Evolution of Two Short Interspersed Elements in *Callorhinchus milii* (Chondrichthyes, Holocephali) and Related Elements in Sharks and the Coelacanth

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**Abstract**

Short interspersed elements (SINEs) are non-autonomous retrotransposons. Although they usually show fast evolutionary rates, in some instances highly conserved domains (HCDs) have been observed in elements with otherwise divergent sequences and from distantly related species. Here, we document the life history of two HCD-SINE families in the elephant shark *Callorhinchus milii*, one specific to the holocephalan lineage (CmiSINEs) and another one (SacSINE1-CM) with homologous elements in sharks and the coelacanth (SacSINE1s, LmeSINE1s). The analyses of their relationships indicated that these elements share the same 3′-tail, which would have allowed both elements to rise to high copy number by exploiting the *C. milii* L2-2_CM long interspersed element (LINE) enzymes. Molecular clock analysis on SINE activity in *C. milii* genome evidenced two replication bursts occurring right after two major events in the holocephalan evolution: the end-Permian mass extinction and the radiation of modern Holocephali. Accordingly, the same analysis on the coelacanth homologous elements, LmeSINE1, identified a replication wave close to the split age of the two extant *Latimeria* species. The genomic distribution of the studied SINEs pointed out contrasting results: some elements were preferentially sorted out from gene regions, but accumulated in flanking regions, while others appear more conserved within genes. Moreover, data from the *C. milii* transcriptome suggest that these SINEs could be involved in miRNA biogenesis and may be targets for miRNA-based regulation.

**Key words:** CORE domain, Deu domain, miRNA, SINEs, SINE highly conserved domain, SINE replication waves.

**Introduction**

Transposable elements (TEs) are DNA sequences that multiply either by producing copies that will insert in different genomic locations or by simply repositioning themselves elsewhere in the genome. TEs can be grouped into two major classes: Class I includes all elements replicating via an RNA intermediate (retrotransposons), and Class II comprises elements that move via DNA intermediates (transposons; Wicker et al. 2007). In both classes, autonomous and non-autonomous elements occur. The autonomous elements encode all the enzymes that are required for their retrotransposition, while non-autonomous elements are basically molecular parasites. In fact, for their mobilization, they exploit the enzymatic machinery encoded by autonomous TEs (Ohshima and Okada 2005; Yang et al. 2009).

The best-known representatives of such quasi-parasitic relationships are the non-autonomous short interspersed elements (SINEs) and their partners, the long interspersed elements (LINEs; Ohshima and Okada 2005). LINEs are ubiquitous Class I elements which code for a polyprotein with a reverse transcriptase (RTase) domain, responsible for the reverse transcription of the RNA intermediates and the integration of cDNA into chromosomal locations. SINEs are small elements with a well-defined modular structure: a small RNA-derived head, an anonymous body (i.e., a sequence without any relationship with known protein-coding domains) and a tail terminating with an oligo-(A) or a microsatellite motif (Kramerov and Vassetzky 2011). In SINEs, the functionally relevant modules are the head and the tail. The former contains the promoters for the RNA polymerase III (pol-III) and is thus responsible for the transcription of the SINE-derived RNA that will constitute the template for the reverse transcription. The tail module, on the other hand, is homologous to the tail of the partner LINE. It is this region of the SINE RNA that binds the LINE-encoded protein (Ohshima and Okada 2005). The replication rate and survival of a SINE is
therefore strictly dependent on the partner LINE activity, so that if the latter undergoes silencing, the former will also stop replicating. A way for SINEs to escape silencing is to change the mobilizer LINE; this can be achieved by switching the tail module through recombination with a different LINE or with another SINE carrying a tail matching that of an active LINE.

It is well-known that distinct SINE families have exchanged functional modules (Takahashi and Okada 2002; Dergon and Zhang 2006). This led to the development of the “module exchange” model which predicts that recombination between different SINEs and/or other DNA elements may promote the switch of functional modules of the element, assuring the retrotransposition competency and, eventually, leading to SINES diversity (Kramerov and Vassetzky 2011; Luchetti and Mantovani 2013).

