RESEARCH



Open Access

Examples of sequence conservation analyses capture a subset of mouse long non-coding RNAs sharing homology with fish conserved genomic elements

Swaraj Basu¹, Ferenc Müller², Remo Sanges^{1*}

From Ninth Annual Meeting of the Italian Society of Bioinformatics (BITS) Catania, Sicily. 2-4 May 2012

Abstract

Background: Long non-coding RNAs (IncRNA) are a major class of non-coding RNAs. They are involved in diverse intra-cellular mechanisms like molecular scaffolding, splicing and DNA methylation. Through these mechanisms they are reported to play a role in cellular differentiation and development. They show an enriched expression in the brain where they are implicated in maintaining cellular identity, homeostasis, stress responses and plasticity. Low sequence conservation and lack of functional annotations make it difficult to identify homologs of mammalian IncRNAs in other vertebrates. A computational evaluation of the IncRNAs through systematic conservation analyses of both sequences as well as their genomic architecture is required.

Results: Our results show that a subset of mouse candidate IncRNAs could be distinguished from random sequences based on their alignment with zebrafish phastCons elements. Using ROC analyses we were able to define a measure to select significantly conserved IncRNAs. Indeed, starting from ~2,800 mouse IncRNAs we could predict that between 4 and 11% present conserved sequence fragments in fish genomes. Gene ontology (GO) enrichment analyses of protein coding genes, proximal to the region of conservation, in both organisms highlighted similar GO classes like regulation of transcription and central nervous system development. The proximal coding genes in both the species show enrichment of their expression in brain. In summary, we show that interesting genomic regions in zebrafish could be marked based on their sequence homology to a mouse IncRNA, overlap with ESTs and proximity to genes involved in nervous system development.

Conclusions: Conservation at the sequence level can identify a subset of putative lncRNA orthologs. The similar protein-coding neighborhood and transcriptional information about the conserved candidates provide support to the hypothesis that they share functional homology. The pipeline herein presented represents a proof of principle showing that a portion between 4 and 11% of IncRNAs retains region of conservation between mammals and fishes. We believe this study will result useful as a reference to analyze the conservation of IncRNAs in newly sequenced genomes and transcriptomes.

* Correspondence: remo.sanges@szn.it

¹Laboratory of Animal Physiology and Evolution, Stazione Zoologica Anton Dohrn, Villa Comunale, 80121, Naples, Italy

Full list of author information is available at the end of the article



© 2013 Basu et al.; licensee BioMed Central Ltd. This is an open access article distributed under the terms of the Creative Commons Bio Med Central Attribution License (http://creativecommons.org/licenses/by/2.0), which permits unrestricted use, distribution, and reproduction in any medium, provided the original work is properly cited.

Background

Long non-coding RNAs (lncRNAs) were firstly reported as transcripts expressed in large numbers in mammalian transcriptomes [1,2]. They were shown to constitute more than half of all the transcriptional outputs of mammalian genomes [3]. Prior to these reports Xist in mammals and *Xlsirt* in amphibians were the only well characterized IncRNAs described to function in X; chromosome inactivation and the formation of cytoskeleton [4,5]. Recently, several aspects of development and disease have been associated with lncRNAs function albeit to a small proportion of them. Processes associated with lncRNAs include epigenetic regulation of multi-gene loci [6,7], apoptosis and cell cycle [8], regulation of gene splicing [9] and tumor suppressor activity [10]. There are also examples of IncRNAs playing a role in the adaptive immunity of mammals [11], being differentially expressed in response to carcinogens [12] and functioning as enhancers [13].

