ZF genes evolved to repress retroviral or LTR retroelement activity.

Results

LTR retroelement correlation with ZF domains and genes

Endogenous retroviruses and LTR retrotransposons differ primarily in the presence or absence of an envelope (ENV) coding sequence. These two types of retroelements interconvert by gain or loss of ENV sequences, and the ENV-coding sequence is extremely diverse and rapidly evolving, making it difficult to distinguish these two groups without detailed analysis. We will refer to both types of elements as LTR retroelements, and we did not attempt to distinguish them in our analysis. Our initial goal was to test for correlation between LTR retroelement load and tandem ZF-coding potential across a wide range of vertebrate genomes. Existing genome annotations are uneven, so we implemented genome searches to detect both LTR retroelements and ZF domains in a manner independent of annotation status and phylogeny. In the case of retroelements, this was made possible by the fact that LTR retroelements include ancient protein-coding domains that distinguish them from all other known genomic features (e.g., for review, see Gogvadze and Buzdin 2009). In the case of ZF domains, this was made possible by the fact that the C2H2 ZF domain has the same length and sequence profile throughout animals (Emerson and Thomas 2009). Details of both searches are given in the Methods section and the Supplemental Methods.

We found a striking correlation across vertebrates between the number of LTR retroelements and the number of ZF domains. This correlation holds within mammals and outside of mammals, and when all 26 taxa are combined (Fig. 1; Supplemental Fig. S1). To account for the fact that shared phylogenetic history probably accounts for some of this correlation, we computed corrected correlations and P-values (Fig. 1; Table1) using the method of independent contrasts (IC) (Felsenstein 1985; Garland et al. 2005). The IC-corrected correlations remained strong and highly significant and were robust to widely different score thresholds for counting LTR retroelements and ZF domains, and to counting the total number ZF domains or the number of putative tandem ZF genes (Table 1; Supplemental Table S1). Given existing evidence that KRAB ZF genes can repress retroviral and LTR retrotransposon transcription, the most obvious inference is that the historical LTR retroelement content of each genome has driven the number of ZF

domains, but we considered other possible explanations. First, it seemed possible that each genome has a characteristic rate of segmental duplication or duplicate retention, and that this rate drives both LTR retroelement and ZF domain content. We examined this possibility by testing for IC-corrected correlations of non-LTR (LINE-like) retroelements with ZF domains and of LTR retroelements with other large dynamic protein domain families (olfactory receptor and immunoglobulin C1 and V domains). None of these correlations were statistically significant (Table 2). Second, it seemed possible that unknown constraints on genome size influence the potential for LTR retroelement and ZF domain content in each genome. We tested this possibility by normalizing LTR retroelement content to genome size and testing the normalized correlation to ZF domains. The correlation remained highly significant (Table 2). Given that LTR retroelements are a significant contributor to genome size, it is unsurprising that genome size itself positively correlates with LTR retroelement and ZF domain content, though these correlations were weak and statistically nonsignificant after IC correction (Table 2).

Other features of these data can be explained by a model in which LTR retroelements drive tandem ZF gene evolution. First, testing various score cutoffs for counting ZF domains showed that the correlation to LTR retroelements is strongest very near the score that best distinguishes human ZF domains in known genes from those in pseudogenes (Table 1; Supplemental Fig. S2). This result suggests that the correlation is stronger for functional ZF genes than for pseudogenes. Second, LTR retroelement correlation to total ZF domain number was slightly stronger than to the number of putative tandem ZF genes (Table 1). This result suggests that the total DNA-binding potential of ZF genes is more important than the number of genes. Finally, the most prominent correlation outlier in mammals is mouse, which has fewer ZF domains than predicted by its LTR retroelement content (Fig. 1). The mouse genome is known to have several groups of recently and currently active ERVs (Stocking and Kozak 2008), suggesting the possibility that the host ZF response is lagging behind a recent burst of LTR retroelement activity. Alternative explanations of this mouse result are considered in the Discussion.

If the major function of ZF genes is to transcriptionally repress LTR retroelements, then the sequence diversity of LTR retroelements should be an important factor in driving ZF number. We estimated the relative sequence diversity of LTR retroelements in each species by extracting their reverse-transcriptase coding regions and measuring their total protein tree length. Unsurprisingly, we found that retroelement diversity correlates strongly with retroelement number, so it was difficult to distinguish the influence of copy number and copy diversity. As expected given this result, retroelement diversity also strongly correlated with ZF number, though not quite as strongly as did retroelement copy number (Supplemental Table S1).

The 26 genome assemblies analyzed above are all based on more than fivefold sequence read coverage, but they vary in read coverage and in the degree of assembly finishing (Supplemental Fig. S1). This variation could affect the apparent retroelement and ZF gene content differentially, for example, by collapsing recent ZF gene duplicates into apparent single genes. Such variation in assembly quality is difficult to detect and control for, but we made one simple test by repeating the analysis only on the 16 published genomes (higher than average read coverage and finishing effort). Using the LTR retroelement and ZF domain cutoffs that gave the best correlation across all species, the IC-corrected correlation for published genomes was higher than for all genomes (R² 0.71 vs. R² 0.67) and the correlation remained highly significant despite the



Figure 1. Correlation of genomic LTR retroelement and ZF domain content: Three panels show the number of detected LTR retroelements plotted against the number of detected ZF domains in different vertebrate groups or all groups combined. The lines show the linear least-squares best fit with its squared correlation coefficient R². The fourth panel (*lower right*) shows the combined data after correction for phylogenetic relatedness by the method of independent contrasts. The line is the linear least-squares best fit forced to go through the origin and its associated R² and *P*-value. The summed score cutoff for LTR retroelements was 50 (see Supplemental Methods). The ZF number was determined from all genomic open reading frames with four or more ZF domain matches with a minimum average score of 55. These counting criteria gave the maximum correlation for combined data, but a wide variety of other counting criteria also gave highly significant correlations (Supplemental Table S1).

smaller data set (*P*-value 7.2×10^{-5}). This result suggests that improved genome assemblies will most likely improve the observed correlations.