As mentioned above, the body sequence of a SINE is apparently not functional for retrotransposition. Nonetheless, some SINE families carry highly conserved domains (HCDs) within the body sequence; this has been observed for widely different SINE families, isolated from a variety of host species (Gilbert and Labuda 1999; Og iwara et al. 2002; Nishihara et al. 2006; Piskurek and Jackson 2011; Luchetti and Mantovani 2013; Matetovici et al. 2016; Nishihara et al. 2016; Luchetti and Mantovani 2016). So far, the discovery of SINE domains conserved and vertically inherited for long periods (up to 800 Ma; Luchetti and Mantovani 2016) remains to be explained. Three models have been proposed. Gilbert and Labuda (1999) suggested that HCDs may help in maintaining the interaction of a SINE with the partner LINE. Alternatively, HCDs may serve as recombination hot spots allowing the generation of new SINES, in line with the module exchange model (Gilbert and Labuda 1999; Luchetti and Mantovani 2013). A third option points to a possible functional role of HCDs conferring some selective advantage to the host genome (Nishihara et al. 2006; Dergon 2012).

SINE insertions may have a deep impact on the host genome by inducing structural genomic variation and/or modifying gene expression profiles (reviewed in Schmitz 2012). Remarkably, some SINE insertions appear to have triggered major evolutionary changes through exaptation, being included as functional components into the host genome (reviewed in Dergon 2012). In general, TE activity is considered one of the major facilitators of genome evolution, suggesting a link with species diversity and evolution as well (TE-Thrust hypothesis; Oliver and Greene 2011).

In a wide attempt to understand the evolution of HCD-SINES, we collected several elements (either previously isolated or newly characterized) looking for possible internal homologies with known conserved domains. We here present the results for the elephant shark Callorhinchus milii genome (Venkatesh et al. 2014), which shows a SINE family with the CORE HCD. Moreover, we analyzed the relationships with coexisting SINES and the potential impact of detected insertions on the host genome evolution. The data helped to depict the life history of two SINE families in the C. milii genome and contribute to the knowledge on internal domain conservation.

Materials and Methods

Elephant shark’s SiNE2-1_CM and SiNE2-1B_CM element consensus sequences, found in the Callorhinchus milii genome assembly 6.1.3 (Venkatesh et al. 2014), were downloaded from Repbase (Bao et al. 2015). Presence/absence of SINE copies in the other genomes has been assayed by BLAST (Altschul et al. 1990), selecting hits with an e-value ≤ 1e−10.

Genome-wide detection of SINE insertions was performed using RepeatMasker v. 4.0 (Smit et al. 2013–2015). We excluded from the analysis all those insertions that could not be unequivocally attributed to a single SINE family, due to the close sequence similarity. These amounted to ~5% of total scored insertions, except the LmeSiNE1c family, for which 41% of scored insertions could not be unequivocally attributed.

The age analysis of SINE insertions was calculated using two approaches. First, transposition in transposition (TinT; Churakov et al. 2010) analysis was used to evaluate the relative age of activity within genomes. For this analysis, a RepeatMasker outfile was generated using a collection of TEs downloaded from the RepBase databank (accessed on December 2016), which included elements both specific to C. milii and ancestral ones (i.e., shared with other taxa). We, then, added to this library the consensus sequence of the newly found SacSiNE1-CM. A second approach was based on sequence divergences. The age of SINE copies was estimated from their divergence from the relative consensus sequence using the Jukes-Cantor substitution model, which accounts for multiple nucleotide substitutions.

The SINE genomic distribution was tested in C. milii and in L. chalumnae, the only species, among presently examined ones, with well-annotated genomes (obtained from NCBI database, accessed on March 2016; Amemiya et al. 2013; Venkatesh et al. 2014). The overlap between SINE insertions and annotated genes was evaluated with BEDTools v. 2.17 (Quinlan and Hall 2010) using the intersectBed function. To assess the significance of observed SINE distribution we performed a simulation. The genomic locations of SINE insertions were randomly permuted 10,000 times using the BEDTools shuffleBed function (excluding genome gaps); their genomic distribution was then determined with intersectBed and considered as a null distribution against which observed data were compared. Finally, we calculated an empirical P value of obtaining an equal or greater number of overlaps in observed data.

Results

CmSiSINE Sequence Characterization

SiNE2-1_CM and SiNE2B-1_CM heads have a clear tRNA origin, being very similar to C. milii Alanine tRNA
(similarity = 81.9–90.3%, respectively); they can also be folded into the typical tRNA clover-leaf secondary structure and retain conserved RNA pol-III promoters, the A and B boxes (fig. 1). The two SINEs differ for the 5.1%; due to their close similarity, and for reasons of clarity, they will be collectively referred to as CmiSINEs.