A probe into the functional role of lncRNAs requires knowledge of the specificity of their expression across developmental stages and different tissues. Microarray technology was initially used to detect lncRNAs in differentiating mouse embryonic stem cells. Many lncRNAs showed correlation of expression with the flanking protein-coding genes that associated with developmental functions [6]. Similar transcriptomics studies identified many lncRNAs expressed in CD8+ cells and during neuronal lineage specification in mouse [11,14]. However, cross-hybridization, background noise and limited genome coverage associated to array based detection techniques were often used as arguments against the pervasive existence of lncRNAs. Today these problems have been solved thanks to the advent of the RNA sequencing technology (RNAseq) [15]. RNAseq deals with direct quantification of a cDNA population and is not limited to transcripts mapped to known genomic sequences. An RNAseq experiment on differentiating human neurons led to the identification of about 1,600 lncRNAs with dynamical expression levels [16]. Recently, sequencing studies have even identified hundreds of lncRNAs expressed during early developmental stages in zebrafish and C. elegans finally demonstrating that the pervasiveness of non-coding transcription is not an exclusive feature of mammalian transcriptomes. Pauli et al identified about 1,100 non-coding RNAs expressed during embryogenesis in zebrafish [17]. Ulitsky et al reported a set of about 700 long intergenic non-coding RNAs (lincRNAs) expressed during zebrafish development; they show low sequence similarity but conserved genomic locations with their mammalian counterparts [18]. Finally Nam et al reported a catalog of 230 lncRNAs in C.elegans [19] which show similar features to vertebrate lncRNAs in terms of lack of general sequence conservation as well as stage specific expression patterns. The execution of large-scale studies about the discovery and characterization of candidate lncRNAs is facilitating the establishment of a catalog of these important molecular players which, in turn, permit to annotate them with specific biological validations and comparative analyses improving our knowledge of the lncRNAome [20].

However, lack of sequence homology and lack of deep tissue and stage specific expression data have been one of the obstacles in defining a proper catalog of lncRNAs among different species. Computational methods for the discovery and the annotation of lncRNAs are sparse and mainly limited to mammalian genomes. Parameters like open reading frame length, lack of homology to proteincoding genes and protein domains, nucleotide composition and substitution rates have been used previously to define computational lncRNA discovery pipelines [21-24]. Sequence conservation has not been often used as support to justify the presence of lncRNAs in an organism because the lncRNAs identified in human, mouse, zebrafish and C. elegans showed little or no sequence conservation in the majority of the population [17-19,25]. In addition, Pauli et al stated that the level of conservation for the majority of zebrafish lncRNA is comparable to that of introns and that only few of them are really conserved [17]. However, short spans of sequence conservation for specific lncRNAs were reported [18]. Four evolutionary constrained mouse IncRNAs were shown to be conserved in sequence and expression between mouse, opossum and chicken [26]. Forty three putative long non-coding sequences from chicken ESTs were found to share sequence homology with human, rat and mouse transcripts [27]. Xist, the IncRNA responsible for the X chromosome inactivation in eutherian mammals shares sequence homology in 14 vertebrates [28]. Sox2ot (Sox2 overlapping transcript) and Har1F (Human accelerated region 1F) are conserved amongst vertebrates [29,30]. The biggest group of IncRNAs constrained in terms of their nucleotide substitution rates between mouse and human is represented by a set of 659 mouse transcripts [31]. This set showed a tendency to juxtapose genes involved in the regulation of transcription and development. A subset of these lncRNAs expressed in the mouse brain and defined as CNS-specific appear to show tissue specific expression pattern similar to their proximal protein coding genes. Despite these evidences, a lack of general sequence conservation associated with lncRNAs has been proposed and this becomes more evident when considering long evolutionary distances such as the one separating mammals from fishes, even if fragments of sequence conservation have been detected [18,27].