In addition to LTR retroelements containing part or all of their internal sequences, vertebrate genomes contain large numbers of solo LTR sequences that arise by recombination between flanking LTRs (Copeland et al. 1983). It is very difficult to obtain unbiased counts of solo LTRs, because they have no generally shared sequence features. In addition, solo LTRs are more abundant for older retroelements, because they have had more time to recombine since their original insertion. Nevertheless, we assessed correlation between ZF sequences and total annotated LTR retroelement sequence content (including solo LTRs) for the 16 published vertebrate genomes, since they have the best annotated general repeat content. The IC-corrected correlation was significant ($\mathbb{R}^2 0.36$, P =0.018), though weaker than for elements with internal sequence. This lower correlation could result from uneven annotation of solo LTR sequences, the expected skew toward older retroelements that are less reflective of recent selective pressure on ZF genes, or other unknown factors.

Though mammalian tandem ZF genes are dominated by those encoding a KRAB domain, this domain association is less common in other tetrapods and is absent in fish (Supplemental Table S1; Looman et al. 2002; Birtle and Ponting 2006; Emerson and Thomas 2009). The fact that ZF domain content correlates strongly with LTR retroelement content throughout all of these groups suggests that the function of non-KRAB ZF genes in other tetrapods and in fish is related to the function of KRAB ZF genes in mammals. This inference is also supported by similarities in sequence evolution of tandem ZF genes in each of these groups (see below; Emerson and Thomas 2009). We speculate that other domains in these taxa play a role analogous to the KRAB domain in mammals, or that tandem ZF proteins bound to DNA can directly repress transcription.

Recent LTR retroelement activity

The data above reflect an historical aggregate of LTR retroelement activity and tandem ZF gene duplications. To test whether these characters are temporally correlated, we estimated the age of LTR retroelement insertions based on divergence between the two long terminal repeats of each retroelement (Johnson and Coffin 1999) and the age of ZF gene duplicates based on synonymous site divergence (d_s). Though mutation rates surely vary among the species, this variation should not affect relative divergence rates of ZF

Coevolution of retroelements and zinc finger genes

Table 1.	Statistical tests for correlation of LTR retroelement counts and ZF domain counts
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		ZF min	Score 40 ^a	ZF mi	nScore 45	ZF mi	nScore 50	ZF mi	nScore 55	ZF mi	nScore 60
LTR cutoff ^b	ZF count type ^c	R squared ^d	<i>P</i> -value ^d	R squared	P-value	R squared	P-value	R squared	P-value	R squared	P-value
minScore 80	ORF count ZF count	0.300 0.517	$\begin{array}{c} 3.8 \times 10^{-03} \\ 3.5 \times 10^{-05} \end{array}$	0.318 0.548	$\begin{array}{c} 2.7 \times 10^{-03} \\ 1.6 \times 10^{-05} \end{array}$	0.363 0.606	$\begin{array}{c} 1.1 \times 10^{-03} \\ 2.8 \times 10^{-06} \end{array}$	0.436 0.671 ^e	$2.4 imes 10^{-04}$ $3.1 imes 10^{-07}$	0.575 0.668	7.2×10^{-06} 3.4×10^{-07}
minScore 150	ORF count ZF count	0.243 0.455	1.0×10^{-02} 1.6×10^{-04}	0.261 0.486	$7.7 \times 10^{-03} \\ 7.6 \times 10^{-05} \\ 0.02 \\ 0$	0.303 0.546	3.6×10^{-03} 1.6×10^{-05}	0.375 0.617	$\frac{8.8\times10^{-04}}{2.0\times10^{-06}}$	0.526 0.638	$\begin{array}{c} 2.7\times 10^{-05} \\ 9.8\times 10^{-07} \end{array}$
minScore 200	ORF count ZF count	0.222 0.424	1.5×10^{-02} 3.1×10^{-04}	0.238 0.455	1.1×10^{-02} 1.6×10^{-04}	0.281 0.516	5.4×10^{-03} 3.6×10^{-05}	0.352 0.590	1.4×10^{-03} 4.6×10^{-06}	0.513 0.622	3.9×10^{-03} 1.7×10^{-06}

^aThe minimum rpsblast score required for each ZF domain match, as used for the two types of ZF counts. For example, if minScore is 40, then ORF count is the number of ORFs with four or more ZF domains at or above score 40, and ZF count is the number of ZF domains in all ORFs at or above score 40. ^bScore cutoff for counting an LTR retroelement match in each genome.

^cZF counts were made either using the number of ORFs containing four or more ZF domains (ORF count) or by counting the total number of ZF domains in ORFs with four or more ZF domains (ZF count). $^{d}R^{2}$ and *P*-values were computed by the method of independent contrasts (Midford et al. 2005; Maddison and Maddison 2010).

^eFigure 1 is a graph of the data for these cutoff values (the peak for the values in this table).

genes and LTR retroelements within a species. Using 5% and 10% divergence cutoffs, we found significant IC-corrected correlations between recent LTR retroelement activity and recent tandem ZF gene duplications across the combined taxa (Supplemental Table S3). The highest correlation was between sequence diversity among recently active LTR retroelements and the number of ZF domains in recent tandem ZF gene duplicates (Fig. 2), but comparisons of the numbers of LTR retroelements and ZF gene duplicates were also highly significant (Supplemental Table S3). Mouse, opossum, and lizard show evidence of especially high recent LTR retroelement activity, and all three species have correspondingly high numbers of recent tandem ZF gene duplicates (Fig. 2, compare the steepness of the curves near the origin). In addition, all three species show possible evidence of an earlier period of relatively quiescent LTR retroelement activity associated with fewer tandem ZF gene duplicates, mental Methods). We could divide the primate lineage into six distinct branches based on available sequences: a basal primate branch (before the divergence of basal primates), a Simian branch (before the divergence of New World from Old World monkeys), a Catarrhine branch (before the divergence of Old World monkeys from apes), a Hominoid/Hominid branch (before the divergence of orangutan from human; this branch is bisected by gibbons, which currently lack a whole-genome assembly), a Hominina branch (before the divergence of chimpanzee from human), and a humanspecific branch. Using a combination of insertion-site analysis and sequence trees of retroelement internal sequences, we defined the branch on which each of 48 primate-specific LTR retroelement families first appeared (Fig. 3; Supplemental Table S4). Another four retroelement families were imperfectly resolved, appearing just before or just after the divergence of New World monkeys.

curve. Alternatively, these plateaus could result from a higher rate of deletion removing older LTR retroelements. Except for a very recent drop in LTR retroelement activity, the patterns on the human lineage suggest relatively slow and constant rates of LTR retrotransposition and ZF gene duplication. Other genomes with relatively low recent LTR retroelement activity (e.g., horse, elephant, and medaka) have curves broadly similar to that of human (data not shown).

