

# ***De-novo* emergence and template switching of SINE retroposons during the early evolution of passerine birds**

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## Abstract

Passeriformes (“perching birds” or passerines) make up more than half of all extant bird species. Here, we resolve their deep phylogenetic relationships using presence/absence patterns of short interspersed elements (SINEs), a group of retroposons which is abundant in mammalian genomes but considered largely inactive in avian genomes. The resultant retroposon-based phylogeny provides a powerful and independent corroboration of previous indications derived from sequence-based analyses. Notably, SINE activity began in the common ancestor of Eupasserres (passerines excl. the New Zealand wrens Acanthisittidae) and ceased before the rapid diversification of oscine passerines (songbirds). Furthermore, we find evidence for very recent SINE activity within suboscine passerines, following the emergence of a SINE via acquisition of a different tRNA head as we suggest through template switching. We propose that the early evolution of passerines was unusual among birds in that it was accompanied by activity of SINEs. Their genomic and transcriptomic impact warrants further study in the light of the massive diversification of passerines.

## Introduction

Short interspersed elements (SINEs) are the most abundant group of the reverse-transcribed retroposons in mammalian genomes (Sotero-Caio, et al. in revision). They rely on *trans*-mobilization by the enzymatic machinery of long interspersed elements (LINEs) (Ohshima, et al. 1996), a *parasitic* interaction so successful that the human genome contains >1,500,000 SINEs compared to <900,000 LINEs (Lander, et al. 2001). On the other hand, SINEs are scarce in avian genomes, which has been noted as one of the most peculiar genomic features of birds (Hillier, et al. 2004; Warren, et al. 2010; Zhang, et al. 2014). While LINEs exhibit up to 700,000 copies in avian genomes, there are only 6,000-17,000 SINEs per avian genome (Zhang, et al. 2014), most of these ancient and heavily degraded (Kapusta and Suh 2016).

Presence/absence patterns of SINEs in orthologous genomic loci are rare genomic changes appreciated widely as virtually homoplasy-free phylogenetic markers (Shedlock, et al. 2004; Ray, et al. 2006). Given the aforementioned scarcity of SINEs, it is not surprising that the emergence and activity of SINEs has never been studied in birds. On the other hand, other types of retroposed elements (REs; LINEs from the chicken repeat 1 superfamily, CR1, and long terminal repeat elements, LTRs) have helped resolve the relationships of various groups of birds, such as Galliformes (Kaiser, et al. 2007; Kriegs, et al. 2007; Liu, et al. 2012), Neoaves (Suh, Paus, et al. 2011; Matzke, et al. 2012; Suh, Smeds, et al. 2015), Palaeognathae (Haddrath and Baker 2012; Baker, et al. 2014), and others (St. John, et al. 2005; Watanabe, et al. 2006; Suh, et al. 2012; Kuramoto, et al. 2015). In the meantime, the sequencing of dozens of avian genomes has revealed SINEs with putative lineage specificity (Warren, et al. 2010; Kapusta and Suh 2016; Suh, et al. 2016) and thus the potential for conducting presence/absence analyses in specific groups of birds.

Here we conduct, to our knowledge, the first study of the emergence and activity of SINEs in birds. We focus on the deep phylogenetic relationships of passerines, the largest radiation of birds with nearly 6,000 extant species (Barker, et al. 2004), using 44 presence/absence markers of SINEs and other REs. In contrast to the only previous study of retroposons in passerines with a single RE marker (Treplin and Tiedemann 2007), our multilocus dataset permits the reassessment of sequence-based phylogenies [e.g., (Barker, et al. 2004; Selvatti, et al. 2015; Moyle, et al. 2016)] and, simultaneously, the reconstruction of the temporal activity of SINEs and other REs during early passerine evolution.

## **Results and Discussion**

We initially chose RE marker candidates from selected retroposon families of zebra finch [including TguSINE1, (Warren, et al. 2010)] in October 2009, a time when genome assemblies were available only for chicken and zebra finch (Hillier, et al. 2004; Warren, et al. 2010). Candidates for presence/absence loci were therefore identified via pairwise alignment of RE-flanking sequences from zebra finch to orthologous regions in chicken (Materials and Methods). This was followed by *in-vitro* presence/absence screening of RE marker candidates as detailed elsewhere (Suh, Kriegs, et al. 2011; Suh, Paus, et al. 2011) using a representative taxon sampling of all major groups of passerines *sensu* Barker, et al. (2004) (Supplementary Table S1). We complemented this with a screening of GenBank (<http://www.ncbi.nlm.nih.gov/genbank/>) for additional SINEs, which identified a TguSINE1-like insertion in *myoglobin* intron 2 of *Pitta anerythra* (accession number DQ785977) that is absent in the orthologous position of other *Pitta* species (Irestedt, et al. 2006). We termed this element “PittSINE” and identified PittSINE marker candidates in a DNA sample of *Pitta sordida* via inter-SINE PCR [(Kaukinen and Varvio 1992); Materials and Methods]. This was followed by cloning and sequencing of the 500-bp to 1,000-bp fraction of PCR amplicons, alignment to chicken and zebra finch genomes to reconstruct the left and right SINE-flanking regions, and then *in-vitro* presence/absence screening of PittSINE marker candidates.

Next, we characterized the structural organization of passerine SINEs (Fig. 1) using the available TguSINE1 consensus sequence (Warren, et al. 2010) and after generating a majority-rule consensus of PittSINE insertions in our sequenced presence/absence markers (Supplementary Data S1). Both SINEs have highly similar, CR1-derived tails (Fig. 1) which exhibit the typical hairpin for putative binding by the CR1 reverse transcriptase and an 8-bp microsatellite at their very end for target-primed reverse transcription (Suh 2015). However, the heads of these SINEs are derived from different tRNA genes, namely tRNA<sup>Ile</sup> in

TguSINE1 and tRNA<sup>Asp</sup> in PittSINE (Fig. 1). Sequence alignment suggests that the tRNA-derived SINEs heads are more similar to the respective tRNA genes than they are to each other (Fig. 1C). However, the opposite is the case for the CR1-derived SINE tails, which exhibit four diagnostic nucleotides distinguishing them from the highly similar 3' end of CR1-X1\_Pass (Fig. 1C).