Despite these evidences, a systematic analysis of sequence conservation of vertebrate lncRNAs is still

lacking and it remains unclear whether and to what extent there is general sequence conservation of lncRNAs between mammals and fishes. In order to fulfill this lack, the work here presented tries to define the level of conservation of mouse lncRNAs in the zebrafish genome using an unbiased choice for the comparison parameters, taking advantage of randomizations and receiver operating characteristic (ROC) analyses. Our approach is centered around subsets of mouse constrained and/or well annotated lncRNAs, based on the assumption that such transcripts constitute a representative set of lncRNAs ideally containing small amounts of transcriptional noise. We would like to specify that, although our analysis consider a subset of already published and very well annotated lncRNAs, we cannot rule out the possibility that some of them might actually be coding for short peptides. Recent studies suggest that few candidate lncRNAs can produce short peptides and there may exist a class of bifunctional RNAs encoding both mRNAs and functional noncoding transcripts [32-34]. Therefore, specific biological validations remain a fundamental step for a proper characterization of these elements. Nevertheless, the special care taken in the choice of the datasets to analyze and the literature about them, makes us confident that we are currently using, the best sets of candidate lncRNAs. We compared these transcripts against the set of zebrafish phastCons elements [35] reported to be significantly conserved among fishes. The phastCons program uses a hidden Markov model-based method that estimates the probability that each nucleotide belongs to a conserved element, based on multiple alignments of selected species. We used the phastCons6way track to select elements conserved among fishes. These are based on scores built on multiple alignment of the zebrafish genome with tetraodon, stickleback, human, mouse and western clawed frog. It is important to point out here that, considering the way in which they are built, these elements represent the best selection of sequences conserved, in first instance, among fishes, but many of them can also result conserved among vertebrates. This choice implicitly adds more genomes to our analyses and is based on the assumption that lncRNAs conserved between mouse and zebrafish are expected to be primarily conserved among teleosts. For this pilot study, the reduction in the dataset dimension, given by such choice, limited the zebrafish genomic search space to the phast-Cons sequences, rather than to the full genome, making it feasible to use several randomizations steps (shuffling of the query sequences) to specifically identify the levels of conservation proper of lncRNAs.

Here we show that the usage of the BLASTn e-value and alignment length as cut-offs is sufficient to distinguish conservation of mouse lncRNAs against zebrafish phastCons elements as compared to shuffled sequences. From an initial dataset of about 2,800 mouse lncRNAs we demonstrate that between 4 and 11% of them contain fragments significantly conserved in zebrafish in agreement with the results by Ulitsky et al [18] on a smaller dataset. Gene ontology enrichment analyses of proteincoding genes flanking the conserved elements, identified similar functional classes significantly enriched in both species, such as regulation of transcription and development. These coding genes exhibited enrichment for expression in the brain in both mouse as well as zebrafish. The lncRNAs shown to be conserved are deemed to be functionally important and suggested for further experimental validation of their function.

Results and discussion

Selection of conservation parameters to select significantly conserved IncRNAs

We developed a pipeline to identify conserved mouse lncRNA fragments in zebrafish using sequence identity, randomization and the identification of an unbiased threshold to detect significant levels of conservation (Figure 1 A). In order to identify the optimal parameters capable to select conserved lncRNA sequences, we used receiver operating characteristic (ROC) like analyses on the distribution of the following BLASTn alignments result measures: 1) query coverage, 2) query alignment length, 3) percentage identity and 4) e-value. ROC like analyses were performed on the results of the following BLASTn searches: 1) mouse lncRNA against zebrafish phastCons elements (true positive set), 2) shuffled mouse IncRNA sequences against zebrafish phastCons elements (false positive set). In order to select significant results we defined a specific cut-off showing less than 0.05% false discovery rate (FDR) for each parameter. The analysis was applied to different datasets and, after the application of the identified filter, between 4 to 11% of the sequences in the true positive datasets resulted to be significantly conserved. Conserved fragments show a mean length of about 160 nucleotides and an average identity of about 80% with their corresponding mouse lncRNA fragments (Figure 1B and 1C). Specifically, mouse candidates lncRNAs from two sources representing three datasets, were used to determine sequence conservation. Mammalian constrained lncRNAs from mouse (659 transcripts defined as CNS/NCNS dataset) [31] were divided into Central Nervous System specific (239 CNS transcripts) and non-CNS specific (420 NCNS transcripts) giving rise to the first and second datasets. LncRNAs identified in the mouse genome by the Ensembl lincRNA annotation pipeline [36,37] (2,147 EnsEMBL transcripts, EnsEMBL version 62) formed the third dataset. An initial exploratory analysis was performed by using BLASTn with word lengths ranging from 8 to 11 nucleotides on the CNS dataset. ROC curves, plotting the distributions