as evidenced by the plateaus on each

Primate LTR retroelements and tandem ZF gene duplicates

Among existing vertebrate genome sequences, primates provide the densest phylogeny and the best annotation of LTR retroelements. Using RepeatMasker annotations and tandem ZF gene annotations in humans as a starting point, we investigated in detail the appearance of new LTR retroelements and tandem ZF genes on the primate lineage (Supple-

Table 2. Control correlations

	R squared ^a	<i>P</i> -value ^a
LINE-like ^b ZF4 count minScore 55 ^c	0.063	0.26
LTR minScore 80 ^d mammal Olf genes ^e	0.015	0.67
LTR minScore 80 mammal Olf pseudogenes ^e	0.003	0.84
LTR minScore 80 IG C1 domain count ^r	0.089	0.14
LTR minScore 80 IG V domain count ^f	0.027	0.42
ZF4 count minScore 55 LTR minScore 80 normalized to genome size ⁹	0.581	$6.0 imes 10^{-06}$
LTR minScore 80 genome assembly size ^h	0.048	0.29
ZF4 count minScore 55 genome assembly size	0.385	0.11
LTR minScore 80 LINE-like	0.183	0.05

^aR² and P-values were computed on the transformation by the method of independent contrasts. ^bLINE-like elements counted for each genome (Supplemental Methods).

^cZF4 counts for ORFs with four or more ZF domains as described for Table 1.

^dLTR retroelements counted for each genome with a minimum score of 80 (Supplemental Methods).

^eMammalian olfactory gene and pseudogene counts were taken from Hayden et al. (2010). Data were available for all the mammals except baboon and marmoset.

^fIG C1 (immunoglobulin constant domain type 1) and IG V (immunoglobulin variable domain) domain counts were made from genomic searches with profiles PF07654 (C1-set) and PF07686 (V-set) with minimum rpsblast scores of 40 and 35, respectively. Score cutoffs were chosen to reflect the approximate number of each domain in the human genome.

⁹LTR retroelement counts were divided by the genome assembly size before computing the correlation statistics.

^hGenome assembly size was computed by counting the number of A, C, G, and T residues directly from the genome assemblies used for all analyses.



Figure 2. ZF gene duplicate and LTR divergence time courses. Three panels show cumulative histograms of LTR nucleotide divergence and closest ZF paralog d_s for the indicated species. The axes have been scaled to best display the full curve for both data sets. The human data are included in all three panels for comparison. The fourth panel (*lower right*) shows statistical analysis at or below one divergence point (0.05 LTR divergence/0.05 paralog d_s) for all species combined after correction by independent contrasts (see Supplemental Table S3). The line is the linear least-squares best fit forced to go through the origin and its associated R² and *P*-value.

Starting with annotated human tandem ZF genes, we used a combination of genome sequence searches, maximum-likelihood trees, and synteny to determine the branch on which each new tandem ZF gene duplicate appeared. Among ZF gene duplicates, we distinguished between those that diverged by at least 5% in amino acid sequence in an attempt to distinguish between selected duplicates and possibly neutral copy-number variation. Since events on each branch of the phylogeny are statistically independent, we analyzed correlations without using independent contrasts. Correlation between appearance of new LTR retroelement families and new tandem ZF genes was remarkably strong and statistically significant regardless of whether or not ZF genes with low divergence were included and regardless of branch assignment of the four ambiguous retroelement families (Supplemental Table S7). The correlation was highest when ZF genes with low divergence were excluded and the ambiguous retroelement families were split equally on the two possible branches (Fig. 3, statistics). These results are consistent with our global analysis of vertebrate correlations, suggesting that many or most tandem ZF genes in primates arose in response to the appearance of new families of endogenous retroviruses. The most prominent deviation from perfect correlation is on the Hominoid/Hominid branch, where no new LTR retroelement families, but 14 new tandem ZF genes appeared. The immediately preceding Catarrhine branch was subject to a particularly intense burst of new LTR retroelements (Fig. 3); we speculate that some of the 14 Hominoid/ Hominid ZF genes arose in response to this slightly earlier burst. In contrast to the human-specific branch, new LTR retroelement families have arisen on the chimpanzee- and macaque-specific branches (Jern et al. 2006; Polavarapu et al. 2006; Rhesus Macaque Genome Sequencing and Analysis Consortium 2007), but the number of families is small, and we lack statistical power to test for tandem ZF gene response.

Predictions for duplicate gene divergence

The hypothesis that most tandem ZF genes function to repress LTR retroelements predicts certain patterns of molecular evolution driven by the epidemiology of retroelements. First, each host genome should acquire distinct expansions of tandem ZF genes in response to lineage-specific retroelement challenges. Second, the duplicate genes that comprise these expansions should be subject to positive selection to modify their DNA-binding specificity as they adapt to new retroelements. These two predictions have already been confirmed for several of the genomes we analyzed here,



Figure 3. Primate phylogeny with the appearance of new endogenous retroviral families and new tandem ZF genes. Data were derived by tracking the first appearance of human ERV families and tandem ZF genes (Supplemental Methods). On each branch leading to the human, the top number indicates the number of tandem ZF gene duplicates (with additional duplicates that diverged by <5% in amino acid sequence in parentheses) and the bottom number indicates the number of new ERV families. Four families of ERVs could not be confidently assigned to a specific branch and are shown straddling the Simian/Catarrhine branch point. The R² and ANOVA P-value shown are for the peak correlation based on various criteria for partitioning the data (see Supplemental Table S7); all other partitions were also significant. The gorilla genome is low coverage and was not systematically analyzed, but the single ERV (HERV-Fc1) that appears on the branch leading to human and chimpanzee is clearly present in gorilla (data not shown). The Hominoid and Hominid branches are split by the gibbons, for which there are currently no genome assemblies.

including human, mouse, cow, frog, fugu, and zebrafish (Emerson and Thomas 2009). We extended these analyses to several additional species, namely, rat, horse, elephant, opossum, lizard, tetraodon, medaka, and lamprey. In all cases, we found large species-specific clades of tandem ZF genes with overwhelming evidence of positive selection affecting predominantly nucleotide contact residues of ZF domains (Supplemental Table S8; Supplemental Fig. S6).