We then checked whether the body aligns to one of the known HCDs (Vassetzky and Kramerov 2013). It emerged that CmiSINEs belong to the CORE-SINE superfamily, having a body 66.78–68.5% similar to 58 out of 65 bp of the CORE consensus sequence (fig. 1).

As SINEs exploit the partner LINE enzymatic machinery by sharing homology at the 3' end of the tail, we also searched for known C. milii LINEs whose tail matches those of CmiSINEs. The analysis of RepBase non-LTRs collection evidenced a 78.6% similarity with the tail of the L2-2_CM LINE (fig. 1; supplementary fig. S1, Supplementary Material online).

![Fig. 1.—CmiSINEs sequence characterization. (A) In the head module, A and B boxes of the RNA pol-III promoter are indicated by black lines. Above SINE sequences, the C. milii Alanine tRNA, the CORE HCD consensus sequence (taken from Vassetzky and Kramerov 2013) and the L2-2_CM tail are aligned. Identical residues are shaded in grey; the “//” symbol indicates that part of the sequence is omitted as non-homologous. The region shared with the SacSINE1-CM family is boxed. (B) The cloverleaf secondary structure of the SINE2-1_CM tRNA-like head; nucleotide substitutions observed in SINE2B-1_CM are reported circled. The anticodon is shaded in grey.](image-url)
Search of CmiSINEs in Related Genomes and Relationship with Other Elements

We searched CmiSINEs in other cartilaginous fishes: Squalus acanthias (whole genome shotgun clone traces archive, N = 12,873; EST library, N = 32,562), Rhincodon typus (assembly Rhincodon v1; Read et al. 2015) and Leucoraja erinacea (assembly LER_WGS_1; King et al. 2011). Representatives of bony fishes were also checked: Danio rerio (assembly GRCz10; Howe et al. 2013), Takifugu rubripes (assembly FUGU5; Aparicio et al. 2002) and Latimeria chalumnae (assembly LatCha1; Amemiya et al. 2013). We also searched in the cyclostome Petromyzon marinus (assembly 7.0; Smith et al. 2013).

These searches evidenced the absence of CmiSINES in the assayed genomes; however, the 3’ end of CmiSINEs aligned significantly with clones from Squalus acanthias (~140 bp, similarity = 77.4–75.5%). Alignment of positive clone fragments obtained from S. acanthias indicated that they were copies of another SINE family with a tRNA-related head: when searched in Repbase, it matched with a known element carrying the Deu HCD and isolated in S. acanthias: SacSINE1 (Nishihara et al. 2006). The latter element was, then, searched and found in the C. milii genome (henceforth, it will be called SacSINE1-CM), with copies showing 74.1% average identity with the SacSINE1 consensus sequence. The comparison between SacSINE1-CM and CmiSINEs confirmed 94.3–95.7% similarity only at their 3’ end (~160 bp), including the region homologous to the 3’ end of L2-2_CM LINE (fig. 2A; supplementary fig. S1, Supplementary Material online).

We found SacSINE1 also in the R. typus genome, where detected copies showed lengths ranging from 97 to 463 bp and a similarity score to the SacSINE1 consensus of 75.4%. The L. erinacea genome also harbors SacSINE1-like elements, with length spanning from 185 to 425 bp and an average identity to the SacSINE1 consensus of 73.7%.

Apart from T. rubripes, SacSINE1 showed significant matches with cyclostome and the other bony fish genomes (table 1). The similarity with P. marinus occurred in a region homologous to the SacSINE1 Deu domain but, apparently, it does not belong to any known repeat, nor it was possible to isolate a new one. Cyclostomes harbor Deu-SINEs (Nishihara et al. 2006), therefore, it is possible that these similarities involved fragments of an extinct SINE lineage. In D. rerio positive hits are located within the Deu HCD, that is part of the SS rDNA-derived element SINE3-1 (Kapitonov and Jurka 2003; Nishihara et al. 2006). Hits on the L. chalumnae genome, though, span from the 5’ end to the Deu domain, thus including the tRNA-like head and the 5’ half of the body. These hits correspond to the coelacanth LmeSINE1 family (Nishihara et al. 2006), comprising five different lineages (LmeSINE1a-e). The alignment of consensus sequences revealed that LmeSINE1s, SacSINE1, and SacSINE1-CM share similarity across the tRNA-like head, the Deu domain and the tail. Only LmeSINE1a and LmeSINE1d lineages showed a large deletion in the body as a main structural difference (fig. 2A; supplementary fig. S2, Supplementary Material online).