of the indicated measures (Figure 2 A) suggest that the reciprocal of the e-value (1/e-value) is the factor capable to better segregate results between the true positive and

false positive sets (area under curve, AUC = 0.79). We plot the reciprocal of the e-value (1/e-value) because plotting the e-value produced curves significantly skewed

0.

0.6

0.4

0.2

0.0

0

0.8

0.6

0.4

0.2

0.0

1.0 8

0.8 2

9.0

0.4

0.2

0.0

1.0

0.8 36 e positive rate

0.6

0.4

0.0 0.2 0.4 0.6 0.8 1.0

0.79 0.2 0.0

74

0.0 0.2 0.4

True positive rate

1.0

0.96

True positive rate

0.82

0.78

1.0

0.18

0.15 0.8

True positive rate

QCoverage

False positive rate

Pldentity

False positive rate

QCoverage

False positive rate

Pldentity

0.

0.8

0.6

0.4

0.2

0.0

0.1

0.8

0.6 0.4

0.2

0.0

1.0

0.8

0.2

0.0

1.0

0.8

0.4

0.2

0.0

0.0 0.2 0.4 0.6 0.8 1.0

rate 9.0 0.0

tive rate 0.6

True posit 0.4 0.0

0.2 0.4 0.6 0.8

0.2 0.4 0.6 0.8

Frue positive rate

A

0.0 0.2 0.4 0.6 0.8 1.0

ale

True positive





1085.6

729.4 373.2

1.0

0.6

False positive rate

1/EValue

0.8

below the diagonal line [38]. In addition, alignment length presents an AUC of 0.64 and, at a manual inspection of results, we noticed that this measure is capable to filter out low complexity (repeated) regions that, in few results, show multiple hits with a small e-value and are hence retained by the exclusive e-value filter. It is now becoming evident that repeats are enriched in lncRNAs [39,40] At the light of such considerations, we decided to combine the 2 measures in order to select significantly conserved lncRNAs, avoiding to obtain low complexity regions in the set of results. Combining the two parameters for filtration gave us zero false positives for each dataset. Interestingly, the change in word size does not affect the performance of the classifier (Additional File 1). Therefore, word size of 11 nucleotides is used in all subsequent analyses. We selected as significantly conserved, lncRNAs sequences showing <0.05% FDR for each of the 2 parameter (e-value and alignment length). Cut-off values were calculated in order to consider significant a percentage of false positives smaller than 0.05% when the same filter is applied to the randomized data. An e-value cutoff of 5e-05 and an alignment length cutoff of 70 nucleotides satisfied this criteria resulting in 11 lncRNAs from the CNS dataset significantly conserved within the zebrafish phastCons elements (Table 1). The BLASTn search was repeated for the NCNS and the Ensembl datasets (Table 1). ROC curves (Figure 2 B, C) and manual inspection of data confirmed the e-value and query alignment length as the best parameters to successfully identify significantly conserved lncRNAs (AUC NCNS: e-value 0.76, alignment length 0.66; AUC EnsEMBL: e-value 0.82, alignment length 0.70). The identified cutoffs are as follows: NCNS) e-value 1e-04, alignment length 66; EnsEMBL) e-value 2e-04, alignment length 62. The results and the annotations of the homology searches for all 3 datasets can be found in Additional File 2. We also performed secondary structure analyses to test if we could segregate the false and true positives more efficiently using RNAz on the BLASTn alignments. RNAz is an efficient method for detecting functional RNAs combining comparative sequence analysis and structure prediction. The program performs two basic calculation: 1) the measure for RNA secondary structure conservation and 2) the measure for thermodynamic stability [41]. The 3 RNAz result parameters used to build the ROC curves were: ratio of pairwise identity by sequence conservation index, Z score and P value (1/P value) (Additional File 3). The sequence conservation index demonstrated a positive performance (AUC 0.74) in agreement with previous reports about structural conservation of conserved lncRNAs [42,43]. However, its performance is lower than the e-value and it is not able

to filter out low complexity regions. Therefore, we did not use this measure in the rest of the analyses.