A final prediction of our hypothesis is that often when a new divergent duplicate tandem ZF gene pair arises, one of the duplicates will retain the ancestral DNA-binding specificity, while the other duplicate acquires a new or modified DNA-binding specificity that targets a new retroelement. After optimizing its new DNA-binding function, the divergent duplicate should be subject to purifying selection. These patterns are expected in cases in which repression of an ancestrally targeted retroelement remains selectively significant, so that only one copy of a duplicate gene pair is free to alter specificity to protect against a new challenge. These patterns are also predicted when the ancestral tandem ZF gene has been exapted for host transcriptional regulation and by some other hypotheses for ZF gene evolution (see Discussion). As described in the next sections, we explored these predictions among human duplicate KZNF genes, since they are best annotated and there exist multiple closely related primate genomes that help resolve the time of duplication and ancestral gene identity.

Tracing the origins of human KZNF genes

We identified 34 cases in which two human KZNF genes were clearly closest relatives (see Methods); each pair is assumed to have arisen by gene duplication from an ancestral gene at some time during tetrapod evolution. In order to trace the evolutionary history of each pair of genes, we used the two human proteins to find all closely related sequences in a set of increasingly divergent mam-

Coevolution of retroelements and zinc finger genes

malian genomes. Because of the patterns of conservation and divergence detailed below, these searches were remarkably effective in unambiguously tracing the ancestry of each gene.

Considering one duplicate pair, one common result was as follows. In one or more of the most closely related species, one clear copy of each gene was found, indicating that both genes were present in the last common ancestor of human and those species. On deeper branches in the tree, each species had only one gene closely related to the two human genes, suggesting that their last common ancestral species had one copy of the gene, which later duplicated and diverged on the human lineage. The other common result was that clear copies of both genes were found back to some point on the phylogenetic tree, but deeper in the tree no specific ancestral genes were found (more accurately, many possible ancestral genes were found, but it was unclear which of them was the true ancestor). Examples of trees reflecting these assignments are given in Supplemental Fig. S8. Below, we consider the latter case first, in which the precise origin of two closely related human genes is unclear, but the pattern of divergence of the two copies from each other can nevertheless be analyzed.

Divergence of gene pairs of uncertain origin

We could unambiguously identify and analyze 19 human duplicate gene pairs with two copies in a number of species but no clear specific ancestor. Table 3 summarizes key features of these duplicates and Supplemental Table S10 gives additional details. The phylogenetic depth of the traceable ancestry of the two genes varied from early in the primate lineage to early in the placental mammalian lineage. It may be presumed that the two genes arose by one or more rounds of duplication and divergence from some specific ancestral gene, but the identity and sequence of the ancestral gene is indeterminate. For example, clear copies of both *ZNF273* and *ZNF680* were identified from all five primate species, but no species outside of primates. A detailed example of sequence divergence patterns for one pair of genes is shown in Supplemental Figure S7.

Three patterns were apparent in most or all cases:

- 1. Orthologs of each gene were subject to purifying selection across the entire set of DNA-binding domains: Both the number of ZF domains and the amino acid sequence of each ZF domain are highly conserved. Averaged across 280 ZF domains from 22 genes randomly selected from these duplicates, the nucleotide and phosphate contact residues are among the slowest evolving (Fig. 4). The most plausible explanation for this pattern is that each orthologous ZF domain is subject to purifying selection to retain its DNA-binding specificity.
- 2. A second pattern is evident when comparing two duplicate genes to each other: Amino acid changes are more abundant in major nucleotide contact residues than elsewhere (18 of 19 duplicate pairs). When the divergence between the duplicates was relatively low, this difference did not reach statistical significance, but in 12 of 19 duplicates, changes were enriched in major nucleotide contact residues with P < 0.01 (Fisher's exact test). Summed over all 222 testable ZF domains from all 19 duplicate pairs, changes between paralogs that are conserved among orthologs occurred in 250 of 666 major nucleotide contact residues (37.5%), but only 489 of 3552 other residues (13.8%), a highly significant enrichment. This pattern is summarized graphically in Figure 4.
- 3. A third pattern is that entire ZF domains were often lost or gained in one duplicate gene relative to the other, consistent with previous observations (Looman et al. 2002; Huntley et al.

Table 3.	Summar	y of duplicate	pairs with an indeterminate ancestral gene	2
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Phylogenetic depth ^a	Duplication depth ^b	Human gene 1	Human gene 2	Inform- ative fingers ^c	Nucleotide contact changes ^d	Nucleotide contact adjacent changes ^e	Other changes ^f	P-val nucleotide contact vs. other ^g	Fingers indel ^h	Fingers defective ⁱ	<i>P</i> -val branch- specific pos selection ^j
Primate specific	> cjac	ZNF273	ZNF680	10	14	15	29	0.001	0	2	0.0005
Primate specific	> cíac	ZNF100	ZNF430	10	7	1	11	0.01	0	0	1.0000
Primate specific	> cjac	ZNF836	ZNF841	15	15	16	29	0.001	2	2	< 0.0001
Eutheria	deep	ZNF570	ZNF583	11	4	19	17	0.75	1	0	0.5398
Eutheria	deep	ZNF383	ZNF829	8	3	8	7	0.2	1	1	0.9287
Eutheria	deep	ZFP30	ZFP82	13	4	10	27	NA	0	0	0.0240
Eutheria	deep	ZNF264	ZNF805	13	5	4	10	0.07	0	0	0.0049
Eutheria	deep	ZNF226	ZNF234	16	12	13	10	<0.0001	0	1	<0.0001
Eutheria	deep	ZFP112	ZNF45	13	28	20	45	<0.0001	0	0	0.0050
Eutheria	deep	ZNF568	ZNF569	15	14	16	26	0.001	2	0	1.0000
Boreoeutheria	deep	ZNF354A	ZNF354B	13	2	5	1	0.07	0	0	0.1160
Eutheria	deep	ZNF619	ZNF621	7	2	8	7	0.43	3	0	0.8065
Primate specific	> cjac	ZNF564	ZNF136	13	25	41	59	<0.0001	0	1	0.2415
Primate specific	> cjac	ZNF124	ZNF670	6	14	14	27	0.0001	0	0	0.0144
Eutheria	deép	ZNF382	ZNF567	9	15	14	35	0.002	3	2	0.0022
Eutheria	deep	ZNF41	ZNF484	13	12	13	31	0.02	2	0	0.0821
Eutheria	deep	ZNF81	ZNF175	12	20	23	28	<0.0001	1	0	0.0831
Primate specific	> mmul	ZNF675	ZNF681	10	17	5	20	<0.0001	0	1	0.0019
Primate specific	> mmul?	ZNF528	LLNL759	15	37	30	70	<0.0001	0	0	< 0.0001
•			total sites:	222	666	1110	3552				
		tota change	al changes: frequency:		250 0.375	275 0.248	489 0.138	<0.0001	15	10	

^aOldest branch identified with an ortholog for either gene of the duplicate pair.