Timing of the SINE Activity

The age analysis performed with the TinT method clearly indicated that CmiSINES experienced replication waves more recently than SacSINE1-CM, with an almost contemporary activity of SINE2-1_CM and SINE2B-1_CM (fig. 3). The inferred activity time of the LINE L2-2 overlaps to that of analyzed SINES.

The same analysis was, then, performed to check the relative time of activity of LmeSINE1 elements, evidencing that replication waves of the five lineages occurred with a slightly different timing (fig. 3), LmeSINE1a and LmeSINE1d being mobilized more recently.

We also performed an age analysis by comparing the extent of similarity of each copy to the relative consensus sequence, based on the principle that more recently produced copies show fewer substitutions than those that were produced more distantly in the past. CmiSINES and SacSINE1-CM consensus sequences were therefore searched in the C. milii genome. We found 302,861 CmiSINE insertions (175,998 for SINE2-1_CM and 126,863 for SINE2B-1_CM). Copies divergence ranged from 0.0% to 42.4% (average = 12.6%) and from 0.0% to 47.7% (average = 17.2%), respectively. Moreover, we found 218,294 SacSINE1-CM insertions, with copies diverging from the consensus by 0.0–49.2% (average = 22.4%). The histogram plot clearly suggests a more recent replication wave of CmiSINES and an older activity of SacSINE1-CM (fig. 4), in full agreement with the TinT analysis.

We, then, attempted to put estimated SINES activity in an absolute timeframe. Considering the presence of CmiSINES in the holocephalan lineage and its absence in the elasmobranch one, we used the split age of these two taxa (421 Ma; Inoue et al. 2010) as the maximum age for CmiSINES. Therefore, the oldest CmiSINE copies, that are the most divergent from the consensus (42.4–47.7%), were considered as being produced 421 Ma and those completely identical to the consensus as being very recently produced elements (0 Ma; fig. 4).

Assuming a strict molecular clock, the substitution rate experienced by the oldest CmiSINE copies ranges between $1.0 \times 10^{-3}$ and $1.1 \times 10^{-3}$ substitutions/site per million year. Based on these estimates, and assuming that each SINE copy likely accumulated substitutions at the same rate, the maximum activity seems to have occurred ~105–118.2 Ma for SINE2-1_CM, and ~144.7–162.8 Ma for SINE2B-1_CM. SacSINE1-CM, on the other hand, appears to have originated about 434.2–488.5 Ma, with a peak in activity around 201.2–226.4 Ma.

Using the elasmobranch substitution rate calculated on nuclear gene synonymous substitutions, $1.6 \times 10^{-4}$ substitutions/site per million year (Martin 1999), we obtained widely different ranges of activity: CmiSINES reached their
peak of activity around 787 and 1,075 Ma, while SacSINE1-CM would have reached it about 1,400 Ma.

We then applied the newly calculated substitution rate to coelacanth LmeSINE1s in order to date their activity. The search for insertions of the five LmeSINE1 lineages in the \( L. \) chalumnae genome resulted in 73,075 (LmeSINE1a), 32,791 (LmeSINE1b), 500 (LmeSINE1c), 70,400 (LmeSINE1d), and 35,616 (LmeSINE1e) hits. Overall,
divergence values ranged from 0.0% to 54.2%. Again, the divergence plot fully agrees with TinT relative time analysis (fig. 3). The age analysis indicated that the replication of the oldest lineage (LmeSINE1e) started at 478.4–538.2 Ma and two main replication waves, corresponding to LmeSINE1d and LmeSINE1a activity, occurred 122.8–138.1 Ma and 57.1–64.2 Ma (fig. 4).

SINEs Genomic Distribution

We checked the distribution of retrieved CmiSINEs and SacSINE1-CM copies within the *C. milii* genome and of LmeSINE1 insertions within the *L. chalumnae* genome, to verify whether insertions are differentially distributed with respect to coding regions. The analysis showed that the two CmiSINEs have about 76.7% (SINE2-1_CM) and 78.1% (SINE2B-1_CM) of insertions distributed within genes or in flanking regions (±10 kbp), while the third element, SacSINE1-CM, showed only 69.3% insertions in these regions (table 2). Comparisons with simulated data indicated that while SacSINE1-CM observed insertions are significantly less common within both genes and flanking regions, CmiSINEs insertions are significantly less common within genes but significantly more represented in flanking regions ($P < 0.05$) (table 2).