Comparison of the genomic contexts of mouse IncRNA and fish phastCons pairs predicted to be conserved

In order to evaluate the locations and shed light on the putative functions of each conserved fragment, we mapped and compared each element in the respective genic context of both analyzed organisms. The 11 putatively conserved IncRNAs in the CNS dataset showed homology to 10 phastCons elements. The NCNS dataset had 23 lncRNAs showing homology to 21 phastCons and the 250 conserved Ensembl lincRNAs showed homology to 209 fragments from 197 phastCons elements. The conserved regions in zebrafish were checked for overlapping features (Table 2, 3). The fragments from the CNS dataset show 6 out of 10 elements overlapping non-coding regions (intergenic, intronic or non-coding exon) in zebrafish and 4 out of 11 in mouse. Regarding the NCNS dataset, 17 out of 22 conserved sequences are present in a non-coding region in zebrafish and 13 out of 23 in mouse. The situation for the conserved Ensembl dataset is different as a minor fraction of elements is present in non-coding regions in zebrafish and in mouse (18% in zebrafish and 27% in mouse). Such difference with the Ensembl data may be explained by considering that CNS and NCNS lncRNAs are curated for being mainly intergenic as compared to the Ensembl IncRNAs. Therefore, in the Ensembl dataset, candidate IncRNA fragments may overlap an external exon of a coding gene in the same chromosomal domain more frequently. However, they must still be considered non-coding because the orientation of the transcripts is in antisense to the protein coding genes they partially overlap (see genes ENSMUSG0000060029 and ENSMUSG00000046413 as an example) and they do not show any significant open reading frame (ORF). Antisense transcripts are reported to be present in large numbers in mammalian genomes and often linked to the regulation of neighboring or overlapping protein-coding genes [44]. Indeed, long non-coding RNAs can influence the expression of protein coding genes in *cis* as suggested in a previous report [31]. They are also reported to be associated with enhancers of neighboring

Table 1 LncRNAs conservation

Dataset	Word Size	Number conserved IncRNAs	Percentage conserved IncRNA	Percentage conserved shuffled
CNS (239)	11	11	4.60%	0.0%
	10	11	4.60%	0.0%
	9	11	4.60%	0.0%
	8	11	4.60%	0.0%
NCNS (420)	11	23	5.40%	0.0%
Ensembl (2,147)	11	250	11.6%	0.0%

The number of lncRNA putatively conserved in each dataset (CNS, NCNS, Ensembl) after applying the query alignment length and e-value cutoffs on the produced alignments.

Dataset	Total aligned regions	Coding exon overlap	Noncoding exon overlap	Intron overlap	Intergenic
CNS	11	7	0	3	1
NCNS	23	10	3	5	5
Ensembl	250	183	31	17	19

Table 2	2 Genomic	location	of	conserved	regions	in	mouse
---------	-----------	----------	----	-----------	---------	----	-------

The genomic locations for the number of mouse IncRNA fragments found to be conserved. The location is deduced with respect to the coding region of the mouse genome in the area of alignment.

coding genes in mouse neurons [45] and human [13]. In order to test if the function of flanking coding genes corroborates the functional conservation suggested for each mouse and zebrafish conserved non-coding pair we identified the coding genes flanking and overlapping the aligned region in zebrafish and their mouse counterparts and evaluated their homology relationships. The search for orthologs was performed scanning a window of 1 megabase flanking the conserved elements in either direction in the 2 genomes (see methods). Results are depicted in3. The figure shows the percentages of conserved mouse lncRNAs sharing orthologous coding gene in the corresponding zebrafish genomic context. All the lncRNA conserved fragments showed at least one ortholog pair from the CNS, the NCNS along with 80% of the Ensembl datasets supporting the hypothesis of syntenic conservation.