We further investigated this peculiar pattern using phylogenetic analyses of the CR1-derived SINE tails and avian CR1 subfamilies *sensu* Suh, Churakov, et al. (2015), which again suggests that TguSINE1 and PittSINE have a single SINE ancestor which derived its tail from CR1-X1\_Pass (Fig. 2A). Assuming that SINEs are *trans*-mobilized by LINE reverse transcriptase enzymes due to high sequence similarity between SINE tails and LINE 3' ends (Ohshima, et al. 1996) and thus depend on LINE activity, the most likely candidate for SINE mobilization is the CR1-X1\_Pass subfamily. This is further supported by temporal overlap of TguSINE1 and CR1-X activity in RE landscapes of the zebra finch genome (Fig. 2B). Additionally, we detected direct evidence for temporal overlap of TguSINE1 and CR1-X1\_Pass activity through our presence/absence analyses (Fig. 3A, Supplementary Table S1).

Our extensive RE presence/absence analyses yielded 19 TguSINE1 markers, 6 PittSINE markers, 13 CR1 markers, and 6 LTR markers which we could trace across a representative taxon sampling of the major groups of passerines *sensu* Barker, et al. (2004). Careful inspection of presence/absence alignments using strict criteria (see Materials and Methods) yielded a conflict-free set of RE markers, which we mapped on a maximum likelihood tree constructed from concatenated RE-flanking sequences from the same data set (Fig. 3A). For three of the deepest passerine branching events, we found a multitude of RE markers and thus statistically significant support in available RE marker tests (Waddell, et al. 2001; Kuritzin, et

al. 2016). These relationships are the respective monophyly of passerines and oscines, as well as the monophyly of Eupasserres (Mayr and Manegold 2004), a group comprising all passerines except the New Zealand wrens Acanthisittidae. The Eupasserres/Acanthisittidae split was first observed in sequence analyses of few nuclear genes (Barker, et al. 2002; Ericson, et al. 2002) and has since been recovered in ever-growing nuclear sequence analyses [e.g., (Barker, et al. 2004; Ericson, et al. 2014; Selvatti, et al. 2015; Moyle, et al. 2016)]. Our analysis of rare genomic changes thus provides the first assessment of this group using an independent marker type and phylogenetic method. None of our RE markers inserted during the rapid radiation of oscine passerines, however, sequence analysis of the RE-flanking regions yielded a topology identical to the aforementioned previous studies. Of particular interest are the four deep-branching oscine lineages Menuridae (e.g., *Menura novaehollandiae*), Climacteridae (e.g., *Climacteris picumnus*), Maluridae/Meliphagidae (e.g., *Malurus cyaneus* and *Myzomela eques*), and Pomatostomidae (e.g., *Pomatostomus superciliosus*) because these four lineages together have been rarely included in passerine phylogenetic studies. We find a branching order (Fig. 3A) which recapitulates previous phylogenetic estimates based on few nuclear genes (Barker, et al. 2004) or ultraconserved elements (Moyle, et al. 2016). This suggests that the rapid radiation of oscines can be congruently resolved even with non-genome-scale data. We note that this is in contrast to the neoavian radiation, which appears to be partially irresolvable even with retroposon markers [reviewed by Suh (2016)]. Within passerines, we further note that the conflict between single-RE support for a Picathartidae/Corvidae clade (Treplin and Tiedemann 2007) and sequence-based phylogenies (Han, et al. 2011) results from incorrect placing of this RE marker on the passerine Tree of Life (see legend of Supplementary Fig. S2 for more information).

We then traced the emergence and activity of SINEs across the passerine Tree of Life. Given

that RE marker candidates were initially chosen on chicken/zebra finch alignments, we expect no bias in the distribution of RE markers on the lineage leading to zebra finch. TguSINE1 was mostly active in the ancestor of oscines and, to a lesser extent, in the ancestor of Eupasserines. Interestingly, we find no evidence for TguSINE1 activity in the common ancestor of passerines or during/after the radiation of oscines and therefore hypothesize that TguSINE1 emerged in Eupasserines and became extinct in the oscines ancestor (Fig. 3A). The emergence of TguSINE1 is thus the first “genome morphology” character for the monophyly of Eupasserines and supplements support from skeletal morphology, which is limited to the presence of a ‘six-canal pattern’ in the hypotarsus (Manegold, et al. 2004).

In contrast to the situation in oscines, the activity of TguSINE1 appears to have been longer in suboscines, postdating the divergence between Old World and New World suboscines (i.e., pitta and phoebe in Fig. 3A). This recent, potentially lineage-specific activity coincides with the putative restriction of PittSINEs to Old World suboscines (e.g., *Pitta* spp.). The aforementioned indication for a common SINE ancestor of TguSINE1 and PittSINE evidenced by four diagnostic nucleotides in their CR1-derived SINE tails (cf. Fig. 1C and Fig. 2A) suggests that the younger PittSINE emerged from the older TguSINE1 after acquisition of a new tRNA-derived head. Assuming that TguSINE1 and PittSINE were both active on the pitta lineage, we propose that the most plausible mechanism for PittSINE emergence was template switching from TguSINE1 to a nearby tRNA during reverse transcription (Fig. 3B). Template switching has been previously proposed in a wide range of chimeric retroposons [e.g., (Brosius 1999; Gilbert and Labuda 2000; Buzdin, et al. 2002; Nishihara, et al. 2016)] and appears to be a particularly common opportunity for SINEs to *parasitize* different LINES via acquisition of new SINE tails (Ohshima and Okada 2005; Nishihara, et al. 2016). Our data show that template switching may also happen for SINE heads and we speculate that the

acquisition of a new SINE head from a different tRNA gene may provide intact and active promoter components for efficient transcription by RNA polymerase III.