Concerning *L. chalumnae* element LmeSINE1s, the proportion of insertions within genes and flanking regions ranged from 55.0% (LmeSINE1a) to 59.6% (LmeSINE1c). Comparisons with simulated insertions profiles indicated quite different patterns (table 2). LmeSINE1b and LmeSINE1c observed insertions are generally not significantly different from simulated ones, except for LmeSINE1b in the upstream 10 kbp (table 2). On the other hand, LmeSINE1d and LmeSINE1e are both significantly more represented than expected in genes and flanking regions, except for LmeSINE1e in downstream flanking regions (table 2). Finally, LmeSINE1a turned out to be significantly less represented in both genes and flanking regions (table 2).

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SINE Distribution in *C. milii* Transcriptome

BLAST comparison with *C. milii* microRNAs (miRNAs) collection revealed significant similarities with both CmiSINEs and SacSINE1-CM. CmiSINEs showed significant similarity with the precursor of eshark_NOVEL_91 (acc. no. JX994806; identity = 82.6–85.9%) and SacSINE1-CM resulted similar to
precursors of eshark_NOVEL_211 and eshark_NOVEL_234 (acc. nos. JX994610, JX994511; identity $= 76.5\%$ and $78.9\%$, respectively) (fig. 5). The extent of similarity includes the SINE HCD (CORE in the CmiSINE and Deu in SacSINE1-CM) and, in two instances, part of the tRNA-related head. The predicted mature miRNA sequence resulted homologous to a fragment either of the tRNA-related head or of the HCD itself (fig. 5).

We then checked the presence of SINEs within C. milii messenger RNAs (mRNAs). BLAST search retrieved 104 positive hits for CmiSINE and 34 for SacSINE1-CM: all hits are located within the 3' UTR (supplementary tables S1 and S2, Supplementary Material online).

**Discussion**

Short interspersed elements are almost ubiquitous, fast evolving components of the eukaryotic genome that may have a profound impact on the host genome (reviewed in Schmitz 2012). In the present analysis, we describe the life-cycle and genome biology of two SINEs (SINE2-1_CM and SINE2B-1_CM, here collectively referred to as CmiSINEs) within the genome of the elephant shark *Callorhinchus milii*, and their relationships with pre-existing SINEs.

**Evolutionary History and Genome Invasion of CmiSINEs**

We found CmiSINEs only in the *C. milii* genome, coexisting with the newly discovered element SacSINE1-CM. This SINE has homologs distributed in the genome of other cartilaginous fishes (SacSINE1 family in *S. acanthias*, *R. typus*, and *L. erinacea*) and, although with a quite divergent sequence, in the coelacanth (LmeSINE1 family).

We discovered that CmiSINEs belong to the CORE-SINE superfamily, as they show similarity with the CORE domain,
a HCD sequence retrieved among several SINE families isolated in widely different host species (Gilbert and Labuda 1999, 2000; Vassetzky and Kramerov 2013; Nishihara et al. 2016). The CORE domain is assumed to be conserved since the Radiata-Bilateria split, being therefore more than 800 million years old (Vassetzky and Kramerov 2013). The CmiSINEs end, though, is homologous to that of SacSINE1 and LmeSINE1 families, and to the 3' end of a L2 LINE. As the SacSINE1 family harbors the Deu domain, the two SINEs appear to have been originated independently. Yet, their activities appear linked because they share the exploitation of the same LINE.

The de novo origin of a SINE is a chimera between a small RNA gene, such as tRNA, 7SL RNA or 5S RNA, and other sequences (Kramerov and Vassetzky 2011), most likely occurring upon reverse transcriptase template switch. This process is also known to occur in retrovirus recombination (Negroni and Buc 2001) and produced chimeric transcripts in Homo sapiens (Buzdin et al. 2002). This allows RNAs from different origins to merge into a single molecule that can be, then, reintegrated. For example, in the platypus genome, small nucleolar RNAs (snoRNA) formed a chimera with a LINE-related RTE retrotransposon tail; the resulting element was, then, replicated up to 40,000 copies through a SINE-like duplication (Schmitz et al. 2008). Furthermore, it is well known that the highly dynamic nature of SINES is also due to the frequent recombination with other elements, exchanging sequence modules (the head, the body and the tail; Takahashi and Okada 2002; Deragon and Zhang 2006; Kramerov and Vassetzky 2011; Luchetti and Mantovani 2013). As the CORE domain is considered an ancestral component of some SINE families, we can speculate that an undetected/extinct CORE-SINE switched the head with a new tRNA after the holocephalan lineage divergence. Similarly, a proto-CmiSINE acquired a new 3' end, containing the region homologous to the tail of a functional LINE, through recombination with a SacSINE1-CM element. The possibility of recombination or template switching with the L2 LINE element can be ruled out as the region homologous to SacSINE1-CM is larger than that homologous to the L2 tail (fig. 1; supplementary fig. S1, Supplementary Material online). The 3' end switch would have conferred to CmiSINEs the retrotransposition competency and, therefore, the ability to successfully invade the host genome.