Functional enrichment analyses of the protein coding genes proximal to the conserved regions

In order to understand the potential biological role of the identified sequences we performed gene ontology and tissue specific expression enrichment analyses on the coding genes flanking the conserved fragments for the Ensembl dataset. The coding genes flanking and eventually overlapping the conserved regions were considered to be the putative lncRNAs associated genes. Significantly enriched GO biological process categories and tissue of expression for the conserved lncRNAs of the Ensembl dataset in zebrafish and mouse were considered for the analysis using DAVID [46,47] and an EASE score cutoff of 0.05. The EASE score is a *p*-value adjustment method specifically designed for biological large-scale studies. It penalizes the significance of categories supported by few genes and favors more robust categories in respect to the Fisher exact probability. It is more conservative than the pure Fisher exact probability and less conservative than the Benjamini and Hochberg FDR [48]. For genes associated to the conserved lncRNAs of the Ensembl dataset, the enriched GO terms included development, regulation of transcription and nucleic acid metabolism in both the organisms (Figure 4A and 4B) in agreement with previous reports in mouse [6,49-51]. Tissue enrichment analyses were also performed to check if the selected genes showed an enrichment for being expressed in similar specific tissues. From this analysis neural and developmental related tissues resulted to be enriched in both the species (Figure 4C and 4D). These results are consistent with previous studies showing that lncRNAs play an important role in regulation, neural development and plasticity [49,50]. It is important to point out that, in mouse, the genes associated to the conserved lncRNAs show the most significant enrichments for expression in the nervous tissues but also a significant enrichment in lung indicating either a possible sub-functionalization of subgroups of lncRNAs or a richer annotation of the mouse transcriptome in terms of domains of expression. Taken together, these analyses highlight a conserved pattern of functions and expression domains of coding genes associated with conserved lncRNA fragments. CNS and NCNS datasets were not used independently because they are not of reasonable dimensions to perform enrichment analysis, however, if we join the 2 datasets together, similar enrichments are obtained (Additional File 4). In 2004 the presence of ultra conserved elements (UCEs) in the human genome was discovered. These elements show about 100% of sequence identity with mouse and many of them are conserved also in fishes. UCEs are greater than 200 nucleotides in length and observed to lie proximal to coding genes related to development, regulation of transcription [52] and cancer related loci [53]. A small fraction of them overlap protein coding exon, however UCEs are

Table 3 Genomic location of conserved regions in zebrafish

Dataset	Total aligned regions	Coding exon overlap	Noncoding exon overlap	Intron overlap	Intergenic			
CNS	10	4	0	1	5			
NCNS	21	4	2	3	12			
Ensembl	209	171	6	9	23			

The genomic location for the number of phastCons fragments found to be conserved. The location is deduced with respect to the coding region of the zebrafish genome in the area of alignment.



mainly non-coding in nature and, although a large fraction seems to be transcribed and/or to function as enhancer they do not overlap current collections of transcripts [53-55]. In order to check if the identified sequences might belong to the ultraconserved family of elements we checked their overlap with UCEs [52,56]. In total four UCEs were found to overlap conserved regions from lncRNAs of the EnsEMBL dataset while a single lncRNA from the NCNS dataset showed overlap with a single UCE. We conclude that the conserved regions identified in this study are not enriched for and do not correspond to UCEs elements. Therefore, they have not to be considered ultraconserved.

Potential of expression of conserved regions in zebrafish

The presence of expressed sequence tags (ESTs) overlapping the region of conservation might support an active transcriptional output in the given region. In this context we chose to check for the overlap of zebrafish ESTs in the region of conservation. Respectively 60%, 45% and



to the EASE score. At maximum the 10 top-scoring significant classes are present into the plots. Grey bars indicate the percentages of genes associated to the respective functional classes from the group of genes flanking the identified conserved elements. Black bars indicate the percentages from the entire transcriptome of the given species.