## **Conclusions**

Here, we reconstructed the deep phylogenetic relationships of passerines using presence/absence patterns of unusual SINE insertions and other REs. This permitted us to follow the emergence, activity, and extinction of TguSINE1 and PittSINE across the evolution of the most species-rich group of birds. While this SINE activity was considerably lower than, for example, that in mammals, it nevertheless exemplifies that at least some birds have a more diverse repetitive element landscape than previously anticipated. Furthermore, we note that the activity of TguSINE1 appears to coincide with the evolution of vocal learning during early passerine evolution (Suh, Paus, et al. 2011). Previous evidence suggests that ~4% of birdsong-associated transcripts in the zebra finch brain contain retroposons (Warren, et al. 2010) and it thus remains to be seen whether SINE activity influenced the evolution of, for example, vocal learning in oscine passerines.

## **Materials and Methods**

We identified candidates for presence/absence loci for TguSINE1 and other selected zebra finch retroposons via pairwise alignment of RE loci from zebra finch to orthologous regions in chicken. This was done by comparing and extracting the respective RE-flanking sequences in the UCSC Genome Browser (Fujita, et al. 2011), followed by automatic alignment using MAFFT version 6 (Kato and Toh 2008). In order to find PittSINE marker candidates, we conducted inter-SINE PCR (Kaukinen and Varvio 1992) using a single, PittSINE-specific oligonucleotide primer (5'-CTCGTTAGTATAGTGGTGAGTGTC-3') and standard PCR parameters of Suh, Kriegs, et al. (2011) with 50°C annealing temperature. All

presence/absence screenings were done using oligonucleotide primers binding to conserved RE-flanking regions in chicken/zebra finch alignments (Supplementary Table S2), using the touchdown PCR and cloning protocols of Suh, Paus, et al. (2011).

For each presence/absence marker candidate, we first aligned all sequences automatically using MAFFT (E-INS-I option) and then manually inspected these for misalignments. We considered a marker candidate as phylogenetically informative and reliable “*if, in all species sharing this RE, it featured an identical orthologous genomic insertion point (target site), identical RE orientation, identical RE subtype, identical target site duplications (direct repeats, if present) and a clear absence in other species*” (Suh, Paus, et al. 2011). This led to a total of 44 high-quality RE presence/absence markers (Supplementary Table S1, Supplementary Data S2). All newly generated sequences were deposited in GenBank (accession numbers XXXXXXXXX-XXXXXXX).

All maximum likelihood sequence analyses were conducted using RAxML 8.1.11 (Stamatakis, et al. 2008) on the CIPRES Science Gateway (Miller, et al. 2010). For the CR1 phylogeny, we used the alignment from Suh et al. (2012), excluded grebe-specific CR1 elements, and added the CR1-derived tails of TguSINE1 and PittSINE. For the passerine phylogeny, we removed the RE sequences from our presence/absence alignments and concatenated the remaining RE-flanking sequences into a multilocus alignment (Supplementary Data S3).

## **Abbreviations**

CR1: chicken repeat 1. LINE: long interspersed element. Mb: million basepairs. MY: million years. MYA: million years ago. RE: retroposed element. RT: reverse transcriptase. RTBS:

reverse transcriptase binding site. SINE: short interspersed element.