As evidenced by the age analysis, the copy number of CmiSINEs increased when SacSINE1-CM started to decrease, reaching the peak of activity more recently (figs. 3 and 4). It is possible that a competition between SacSINE1-CM and CmiSINEs was established. What may have determined the success of CmiSINEs over the SacSINE1-CM?

### Table 2
Genomic Distribution of Observed and Simulated SINE Insertions (± Standard Deviation)

| SINE       | Location       | Observed Insertions | Simulated Insertions (± S.D.) | Results of Significant Comparisons* |
|------------|----------------|---------------------|------------------------------|-------------------------------------|
| SINE2-1_CM | Upstream genes | 29,556              | 27,591 ± 162                 | >                                   |
|            | Within genes   | 76,000              | 81,194 ± 213                 | <                                   |
|            | Downstream genes | 29,487             | 27,570 ± 162                 | >                                   |
| SINE2B-1_CM| Upstream genes | 22,418              | 19,825 ± 136                 | >                                   |
|            | Within genes   | 54,015              | 58,364 ± 176                 | <                                   |
|            | Downstream genes | 22,641             | 19,807 ± 135                 | >                                   |
| SacSINE1-CM| Upstream genes | 30,323              | 34,271 ± 177                 | <                                   |
|            | Within genes   | 90,622              | 100,836 ± 233                | <                                   |
|            | Downstream genes | 30,258             | 34,242 ± 179                 | <                                   |
| LmeSINE1a  | Upstream genes | 5,156               | 5,429 ± 71                   | >                                   |
|            | Within genes   | 29,903              | 30,778 ± 143                 | <                                   |
|            | Downstream genes | 5,146              | 5,415 ± 71                   | <                                   |
| LmeSINE1b  | Upstream genes | 2,709               | 2,524 ± 49                   | >                                   |
|            | Within genes   | 13,918              | 13,796 ± 89                  | <                                   |
|            | Downstream genes | 2,589              | 2,516 ± 49                   | >                                   |
| LmeSINE1c  | Upstream genes | 39                  | 38 ± 6                       | <                                   |
|            | Within genes   | 227                 | 209 ± 11                     | >                                   |
|            | Downstream genes | 32                | 37 ± 6                       | <                                   |
| LmeSINE1d  | Upstream genes | 5,585               | 5,429 ± 71                   | >                                   |
|            | Within genes   | 30,584              | 29,641 ± 132                 | <                                   |
|            | Downstream genes | 5,575              | 5,415 ± 71                   | >                                   |
| LmeSINE1e  | Upstream genes | 2,917               | 2,726 ± 50                   | >                                   |
|            | Within genes   | 15,468              | 14,962 ± 93                  | >                                   |
|            | Downstream genes | 2,713              | 2,720 ± 50                   | >                                   |

*Symbols “<” and “>” indicate comparisons where observed insertions are significantly less or more (P < 0.05), respectively, than simulated ones.
was already replicating when CmiSINE ancestors originated; therefore, it is possible that it simply came to the end of its life-cycle by accumulating too many mutations and/or defective copies (Roberston 2002). Another explanation, not necessarily excluding the previous one, could be that the holocephalan genome already established a defense mechanisms against SacSINE1-CM multiplication, like miRNA suppression (Smalheiser and Torvik 2006) and/or histone methylation (Varshney et al. 2015), when CmiSINEs started their activity. This would have favored the replication of CmiSINEs which, at that time, lacked a limiting mechanism, further contributing to the under-replication of SacSINE1-CM.