^bPhylogenetic branch on which the duplication occurred (> cjac = before marmoset, > mmul = before macaque, deep = before Boreoeutherian split). ^cNumber of ZF domains shared between the duplicate copies.

^dChanges that occurred at one of the three major nucleotide contact sites. For the three "changes" columns, changes in fingers are defined as amino acid residues that are invariant among all orthologous copies of a gene and different between the two duplicates.

^eChanges that occurred at a site immediately adjacent to a major nucleotide contact site (there are five such sites, because one is an invariant zinccoordinating H residue).

^fChanges that occurred at one of the remaining 16 sites (16 = 28 - 3 - 5 - 4 invariant zinc-coordinating residues).

⁹Result of a one-sided Fisher's exact test for whether the changed nt contact sites are more frequent than changed other sites, not corrected for multiple testing (NA: not applicable, because they are less frequent).

^hNumber of ZF domains involved in 28 amino acid indel changes between the duplicates (some indels involve more than one adjacent ZF domain). Number of ZF domains in which one duplicate copy has lost one or more zinc-coordinating residue.

^jP-value for positive selection from the branch-site model of codeml, with the branch joining the duplicates labeled (see Methods), not corrected for multiple testing.

2006). When such a difference was observed it was strongly conserved among orthologs of each of the two genes, suggesting that these domain arrangements are also subject to purifying selection. Some such events involved insertion or deletion of ZF domains and others involved point mutations that disrupt the canonical finger structure (Table 3). These results suggest that finger gain and loss contribute to changes in DNA-binding specificity between duplicate genes.

Duplicate divergence is asymmetric

In the 19 cases discussed above, the absence of an identified ancestral gene in the outgroup species precluded analysis of the symmetry of divergence following duplication. In the other 15 duplicate cases, the ancestral gene state could be identified based on the pattern of gene number and gene type in various species (Table 4; Supplemental Table S10). For example, copies of both human *ZNF557* and *ZNF558* were clearly identified in chimpanzee and orangutan, but only one related gene was found in macaque, marmoset, cow, dog, horse, and rodents, suggesting that a single ancestral gene duplicated on the branch leading to great apes. By comparison of the two duplicates with the single gene from outgroup species, we could address whether divergence occurred in one or both duplicate copies. As shown in Table 4, the results usually indicated highly asymmetric divergence of the duplicate genes. Alignments for two examples are shown in Supplemental Figure S7. Amino acid changes following duplication were strongly biased toward nucleotide contact residues: In total, conserved changes occurred in 155 of 504 major nucleotide contact residues (30.8%), but in only 250 of 2688 other residues (9.3%). After an initial period of divergence, the divergent copy became subject to purifying selection, since its copies in all descendant species are very similar in amino acid sequence. These patterns suggest that one duplicate retains the ancestral DNA-binding specificity, whereas the other duplicate acquires a new or modified DNA-binding specificity.

Apparent divergence asymmetry could result from genome assembly artifacts or repeated gene loss in specific lineages. A combination of measuring assembly completeness, dating duplicates by synonymous site divergence, and parsimony analysis of loss events, indicates that these artifacts cannot account for the asymmetry results (Supplemental Methods).

Positive selection following duplication

If new duplicate KZNF genes are subject to selection to acquire new DNA-binding specificities, codon-based methods for analyzing selection might be able to detect branch-specific positive selection. We used the branch-site maximum-likelihood models imple-



Figure 4. Changes in orthologous zinc fingers and duplicate zinc fingers compared with diversity among all fingers. (*Top*) The averaged relative rates of divergence in 280 orthologous ZF domains from 22 randomly chosen KZNF genes (see Methods). (*Middle*) The diversity among all ZF domains from human KZNF genes as a logo plot (http://weblogo.berkeley.edu). (*Bottom*) The number of conserved changes observed among the 390 testable zinc fingers in all 34 duplicate gene pairs analyzed. Circles labeled P indicate residues that make major phosphate contacts and squares labeled N indicate residues that make the major nucleotide contacts.

mented in codeml to test this possibility. For 17 of the 34 duplicates, highly significant evidence (P < 0.01) was obtained for positive selection on the branch connecting the two duplicate copies (Table 4). In many cases, the number of available sequences and their total tree length are well below the optimum for detection of positive selection by this method (Anisimova et al. 2001), so it is possible that divergence of all of the duplicates involved positive selection, but reached statistical significance only in the strongest cases. As expected, the specific residues implicated in positive selection are strongly enriched in the major nucleotide contact residues (data not shown). These results directly support the idea that initial duplicate divergence is driven by selection to acquire new DNA-binding specificity.

Coevolution of retroelements and zinc finger genes

Stable genes

Though duplication is common in the KZNF family, some genes are old and highly conserved. Marsupial mammals diverged from placental mammals about 180 Mya (Kumar and Hedges 1998; Mikkelsen et al. 2007). Using systematic genome searches, we identified 20 human KZNF genes that have clear orthologs in the marsupialopossum genome and are present in all or nearly all placental mammalian genomes (ZKSCAN1, ZNF3, ZNF18, ZNF192, ZNF202, ZNF205, ZNF212, ZNF213, ZNF263, ZNF282, ZNF398, ZNF436, ZNF446, ZNF496, ZNF641, ZNF746, ZNF764, ZNF777, ZNF783, and ZNF786). Each gene was present in single copy in opossum and throughout placental mammals, and their ZF domains were invariant in number and highly conserved. These results indicate that some KZNF genes adopted stable functional roles early in mammalian evolution and that they have subsequently retained the same DNA-binding specificity. Explanations that reconcile this finding with the retroelement hypothesis are given in the Discussion.