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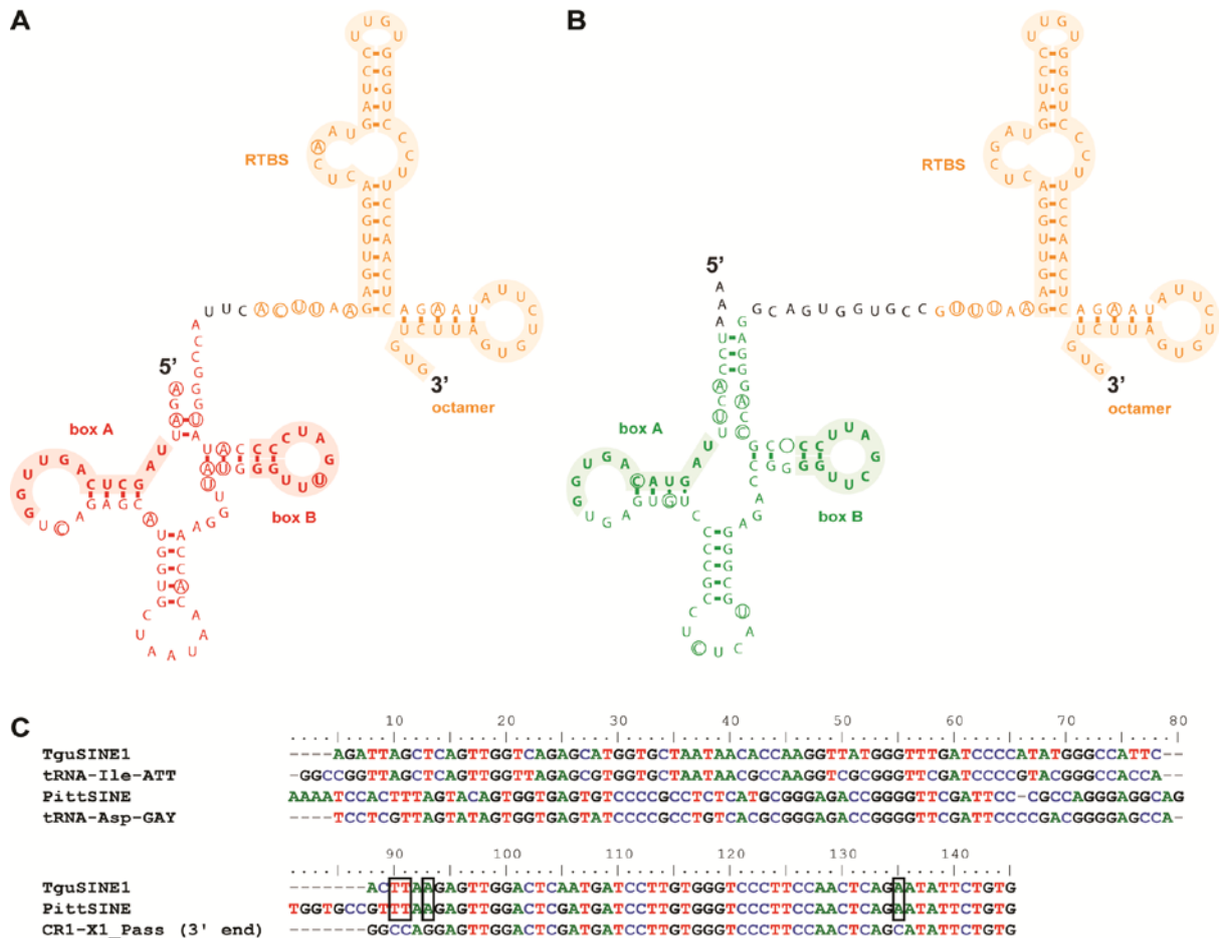
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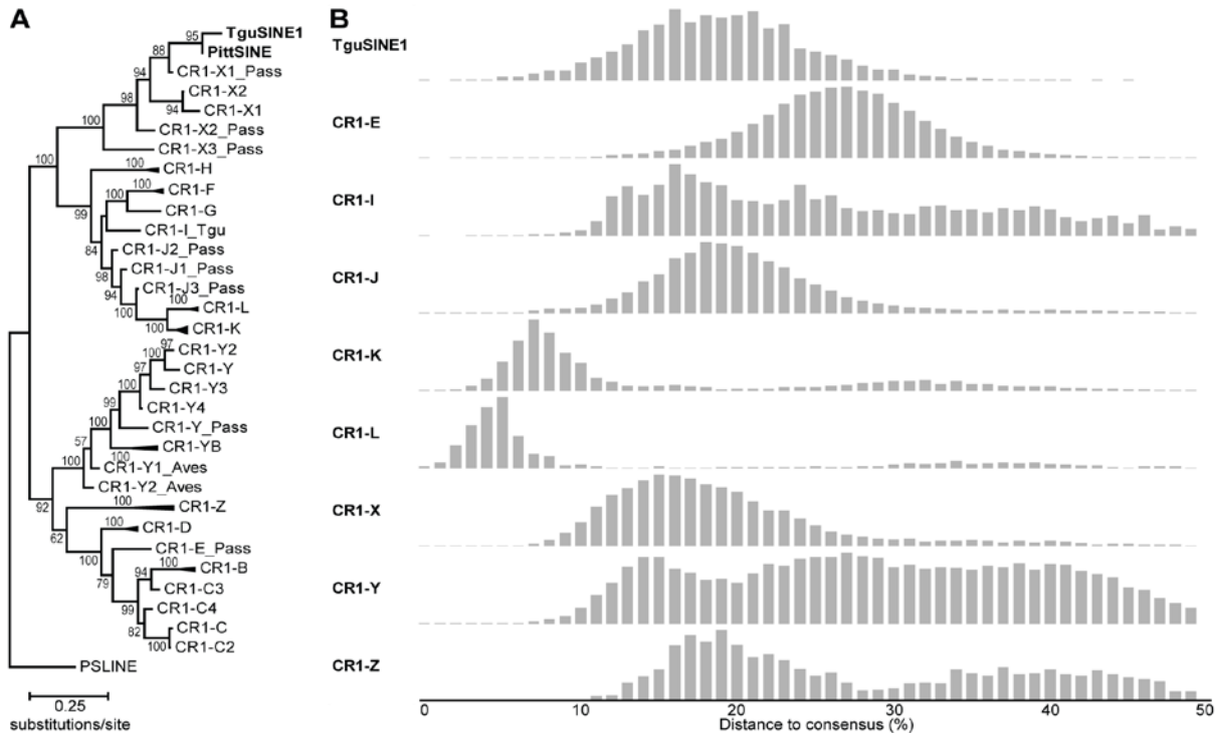
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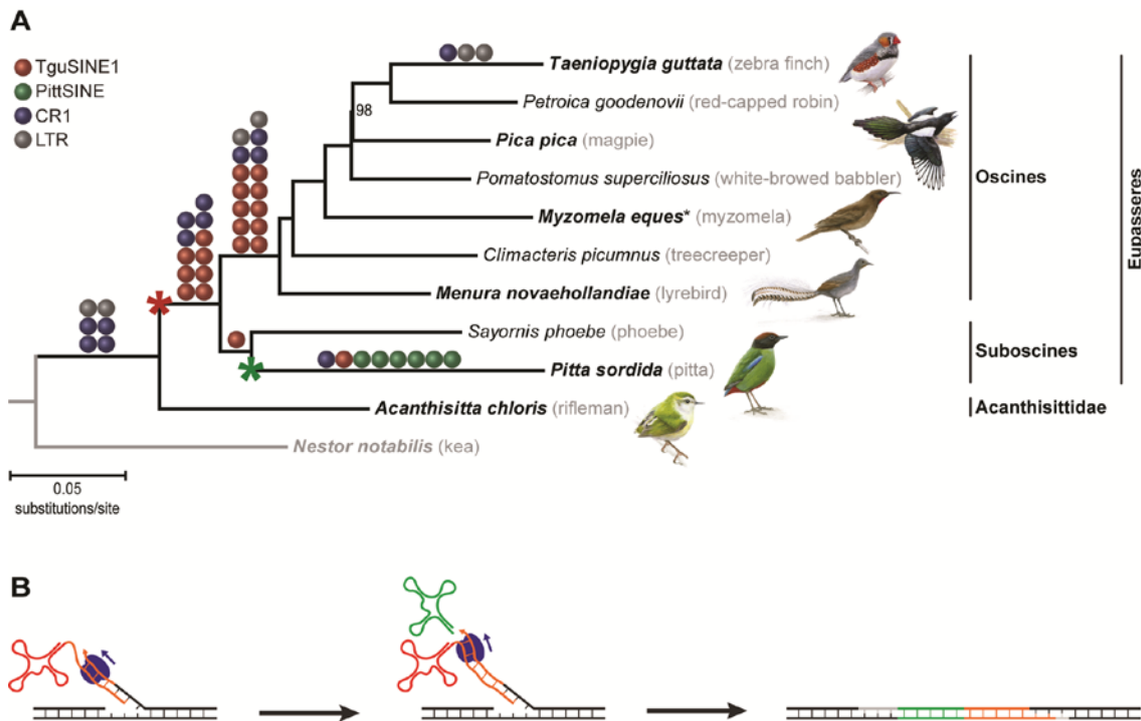
## Figure Legends