SINE Replication Waves and the Evolution of Host Lineages

To analyze the timing of SINEs activity we applied a strict molecular clock to the divergence from the consensus of scored element copies, using two substitution rate estimates. One is calculated on presently analyzed data, using the divergence between Elasmobranchii and Holocephali as a maximum age for CmiSINEs, while the other one was calculated on synonymous substitution rate of three nuclear genes (Martin 1999). The two estimates vary widely, the latter substitution rate being about six times slower: in fact, its use resulted in an estimate of the SacSINE1-CM replication wave older than the metazoan origin (Blair 2009). As this

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**A**

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**B**

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**Fig. 5.**—Blast analysis of C. milii miRNA transcripts against CmiSINE (A) and SacSINE1-CM (B) consensus sequences. Mature miRNA sequences are in bold red; SINE tRNA-related head is shaded in grey and the highly conserved domain sequence (CORE or Deu) is underlined.
appears quite unlikely, we decided to disregard results from this substitution rate. The substitution rate based on presently analyzed data, further, gave age estimates more consistent with the evolutionary history of the considered species.

Since the origin of the holocephalan lineage, about 421 Ma, we detected three replication waves: the SacSINE1-CM one, \( \sim 200-220 \) Ma, and the CmiSINE ones, occurring \( \sim 140-160 \) Ma and \( \sim 100-120 \) Ma. Although having a potentially large error associated with the assumptions we made for the age calculation (i.e., strict molecular clock and constancy of SINE copies substitution rate), this dating suggests that replication waves occurred right after two major events in the evolution of Holocephali: first, the end-Permian extinction about 250 Ma, when, as suggested by fossil records, it seems that a single lineage survived by habitat shifting and gave rise to modern Holocephali (Grogan and Lund 2004); second, the diversification of modern Holocephali, which occurred about 167 Ma (Inoue et al. 2010).

The LmeSINE1 family in the coelacanth genome showed a pattern with two main replication waves, \( \sim 120 \) and \( \sim 50 \) Ma, corresponding to the activity of two elements: LmeSINE1d and LmeSINE1a, respectively. This double-peak activity profile is in line with the analysis carried out on the full TE complement (Chalopin et al. 2014). Interestingly, the most recent replication wave occurred close to the divergence time of the two coelacanth species, \( L. \) chalumnae and \( L. \) menadoensis (30–40 Ma; Inoue et al. 2005).

On the whole, it appears that some SINE mobilization bursts in \( C. \) milii and \( L. \) chalumnae occurred relatively close to major events in the host taxon evolution. It is thus intriguing to ask whether this correlation provides some causal relationships. TE bursts are explosive replication waves of elements that may occur under certain conditions. For example, environmental stresses acting on host species (Capy et al. 2000; Grandbastien et al. 2005; Zeh et al. 2009) or genomic “shocks”, such as host lineage hybridization, may induce TE bursts (O’Neill et al. 1998; Labrador et al. 1999; Wang et al. 2010). Increases of TE copy number may trigger genomic restructuring and it is considered a generator of diversity, possibly underlying speciation events (the TE-Thrust hypothesis; Rebollo et al. 2010; Oliver and Greene 2011; Belyayev 2014). Furthermore, some TE insertions may also have adaptive significance (Gonzalez et al. 2010; Kohler et al. 2015), supporting their role in species diversity and evolution. Examples of correlations between TE expansion and speciation events have been given in mammals (Pace and Feschotte 2007; Ray et al. 2008; Jurka et al. 2011) as well as in bony fishes (De Boer et al. 2007; Symonova et al. 2013). Thus, it could be hypothesized a possible role of SINE insertions in species diversification and adaptation.

Most of the SINE insertions in both \( C. \) milii and \( L. \) chalumnae occurred within genes and their flanking regions. In the elephant shark, though, CmiSINES have accumulated in gene flanking regions, but they appear as preferentially sorted out from genes. On the other hand, SacSINE1-CM resulted significantly less represented in the whole genomic regions (i.e., genes and flanking regions). Conversely, the picture in the coelacanth genome is more complex: the most recent active element, LmeSINE1a, appears as preferentially sorted out from the whole genomic regions, at variance of those elements that burst in the past. In \( L. \) chalumnae both past and recent TE activity has been taken as an indication of a dynamic genome, in contrast to the morphological stasis (Chalopin et al. 2014); moreover, it has been suggested that TE activity affected postspeciation divergence of the two \( L. \) chalumnae species (Naville et al. 2014). Although direct evidence cannot be drawn from this analysis, our data on LmeSINE1 elements genomic distribution appear partially in line with the role proposed for TEs in genome evolution. On the other hand, the replication wave of CmiSINES in the holocephalan genome occurred right after the main lineage diversification, indicating that these elements might be not directly implicated in the modern Holocephali radiation. Though, the preferential insertions accumulation in gene flanking regions could hint at possible roles in the \( C. \) milii lineage evolution.