Discussion

Based on the striking correlations between LTR retroelement content and C2H2 ZF domain content throughout vertebrates and over time, we propose that most tandem ZF genes originate to repress transcription of LTR retrotransposons or retroviruses. The linear-regression lines for the raw correlations pass close to 0 (the origin) LTR retroelements and ZF domains (Fig. 1), suggesting that most or all tandem ZF genes are involved. Consistent with this hypothesis, recent publications show that a mouse KZNF gene represses murine leukemia retrovirus (Wolf and Goff 2009) and that an unknown suite of KZNF genes probably repress a wide variety of IAP and MusD LTR retroelements (Matsui et al. 2010: Rowe et al. 2010). The vast majority of tandem ZF genes have no experimentally determined organismal function, a situation fully compatible with retroelement repression because this function should be difficult to ascertain. Nevertheless, a handful of tandem ZF genes are implicated in other processes, including sex-limited gene expression, imprinting, and mouse embryonic development (Krebs et al. 2005; García-García et al. 2008; Li et al. 2008; Mackay et al. 2008). Similarly, some KZNF genes arose early in mammalian evolution and have been retained throughout Therian mammals with nearly invariant DNA-binding domains. These genes are unlikely to have current-day retroelement-related functions, since there is no evidence for such widely shared retroelements. One plausible explanation is simply that some tandem ZF genes evolved directly to fulfill other host functions and that they were never involved in retroelement repression. Alternatively, it is well established that transcriptional promoters and enhancers present in retroelements are sometimes exapted for host transcription, following chance integration in an appropriate location to confer useful transcriptional regulation on a host gene (for review, see Cohen et al. 2009). A few studies provide indirect evidence that host exaptation of retroelement regulatory sequences may be extremely common (Lowe et al. 2007; Conley et al. 2008). In addition, in at least two cases a retroviral gene itself appears to have been adopted for a host function (Best et al. 1996; Dupressoir et al. 2009). Thus, it is possible that tandem ZF genes that now function as host transcription factors evolved initially to repress LTR retroelements and were later retained on the basis of their regulatory role for a host gene.

In mammals, transposition competence of new ERV families is usually relatively transient, decaying over a period of several million years; after transposition-specific ERV sequences evolve neutrally

Table 4.	Summary	of du	plicate	pairs	with	an	inferred	ancestral	gene
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Phylogenetic depth ^a	Duplication depth	Human gene 1 ^b	Human gene 2 ^b	Inform- ative fingers	Nucleotide contact changes	Nucleotide contact adjacent changes	Other changes	P-val nucleotide contact vs. other	Asymmetry score ^c	Fingers indel	Fingers lost	P-val branch- specific pos selection
Primate specific	> ppyq	ZNF431	ZNF714	12	13	5	15	<0.0001	0.91	0	0	<0.0001
Primate specific	> ppyq	ZNF679	ZNF716	7	3	7	7	0.19	0.41	3	0	0.0377
Primate specific	> mmul	ZNF160	ZNF665	18	15	12	32	0.002	1.00	2	0	0.0180
Primate specific	> ppyq	ZNF468	ZNF28	10	3	4	8	0.24	0.93	7	0	0.0059
Primate specific	> ppyq	ZNF611 ^d	ZNF600	14	8	7	5	0.0001	0.47	3	3	< 0.0001
Primate specific	> ppyq	ZNF799 ^d	ZNF443	13	3	5	7	0.20	0.87	1	0	0.5463
Eutheria	> mmul	ZNF773	ZNF419	9	5	5	1	0.0004	1.00	2	0	0.0104
Eutheria	> mmul	ZNF33B ^d	ZNF33A	16	7	4	4	0.0003	1.00	0	0	0.0018
Eutheria	> cjac	ZNF585A	ZNF585B	21	5	2	2	0.001	0.56	0	0	0.4948
Primate specific	> ppyg	ZNF736 ^d	ZNF727	7	10	10	17	0.002	0.89	2	0	0.0013
Eutheria	> mmul	ZNF558	ZNF557	9	15	9	29	0.05	0.93	1	0	0.0296
Theria	> cjac	ZNF764	ZNF747	4	3	3	3	0.05	0.44	3	0	0.0024
Eutheria	> mmul?	ZNF133	ZNF343	10	23	17	41	<0.0001	0.95	2	1	0.4348
Primate specific	> mmul?	ZNF17	ZNF749	9	22	24	38	<0.0001	0.76	1	3	<0.0001
Euarchontoglires	> cjac	ZIK1	ZNF416	9	20	22	41	<0.0001	0.98	0	2	<0.0001
			totals:	168	155	136	250			27	9	
			sites:		504	840	2688					
		fre	equencies:		0.308	0.162	0.093					

^aMost column headers are as defined in Table 3. Euarchontoglires is the clade that includes rodents, lagomorphs, and primates.

^bThe inferred ancestral gene state is listed in the left column and the divergent duplicate gene in the right column.

^cThe asymmetry score can vary from 0 to 1 and is a measure of the extent to which amino acid changes occurred in the divergent duplicate relative to the conserved duplicate (see Methods). A score of 1 means that changes occurred exclusively in the divergent duplicate, and a score of 0 means that changes were equally distributed between the duplicates. Only changed sites in which all the orthologs had the same amino acid were counted (conserved changes).

^dThese duplicate gene pairs are also described in Nowick et al. 2010.

and eventually lose protein-coding function and transcription competence (e.g., de Parseval and Heidmann 2005; Stocking and Kozak 2008). During this transition, selection to retain specifically protective ZF genes will attenuate. Unless they are exapted for a distinct host function, most such ZF genes should eventually become pseudogenes or be deleted from the genome. This predicted pattern has not been analyzed in detail, but the general expectation of abundant ZF pseudogenes is clearly met in the human genome and probably in other genomes (Supplemental Fig. S2).

The state of ERVs and tandem ZF genes in the mouse is of particular interest, because further experimental tests of our hypothesis are most feasible there. Our data show that the mouse reference genome assembly has an unusually high number and diversity of ERVs relative to tandem ZF genes. One possible explanation is that tandem ZF gene response in mouse lags behind the recent high ERV activity that is known in mouse (Stocking and Kozak 2008), but alternative explanations are possible. First, mouse has a higher rate of genomic deletion than human (Mouse Genome Sequencing Consortium 2002), which should remove older ERVs more quickly, potentially freeing ZF genes involved in their defense for directional positive selection to protect against new ERV challenges. This possibility may be testable by a focused analysis of the patterns of duplication and positive selection among mouse ZF genes. Second, a strong recent evolutionary ZF response to high ERV activity is expected to result in an abundance of unfixed ZF gene duplicates, causing heterogeneity in the number of ZF genes in mouse populations. This possibility predicts that sequences from other wild Mus musculus isolates will vary in ZF gene content, with some isolates having more or fewer ZF genes than the reference genome. Finally, though the mouse genome assembly is one of the highest quality that we analyzed, it is possible that a recent burst of ZF gene duplication would be obscured by assembly collapse of multiple similar paralogs, resulting in an underestimate of ZF gene number.