**Figure 1: Simplified RNA secondary structures of passerine SINEs with CR1-derived tails (orange) and tRNA-derived heads.** The SINE heads are tRNA<sup>Ile</sup> (red) in TguSINE1 (A) and tRNA<sup>Asp</sup> (green) in PittSINE (B). Shaded regions denote promoter boxes A and B in tRNAs, as well as the reverse transcriptase binding site (RTBS) and 5'-ATTGTRTG-3' microsatellite typical for CR1 elements of amniotes (Suh 2015). Circles indicate nucleotide differences between SINE consensus sequences and the respective tRNAs or CR1 they are derived from. The RTBS hairpin structure is also visible in mfold-based predictions of SINE secondary structure (Supplementary Fig. S1). (C) DNA alignment of TguSINE1 and PittSINE with respective tRNA genes and 3' end of CR1-X1\_Pass. Black boxes denote diagnostic nucleotides present in the CR1-derived tails of TguSINE1 and PittSINE.

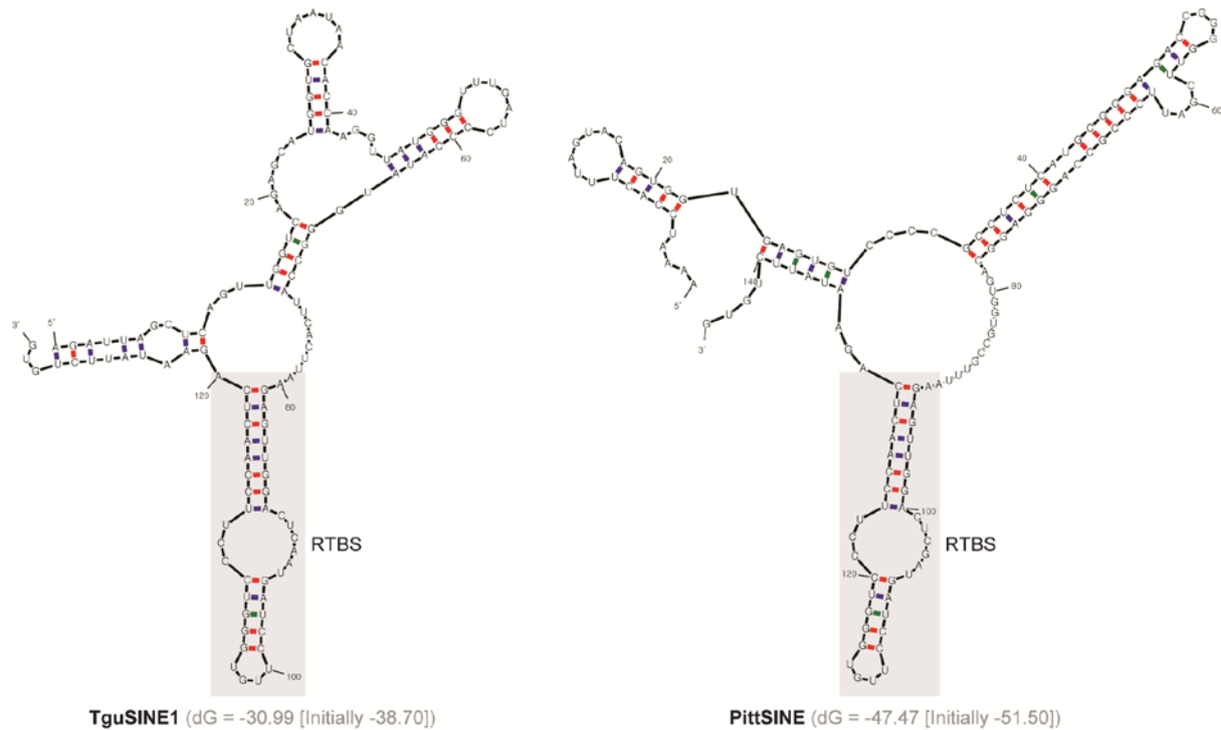


**Figure 2: Passerine SINEs share a common ancestor and are mobilized by CR1-X. (A)** Maximum likelihood phylogeny of passerine SINE tails and avian CR1 subfamilies in Repbase (Jurka, et al. 2005) (GTRCAT model, 1,000 bootstrap replicates) suggests that TguSINE1 and PittSINE arose from the same CR1-X subfamily (CR1-X1\_Pass) and share a common SINE ancestor. Note that the topology of the CR1 phylogeny is identical to that of previous studies (Suh, et al. 2012; Suh, Churakov, et al. 2015). **(B)** Comparison of the TguSINE1 landscape with landscapes of CR1 families (merged subfamilies from panel A) suggests temporal overlap of SINE and CR1-X activity in the zebra finch genome. RE landscapes were generated using the zebra finch assembly taeGut2 following methods detailed elsewhere (Suh, Churakov, et al. 2015).