TE replication waves may be not causative but rather caused by host’s evolutionary history, and may have affected genome evolution subsequently. For example, TE massive burst occurred during the domestication of rice, thus TE insertions were not causative of cultivar evolution even if some insertions became part of the new regulatory network (Naito et al. 2009). In the “Carrier Subpopulation” hypothesis, Jurka et al. (2011) linked TE burst with species fragmentation in subpopulations. As subpopulations can lead to speciation, hence TE replication waves correlate with species diversification, being not causative but rather accompanying the process.

Obviously, this does not exclude that new insertions may trigger species-specific genomic changes (Jurka et al., 2011). From data presented here, we can infer a possible role of \( C. \) milii SINES in the genome functionality and, thus, in its evolution. In fact, both CmiSINES and SacSINE1-CM sequences match miRNA sequences. These are small RNAs that act as post-transcriptional regulators of gene expression by binding target mRNAs at their 3’ UTR (He and Hannon, 2009). Accordingly, we found mRNAs transcribed from 138 genes carrying CmiSINES or SacSINE1-CM insertions within the 3’ UTR. Therefore, our data point out that SINES can be transcribed as regulatory RNAs and that they may regulate the expression of these 138 genes. Post-transcriptional gene expression regulation by SINE-derived miRNA have been found for primate Alus and fish V-SINES: in both instances SINES located at mRNAs 3’ UTR act as binding sites for regulatory miRNAs (Daskalova et al. 2006; Scarpato et al. 2015). Moreover, a possible role for the regulation of SINES activity cannot be excluded, with SINE-derived miRNA binding retrotransposing SINE transcripts. Overall, data presented here reinforce the hypothesis of a functional role for these elements.
SINEs Genomic Distribution and Highly Conserved Domains

HCD-SINEs represent an apparent paradox as they show a long-term conserved element, the HCD in the body module, in an otherwise fast evolving sequence, the SINE itself. Many attempts have been made to explain the conservation of such regions and two main hypotheses, not necessarily mutually exclusive, appear as more reliable: i) HCD are selectively conserved because they may serve as SINE-SINE recombination hotspot, or ii) HCD are selectively conserved because they provide some advantage to the host genome, having some still undetermined, albeit important, function (reviewed by Deragon 2012). Some highly conserved SINE domains have been implicated in exaptation events, being domesticated as enhancers (Bejerano et al. 2006; Santangelo et al. 2007; Sasaki et al. 2008; Nakanishi et al. 2012). If HCDs have some function exploitable by the host genome for gene functionality, one might expect to find them more frequently associated to coding regions. In this study, we provide evidence that two different HCD-SINES, CORE- (CmiSINEs) and Deu-SINES (SacSINE1-CM), may have different genomic distribution in the same genome. In fact, while both HCD-SINES are preferentially sorted out from gene regions, CmiSINEs appear to accumulate in the flanking regions.

Data on SacSINE1-CM are in line with data on another Deu-SINE, the amniote-wide AmnSINE (Hirakawa et al. 2009). On the other hand, elements of the LmeSINE1 lineage—a Deu-SINE family homologous to SacSINE1-CM—may show preferential occupancy within genes and their flanking regions, a random distribution or even a significant depletion from gene regions. Thus, it is likely that HCD-SINES may affect genomes in different ways.

Present data do not allow us to suggest a specific function for the Deu domain of LmeSINE1, but a possible model for both CmiSINEs and SacSINE1-CM can be envisaged. In these instances, HCDs of the two SINES match almost entirely to miRNA precursors, a form that is not simply an intermediate but may have regulatory activity (Roy-Chaudhuri et al. 2014). Therefore, the possibility to be transcribed as miRNA and to act also as a target for miRNA post-transcriptional regulation would imply an active function for their HCD. This could be the first evidence of a specific function for these paradoxical highly conserved domains. Other HCD-SINES have been found to act as miRNA target (V-SINES); on the other hand, it was not specified whether they are also transcribed in miRNA precursors nor which part of the SINE is included in the transcript (Scarpato et al. 2015). It would be, therefore, interesting to check if other HCD-SINES are actively transcribed as miRNA and if the HCD itself is involved in the transcription.

Supplementary Material

Supplementary data are available at Genome Biology and Evolution online.

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