In mammals, the large majority of LTR retroelements are clearly ERVs, as indicated by the presence of a viral envelope gene or close relatedness to a known retrovirus. Outside of mammals, this relationship is less clear. Fish appear to have relatively few ERVs and a large burden of LTR retrotransposons with no clear retroviral connection (Basta et al. 2007, 2009). In chicken and finch, most retroelements are classified by RepeatFinder as ERVs, but detailed analysis is lacking, and in lizard no analysis of LTR retroelements is available. Retroviruses have repeatedly evolved from vertically transmitted retrotransposons by acquisition of an envelope gene (e.g., Doolittle and Feng 1992; Laten et al. 1998; Malik et al. 2000). Conversely, integrated retroviruses can readily convert to vertically transmitted transposons by loss of the envelope gene (Ribet et al. 2008). It is possible that tandem ZF genes repress exogenous retroviruses, endogenous retroviruses, and LTR retrotransposons, but the balance of activities for these groups remains unclear and may vary in different species.

Implications for retroviral repression in mammals

Judging from patterns of endogenized retroviral sequences, mammals have been subject to an ongoing barrage of retroviral infections of diverse types (de Parseval and Heidmann 2005; Blikstad et al. 2008). New retroviral infections in a particular species can arise by a shift or expansion of host range by a retrovirus that infects another species (e.g., Benveniste and Todaro 1974; Chen et al. 1996; Gao et al. 1999; Martin et al. 1999). The consequence for the new host is the occasional appearance of an unpredictable new retroviral challenge. If a retrovirus successfully integrates in the germ cell lineage, it may also result in the spread of a new deleterious ERV in the host genome. It is well established that mammals combat retroviral infection in multiple ways, including attacking viral RNA with APOBEC cytidine deaminases and ZAP, interfering with viral capsid with Fv1 and TRIM5alpha, and preventing viral particle release with Tetherin (for review, see Wolf and Goff 2008). The pervasiveness of retroviruses in vertebrates and the multiple layers of viral restriction by the host support the idea that there should also be strong selection on the host to repress retroviral transcription. The size, diversity, and rapid evolution of the tandem ZF gene family suggests that these genes fill this role.

The sequence divergence patterns of new duplicate genes suggests the following model for the contribution of KZNF genes to host response to a new retroviral infection in mammals. Starting from either a new duplicate gene or a pre-existing copy-number polymorphism, a KZNF gene with significant, even minor, off-target binding to a new retroviral sequence is driven to fixation and starts to evolve improved target recognition by changes in amino acid sequence and changes in ZF number. This pattern of duplicate evolution corresponds in many ways to that proposed for bacterial genes (Bergthorsson et al. 2007). The initial off-target binding to a new retrovirus may arise purely by chance or may result from sequence relatedness of the new retrovirus to a previously encountered retrovirus for which the host has already evolved a cognate KZNF gene. If the previously encountered retrovirus (or its endogenized copies) remain selectively significant for the host, there will be pressure for one copy of the KZNF gene to retain its ancestral DNA-binding specificity and for adaptation to the new retrovirus to act on the other copy. If the previously encountered retrovirus is no longer selectively significant for the host, targeting a new retrovirus could be achieved by directional selection on an ancestral KZNF gene without gene duplication, though we didn't observe any clear instances of this pattern.

Other possible evolutionary drivers

A number of other potential drivers of tandem ZF gene duplication and divergence have been suggested and probably apply in specific cases. Based on the expansion and diversification of the KZNF gene sequence and expression patterns on the primate lineage, it has been suggested that these genes underlie the evolution of novel primate traits, including an enlarged brain (Hamilton et al. 2003; Nowick et al. 2009, 2010). Based on the expansion of genes in a cluster of KZNF genes in mouse that includes two genes that modify sex-limited expression of other genes, it has been suggested that KZNF genes play a role in speciation via modification of sexspecific traits (Krebs et al. 2005). One KZNF gene with an ortholog in mouse (Zfp57) and human (ZFP57) has been shown to be required for genomic imprinting at several loci (Li et al. 2008; Mackay et al. 2008). Since imprinting involves maternal-zygotic conflict (e.g., Smith et al. 2006), this process has the potential to drive KZNF duplication and diversification. Finally, the PRDM9 tandem ZF gene is strongly implicated in specification of recombination hotspots (Baudat et al. 2010; Myers et al. 2010; Parvanov et al. 2010). Though recombination hotspots evolve rapidly, the domain structure and evolution of PRDM9 are clearly different from all other tandem ZF genes (Oliver et al. 2009; Thomas et al. 2009), and it has not been subject to the expansion seen in the genes described here. None of these explanations alone suffice to explain the general correlations between genomic LTR retroelement content and tandem ZF coding potential. In contrast, the established potential for host exaptation

of retroviral regulatory elements provides a plausible mechanism by which tandem ZF genes initially selected for retroelement repression could, over time, adopt a variety of other host functions.

Coevolution of retroelements and zinc finger genes

Methods

Species key

A key for species abbreviations, common names, and genome assemblies is provided in the Supplemental Methods.

Retroelement counts

RepeatMasker data were unavailable for several genomes of interest and misleading for others, apparently because some genomes contain abundant retroelement sequences that do not yet appear in the RepBase sequences used as queries by RepeatMasker (AFA Smit, R Hubley, P Green. 1996–2010. RepeatMasker Open-3.0, http://www. repeatmasker.org). To make counts of retroelements in a manner independent of repeat annotation status and species phylogeny, we used the fact that LTR retroelements are distinguished from all other known sequences by the appearance of a characteristic pattern of conserved coding elements, namely protease, reverse transcriptase, RNaseH, and integrase domains (many retroelements also encode gag and env proteins, but these are poorly conserved across the broad phylogenetic space we wished to analyze). We used patterns of genomic matches to Pfam profiles for these domains to identify LTR retroelements as detailed in the Supplemental Methods.