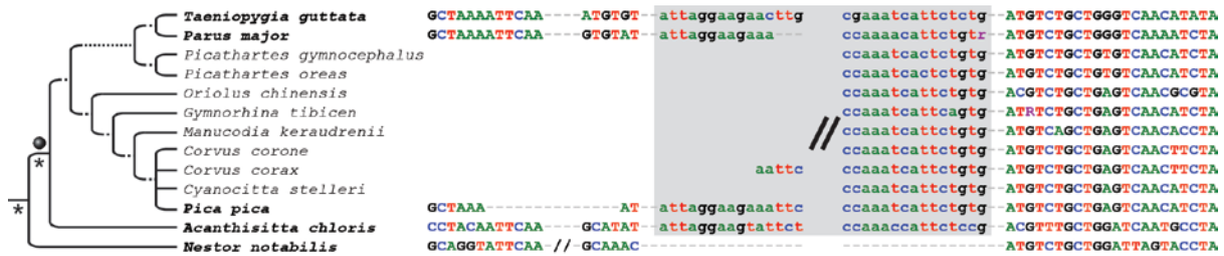


**Figure 3: Emergence and timing of CR1-mobilized SINE activity during early passerine evolution.** (A) Phylogenomic analysis of early passerine relationships using retroposon presence/absence markers (colored balls) mapped on a maximum likelihood phylogeny of concatenated retroposon-flanking sequences (GTRCAT model, 1,000 bootstrap replicates). Our sampling consists of the major deep passerine lineages *sensu* Barker et al. (2004). Red and green asterisks indicate emergence of TguSINE1 and PittSINE, respectively. The black asterisk indicates that for some loci (Supplementary Table S1), *Malurus cyaneus* was sampled instead of *Myzomela eques* to represent the Maluridae/Meliphagidae clade (Barker, et al. 2004). Only bootstrap values <100% are shown and the names of pictured birds are emphasized in bold. (B) A scenario for the emergence of PittSINE. Template switching from TguSINE1 RNA (red, tRNA<sup>Ile</sup> head; orange, CR1 tail) to tRNA<sup>Asp</sup> (green) during target-primed reverse transcription by CR1 reverse transcriptase (blue). The resultant tRNA<sup>Asp</sup>-CR1 chimaera was flanked by a target site duplication (grey) and transcriptional activation gave rise to the PittSINE family.

## Supplementary Material



**Supplementary Figure S1: Predictions of secondary structures of TguSINE1 and PittSINE using mfold (Zuker 2003).**



**Supplementary Figure S2: The RE marker of Treplin & Tiedemann (2007) does not suggest “phylogenetic affinity of rockfowls (genus *Picathartes*) to crows and ravens (*Corvidae*)”.** Our extended phylogenetic sampling instead suggests that the RE insertion occurred in the ancestor of all passerines (grey ball). This discrepancy is because Treplin & Tiedemann (2007) were, due to methodological limitations, unable to detect RE presence in non-oscine passerines (*Acanthisitta chloris*) and RE absence in the parrot outgroup (*Nestor notabilis*). Taxa with bold names were sampled in the present study and the grey box denotes the 5’ and 3’ end of the CR1 insertion.

**Supplementary Table S1: Presence/absence matrix of passerine retroposon markers including RE target site duplication (TSD) motifs and location in the taeGut2 assembly of the zebra finch genome.**

Marker	<i>Taeniopygia</i>	<i>Petroica</i>	<i>Pica</i>	<i>Pomatostomus</i>	<i>Melurus</i>	<i>Myzomela</i>	<i>Climacteris</i>	<i>Menura</i>	<i>Sayornis</i>	<i>Pitta</i>	<i>Acanthisitta</i>	<i>Nestor</i>	RE subfamily	RE orientation	TSD	taeGut2 chromosome	taeGut2 start coordinate	taeGut2 end coordinate	
L-4	+	?	?	?	+	?	?	+	+	?	+	-	TguLTR5d	-	GTAAG	2	107,658,311	107,659,269	
L-4	+	?	?	?	+	?	?	+	-	?	-	d	TguSINE1	+	?	2	107,658,311	107,659,269	
Pso01	d	?	?	?	?	?	?	d	-	+	-	?	PittSINE	-	C	4	43,915,733	43,916,189	
Pso02	-	?	?	?	?	?	?	-	-	+	?	?	PittSINE	+	CTATG	5	27,322,271	27,322,008	
Pso03	+	?	?	?	?	?	?	+	+	+	-	?	CR1-X1_Pass	+	TTTCT	10	11,861,921	11,861,386	
Pso03	-	?	?	?	?	?	?	-	-	+	-	?	PittSINE	-	GAAAT	10	11,861,921	11,861,386	
Pso04	-	?	?	?	?	?	?	-	-	+	?	?	PittSINE	+	GGA	3	38,268,171	38,270,277	
Pso08	-	?	?	?	?	?	?	?	-	+	-	?	PittSINE	-	CACA	10	1,122,922	1,122,063	
Pso10	-	?	?	?	?	?	?	-	+	+	-	?	TguSINE1	+	TG	5	10,716,893	10,716,277	
Pso12	-	?	?	?	?	?	?	-	-	+	?	?	CR1-L_Tgu	-	?	7	30,071,845	30,072,221	
Pso13	-	?	?	?	?	?	?	?	-	+	?	?	TguSINE1	+	?	5	10,715,853	10,716,265	
Tgu02	+	?	+	?	+	?	+	+	+	-	-	-	TguSINE1	+	?	1	10,356,482	10,355,761	
Tgu05	+	?	+	?	+	?	+	+	+	?	-	-	TguSINE1	+	A	1	100,547,887	100,547,056	
Tgu07	+	?	+	?	?	+	?	+	-	-	-	-	TguSINE1	+	A	1	118,418,987	118,418,291	
Tgu07	d	?	-	?	?	-	?	-	-	+	-	-	PittSINE	+	CGATAGTG	1	118,418,987	118,418,291	
Tgu08	+	+	?	?	+	?	+	+	+	?	-	-	TguSINE1	+	ATTATT	2	4,459,063	4,459,958	
Tgu09	+	?	+	?	?	?	+	+	?	-	-	-	TguSINE1	+	AA	2	19,559,154	19,560,021	
Tgu10	+	?	+	?	?	?	+	?	+	?	-	-	TguSINE1	+	?	2	39,949,817	39,949,452	
Tgu13	+	?	+	?	?	?	?	+	+	?	-	?	TguSINE1	+	?	2	66,988,522	66,989,422	
Tgu14	+	?	+	+	?	+	+	+	+	+	-	-	TguSINE1	+	TGAGGAG	2	75,161,831	75,162,667	
Tgu17	+	?	+	?	?	+	?	+	-	?	-	-	CR1-L_Tgu	+	AC	1	20,072,950	20,071,904	
Tgu19	+	?	+	?	?	+	+	+	+	?	-	-	CR1-L_Tgu	+	GAATT	2	63,475,805	63,476,911	
Tgu20	+	?	+	?	?	+	?	+	-	?	d	-	CR1-L_Tgu	+	?	2	137,203,952	137,204,879	
Tgu21	+	?	?	?	+	?	?	+	+	?	+	-	CR1-L_Tgu	+	AAC	3	87,434,090	87,435,182	
Tgu24	+	?	+	?	?	+	+	+	+	?	?	-	TguLTR5c	+	AAGAG	4	38,637,563	38,638,584	
Tgu26	+	?	+	?	?	?	+	+	?	-	-	?	TguLTR5c	+	AATTA	6	17,528,118	17,526,787	
Tgu27	+	?	+	?	?	?	+	+	-	?	-	-	TguLTR5c	+	TGCTC	1A	32,328,895	32,327,600	
Tgu32	+	-	-	?	-	?	-	-	-	?	-	-	CR1-J3_Pass	+	?	2	19,478,341	19,479,587	
Tgu33	+	?	+	?	?	?	+	+	+	?	-	?	CR1-J3_Pass	+	GA	3	24,954,034	24,952,882	
Tgu34	+	?	+	?	+	?	+	+	-	-	-	-	CR1-J3_Pass	+	T	3	32,857,318	32,856,592	
Tgu37	+	-	-	?	?	?	d	?	-	-	-	-	TguERVk9_LTR2h	+	CTAGGT	9	6,881,758	6,882,600	
Tgu39	+	?	+	?	?	?	?	?	+	?	-	-	TguSINE1	+	TA	2	121,403,718	121,402,809	
Tgu42	+	?	+	?	?	+	+	+	-	-	-	-	TguSINE1	+	?	3	11,125,489	11,126,181	
Tgu43	+	?	+	?	?	?	+	+	-	-	-	-	TguSINE1	+	AGGATATTAA	3	98,795,454	98,794,809	
Tgu47	+	+	+	?	?	+	+	+	+	+	-	-	TguSINE1	+	GAATTT	5	24,046,603	24,045,599	
Tgu48	+	?	+	?	+	?	+	+	+	+	+	-	CR1-X1_Pass	+	TATTTA	5	35,475,748	35,475,476	
Tgu51	+	?	+	?	?	?	+	+	+	?	-	?	TguSINE1	+	ATTCAG	6	30,476,496	30,477,321	
Tgu52	+	?	+	?	?	+	+	+	+	?	-	?	TguSINE1	+	AT	6	33,164,987	33,165,899	
Tgu58	+	?	+	?	?	?	+	?	?	-	?	?	TguSINE1	+	CTT	Z	39,585,532	39,586,342	
Tgu60	+	?	+	?	?	?	+	+	+	-	?	-	d	TguSINE1	+	ACAT	13	15,018,269	15,019,192
Tgu61	+	?	+	?	?	?	+	?	+	-	-	-	CR1-X1_Pass	+	?	15	2,160,370	2,160,694	
Tgu62	+	-	?	?	?	?	?	-	-	-	-	-	TguLTR11n	+	TAACC	11	15,647,516	15,648,610	
Tgu65	+	?	?	?	?	?	+	?	?	+	+	-	CR1-X1_Pass	+	AAAT	1	1,206,381	1,206,988	
Tgu72	+	?	?	?	?	?	+	+	?	?	+	-	CR1-X1_Pass	+	?	2	6,160,531	6,161,492	

The character states are '+' (RE presence), '-' (RE absence), or 'd' (unspecific deletion) for each genomic locus. Missing data is indicated by '?'. The TguLTR5d insertion of marker L-4 was first described in Suh, Paus, et al. (2011)

**Supplementary Table S2: Oligonucleotide primer sequences.**

Marker	forward primer (5'-3')	reverse primer (5'-3')
L-4	CAGTTGTGTGAGCTTTCCTTG	CAGAAGAAAAATTGTATGATAATGG
	GCTTTCCTTGTGATATGGTTG	CTGTTTCTTGCTTATTGTGTTGG
Pso01	AAACCAGAGTATCCCCTTGC	CATTAGAAATCAGCTCAAACAGTG
	AGAGTGCCACAGATGCCTG	CTCATTTTTCCATTTCCAGC
Pso02	ACTTGGGATGAGAGGAGAGC	ATAAATTAGGCAGGCAAAGC
	GGATGAGAGGAGAGCTTGC	GCTATTTGTACGAAGATATGTAATGC
Pso03	GTCACAACGCAGTTCTGATG	CCTTAGAAATGGAGAAGCCTG
	GGGTATTTACTACTTTTGTGTC	GCTGTGTGTAGGATCTGTTGC
Pso04	CTCTCTGAAGTGGACTGTGTCTC	AGCTGAGGGCACATAGTAGC
	TCTGAAGTGGACTGTGTCTCATC	TGAGGGCACATAGTAGCTGC
Pso08	ACTCTGTCCTGACCTTATCCAG	CAGAACAAAGCCAAGCAATG
Pso10	TGCTGGCTTTGAATGACAG	TGCCACATTTCTTAGTGATGG
	GGCTTTGAATGACAGCTACTG	TGACCCCATATCAGGAGC
Pso12	GAAGAAACCACAGCTTCAAGG	GTCATGTTCCAGACAGGCAGG
	AGAAACCACAGCTTCAAGGG	ATGTTCCAGACAGGCAGGATG
Pso13	CACGAGTTTTAGAGGCATAACC	GCGAAGTATTTACCATCACTAAG
	CAGACTGTTCCACCAAGCAG	
	AGCTGAGAAAGCACCAAAGTC	GAAATGTAGCATCCAGTGAGAAG
Tgu02	TGTTCAACATTCATTCC TTGG	AAATCTCAGAGGAGCTTCGTAC
	GCCAGTAAGCTCAGTTTTTCAG	CCATAGGAATATCAAAGAACCTG
Tgu05	GCAGGAAATGCCTTTCTG	CACTGTTCTCCATCATCTTCAG
	GAAATGCCTTTCTGTTTTG	CTTTCTAAACTTGGGTTCTATGC
Tgu07	AAGGAGAGGTCAGTGAATTGG	TCATCAGGGATCTGACTTGC
	ACAAAGTATTTGAGAGCCAGC	ATGACCCAGCCCTGTCAG
Tgu08	AACCAACAACCTCTAAAAGCCAC	GCCCAAAGATGACATCGTG
	GGGAAGGAGGAATCTGAATAC	TGGACAGAGTTAAGATGGAAGAC
Tgu09	CAATTAGTGAGTTATAGCCACACAC	CATGACCAACAGTAACCATCAG
	ACACTGCAATACATGTGATAAGTC	TCAGACTTCTAACAGCAAGAGC
Tgu10	TCATATCAATTCAAGGCAAGG	ATCACAAGGCTGGATGAGC
	AGGTTCCCTGTTCTGTTACTG	GGATGAGCTTGCTGTTGC
Tgu13	GCCACCACAACATCCCAC	GGAGAGGGTGACTATTGTCTGTG
	GCAGAAGTGAGATTTTAAGC	GGGTGACTATTGCTGTGAGG
Tgu14	TTCGTAGACTTCACAGAAATCAC	CCTTTCAGTCTGCTACCAG
	CCTGCCTATTTAGTGTAGTTTCAG	CTGCTACCAGTGCAAAGC
Tgu17	ATCTGCCAAGACAGCAAGTC	CCTGATCCAATTCAATCCAG
	AGACCAAGTAGAAAGATTCCTC	CACACACAAAGACTAAACAACCTGG
Tgu19	AACCTCGACTTCCAGATGG	GAAGAAATAACGCTCACATCAG
	TACAAAAGGAAGAGCGATGG	CAGGTCTCTTCAGATTGATG