ZF domain searches

We performed a search for zinc finger domains on selected genomes using the program rpsblast with the -p F option (6-frame translation of DNA query). The search profile consisted of a 28 amino acid weight matrix profile of the ZF domain (including the 7-amino acid linker region upstream of the 21-amino acid ZF core). This profile was generated from the set of functional human tandem ZF proteins using the psiblast program as directed in the NCBI blast documentation. Subsequent analysis showed that this profile is nearly identical to profiles derived from tandem ZF proteins from other species (examples shown in Supplemental Fig. S5). Genome searches were carried out in two forms: one search of the entire genome assembly and a second search of all open reading frame (ORF) segments of 100 codons or longer. From ORF searches, we counted: (1) the number of ORFs with one or more ZF match above some score cutoff, and (2) the total number of ZF matches above some score cutoff. Counts from all searches with various score cutoffs are reported in Supplemental Table S1. To be sure that the ZF domain matches reflect bona fide tandem ZF genes, we determined the ZF domain profile and the number of tandem ZF domains for each genome with large numbers of ZF genes (examples shown in Supplemental Fig. S5). To determine the ZF domain score distribution expected for genes and pseudogenes, all human ZF domain matches were divided into those in known RefSeq genes (plus a few probable genes not yet appearing in RefSeq) (Huntley et al. 2006) and those outside genes (which are likely to belong to pseudogenes and gene fragments). We made a density histogram of the rpsblast scores for each group (representing how well each hit matches the ZF profile) and superimposed the histograms (Supplemental Fig. S2). The pseudogene scores presumably reflect a distribution of times of neutral evolution since pseudogenization. The crossover point where the hit density for genes first exceeds the hit density for pseudogenes occurs at about rpsblast score 57. This crossover point is close to the peak correlations of LTR retroelements and ZF domains (Supplemental Table S1).

Phylogenetic correction by independent contrasts

Comparing two characters in a scatter plot assumes statistical independence, an assumption that is violated when related species are used as data points (Felsenstein 1985; Garland et al. 2005). This can create spurious correlations across broad phylogenies (Whitney and Garland 2010). To account for such phylogenetic concerns we tested our data using Felsenstein's independent contrasts method implemented in the PDAP package (Midford et al. 2005) of Mesquite (Maddison and Maddison 2010). Specifically, we used the positivized x vs. y contrasts (mode 9 in PDAP) to measure the correlation between respective zinc finger metrics and ERV metrics. The tree used in the analysis was based on best estimates for species divergence times derived largely from TimeTree (http://www.timetree. org/; Hedges et al. 2006). Pearson correlations were forced to go through the origin.

Identification of new LTR retroelement families and tandem ZF genes on primate branches

Phylogenetic branches on which new LTR retroelement families were added were determined using a combination of RepeatMasker annotations, shared insertion site analysis, and DNA trees of retroelement internal sequences (Supplemental Methods; Supplemental Fig. S4; Supplemental Table S4). Branches on which new tandem ZF genes were added were determined by identifying primate orthologs of human tandem ZF genes and determining the first appearance of each gene (Supplemental Methods; Supplemental Table S5).

Identification of closest human gene pairs and their orthologs in other species

Pairs of human duplicate KZNF genes were identified as reciprocal best blastp matches or as neighbors on a pairwise distance tree among all human KZNF genes, with further tests to eliminate unclear gene pairs (Supplemental Methods). For each pair of human duplicate genes, a TBLASTN pipeline was used to identify all close relatives in other genome assemblies (Supplemental Methods). Final analysis of candidate sequences was based on maximum-likelihood trees using protein sequence (Supplemental Methods). Most analysis was carried out on partial genes corresponding to the human ZF exon, and thus lacking the KRAB domain. Many identified gene candidates did not correspond to "UCSC known", RefSeq, or Ensembl predicted genes. To be sure that the orthologs identified in nonhuman genomes are bona fide candidate KZNF genes, we used a form of chained TBLASTN search for all coding exons (Supplemental Methods).

Rate of change in orthologous fingers

We measured the relative rates of change of specific amino acid residues in orthologous KZNF domains as follows. A total of 11 KZNF gene pairs were arbitrarily selected from among the duplicate pairs with an indeterminate ancestral gene. For each pair, all available orthologs were gathered from the five primates: cow, dog, and horse. The 22 genes encoded a total of 280 ZF domains. Each orthologous finger protein set was "aligned" (these are gap-free alignments since all fingers conformed to the standard 28-amino acid finger domain), and PhyML 3.0 was used to estimate rates of change at each site (JTT matrix, 20 rate categories, gamma parameter 1.0) (Guindon and Gascuel 2003; Guindon et al. 2009). The "lk" output file from PhyML gives the likelihood that each site belongs in each of the 20 estimated rate categories (Anisimova and Gascuel 2006). The peak likelihood rate value was extracted for each position in each ZF domain. These rates were averaged across all 280 orthologous finger groups. Note that this method does not measure the absolute divergence of the sequences, which varied from gene to gene depending on available orthologs. Since all 28 positions were present in all aligned finger sets, this method does produce an average estimate of the relative rates of change at each ZF site, as plotted in Figure 4.

Asymmetry calculation

For duplicate KZNF genes with an identified ancestral gene, asymmetry of divergence between the two duplicates was computed as follows. For each aligned site, the ancestral state was inferred when all copies of the ancestral gene (the single-copy gene present in early branching species) encoded the same amino acid. At such sites, when all copies of each of the two duplicate genes encoded the same amino acid and at least one gene diverged from the ancestral state (i.e., the site changed and was conserved among orthologs), the site was counted as informative. When duplicate copy 1 (the copy overall most similar to the ancestral state) was divergent, the site received a score of -1; when duplicate copy 2 was divergent, the site received a score of 0. When averaged across all informative sites, the expected score is 0 if divergence is perfectly symmetric.

Tests for positive selection

For species-specific expansion analysis, clades of species-specific tandem ZF exons were collected and analyzed by site models 7, 8, and 8A implemented in codeml PAML 3.15 (Yang et al. 2005; Zhang et al. 2005). Additional details are described in Emerson and Thomas (2009). Results and statistical tests are shown in Supplemental Table S8. Strong evidence of positive selection was detected in 30 of 35 clades. To determine the types of protein sites subject to positive selection, ZF sites with Bayes-Empirical-Bayes P-values of 0.98 or higher were counted, summing over all the clades. Counts for each of the 28 classes of ZF sites are shown in Supplemental Figure S6. For each human duplicate gene pair, branch-site models implemented in codeml were applied to test for branch-specific positive selection (Yang et al. 2005; Zhang et al. 2005). For the control model A1, the foreground d_N/d_S was constrained to be 1.0 on all branches of the tree, whereas for selection model A, the foreground d_N/d_S was allowed to differ on the branch joining the duplicate copies. Statistical analysis was based on a χ^2 test of twice the difference in log likelihoods between the two models with one degree of freedom. Specific codeml output details and statistics are shown in Supplemental Table S9.

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Coevolution of retroelements and zinc finger genes

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