**Supplementary Table S2: Continued.**

<b>Tgu20</b>	TTGCTTGGCACCAGACAC GCCTGACATTATCTTTCCCTG	AAACAGGTGGTCTCTGAGC CAAGGCAATGCTGATGTTT
<b>Tgu21</b>	AGGATCAGTGGAAATTGGTTC GCCCTGTGACAGCACTAATG	CTTTTACAGAAGCCATAAGTGC CACAGTTCAGACCTTCAAATCC
<b>Tgu24</b>	AGGAGTAATAGCCTGTGGAAGA GAGTAATAGCCTGTGGAAGAGAG	CTGGTATTGAGACAGACAGTGG ACAGACAGTGGATCAGAAGAAAC
<b>Tgu26</b>	GGAGTAGCAATGTACCCATAGAC CATAGACATCAATAAAGGAAGAGG	TGAAGTATCAAGTCTGCCTC AGCCCTGTCTTCTAAATCTCAG
<b>Tgu27</b>	GTGGAAAATCATTGAGGACTG AGCCAGTAGGGAGCATCAG	GCAGTTCAAGGGAATAGTAACAG AGTTCAAGGGAATAGTAACAGAAG
<b>Tgu32</b>	GGGAATTTGATCCAGCTATTG	GATACTTACCCCATGCTTTACTG
<b>Tgu33</b>	CCTTACCATCTCCTAATGACCTC	CCATCATCTTTTTTCAGAGCATC
<b>Tgu34</b>	ATAGGTGGTGTATGATGATTGG CCTGGAGCTGGCAAGACTAC	TTCCAGTGTCTGCCATCTG ATCTGACGCTTAAAACCTGC
<b>Tgu37</b>	ATTGAGGGAGTCAAAGGAGC GTCAAAGGAGCCAAGAACTTC	CATCTCAGTACATCTGCTGTGC CAGAAAGGGTGTCTTGTGTCAG
<b>Tgu39</b>	CTTCTGTTCTCAAACCCAC TTCTCAAACCCACCAGAC	CTTGCCTATTCTTCTTTCAG CTTCTTTCAGGTGTTTTGTC
<b>Tgu42</b>	AAATTAAGCCTCTAGTCTGTTCTG ATTAAGCCTCTAGTCTGTTCTGG	CCTCCCTCAGGTTTCCAG CCTTCAGGTTTCCAGTGAAC
<b>Tgu43</b>	GGAAGCCATCACAACACTTG	GTTTATCAGACATACGTCAATGC
<b>Tgu47</b>	CTGCTTGCTTCCCTCTCTG CTTCTTCCCTCTCTGTTTC	ACCTCTGTTTCTACATCAGTGAAG CCCAGCTAGACAGGAAAGATG
<b>Tgu48</b>	TTGAGATGTAGAGCTTGACACTAAC	AGAGATTCTGCCTCTATTCTTTG
<b>Tgu51</b>	ATTGCGAGTCCAGATTTCC CGCAGTCCAGATTTCTG	TTGACAGGAGGGATAACAACC ATCAAACAAGATCCTGTGAGC
<b>Tgu52</b>	TCACCACAAAGCAGATGATG CACCACAAAGCAGATGATGC	GGCAGAGAAATCCTTCTG AGGTTGGATATCAGGAAAAGG
<b>Tgu58</b>	TCTCTCTGCTGCTGGTTC TCCTCTGCTGCTGGTTCTAG	GCCCAGATTGCTAATGAAAG CCCAGGGTAAATACAGATGC
<b>Tgu60</b>	CCAGGCAGAAGGTGAACTG CAGTGCAGGAAGTTGGAGTG	GTCTTGGACTTTTGTGTTGACC TGACCAGATTTGCTCCCTG
<b>Tgu61</b>	ATCAAGAAGCTCAGCACGAC ACGACCAACAAAAACGAGAC	GGGATTGACAGCCTCAGAG GGATTGACAGCCTCAGAGC
<b>Tgu62</b>	CTGGGAGTGTTAATGGTGAGAG GAGTGTTAATGGTGAGAGTGTGC	GATTTATAGTGCCCTTCCATAAGC ATAAGCGTAAAGGTGTTCTGC
<b>Tgu65</b>	GGAGCAGTCACAGTCTTCAGG	TGTGTGTTTTCACAGTCCATAGC
<b>Tgu72</b>	ATATTTGCTACTAGGTTGCTACG	ATTTACCCATTTGCCTGAAG

**Supplementary Data S1: Majority-rule consensus sequence for PittSINE as reconstructed from our PittSINE-bearing presence/absence patterns.**

**Supplementary Data S2: Fasta-formatted alignments of all RE presence/absence markers.**

**Supplementary Data S3: Fasta-formatted multilocus alignment of concatenated RE-flanking sequences used for generating the phylogenetic tree of Fig. 3A.**