70% of the predicted CNS, NCNS and Ensembl conserved regions are covered by at least one EST in zebrafish (Table 4). Interestingly, by randomly selecting ~1,200 non-repeated genomic regions of the same extension from the zebrafish genome we obtained a percentage as small as 8% in overlap with ESTs (*p*-value: 7.5e-08, 5.2e-09 and <2.2e-50 respectively for CNS, NCNS and Ensembl dataset). The results are consistent with the possibility that the majority of the conserved regions predicted in the analysis represent actively transcribed regions of the zebrafish genome. Further, in order to add supporting evidences to the potential of expression of the zebrafish conserved

Tabl	e 4	ESTs	overl	apping	the	region	of	conservation
------	-----	------	-------	--------	-----	--------	----	--------------

Dataset	Conserved	EST overlap	
CNS	10	6	
NCNS	21	10	
Ensembl	209	149	

Number of conserved fragments overlapping ESTs in the zebrafish genome.

fragments herein isolated we performed overlap analysis with the recently published zebrafish candidate lncRNAs resulting from RNAseq experiments [17,18]. The comparison of all the predicted conserved regions with the published lncRNAs resulted in 6% of our conserved regions showing overlap with at least one reported lncRNAs. It is important to point out that no definitive estimation of the number of lncRNAs expressed in an organism is currently possible. Such uncertainty arises from the fact that noncoding RNAs are expressed at lower levels as compared to coding genes [17,25,57]. Computational identification of IncRNA transcripts from next-generation sequencing data remains a "work in progress" in terms of mapping reads to the genome, assembly of new transcripts, definition of background noise and cut-off parameters. Hence, in our analysis a lack of overlap does not signify an absence of transcribed elements in zebrafish, but may reflect on undetected transcripts. In order to test this hypothesis we mapped the raw reads from the study [17] (SRP009426) on the zebrafish genome and computed the overlap between the mapped reads and all the conserved fragments. Interestingly, more than 90% of the predicted conserved regions in the zebrafish genome show overlap with at least one mapped read while only 25% of a set of randomly chosen genomic regions overlap at least one read (*p*-value for difference in proportions <2.2e-50). Checking for regions with more than 1,000 reads overlap, we found that 20% of the conserved regions resulted positive while only 4% of random regions showed such an overlap (p = 1.2e-15; Additional File 5). The highly significant differences between the conserved regions and the random sequences indicate that the RNAseq data supports transcriptional evidences in zebrafish for most of the regions predicted to be conserved lncRNAs. Finally, in order to get information about the expression domains for the conserved sequences we took advantage of the publicly available RNAseq study SRP012923. This study contains RNA samples from nine different tissues (heart, kidney, testis, liver, muscle, skin, gill, eye and brain) of Gasterosteus acu*leatus* (stickleback) a sequenced teleost fish. First, we mapped the conserved zebrafish fragments on the stickleback genome and, as expected, all the zebrafish sequences were mapped. Then, we mapped all the raw reads from the SRP012923 study on the stickleback genome and calculated the overlap with the conserved fragments. Again, more than 85% of conserved elements resulted to overlap raw reads (Figure 5 A) consistently with what we observed in zebrafish using the data from Pauli et al. The stickleback data were generated and made publicly available by the Broad Institute. In agreement with the mouse expression data of the corresponding lncRNAs, the conserved CNS sequences show high levels of expression in the brain (Figure 5 B) also in stickleback. Conversely, the NCNS data result to be transcribed at very low or even background levels (Figure 5 C), while the Ensembl dataset shows low but widespread expression (Figure 5 D). Based on these results we can confirm that the mouse CNS elements are likely to be CNS specific also in fishes. Regarding the NCNS dataset we cannot propose conservation of expression between mouse and stickleback, more and deeper sequencing data will probably clarify this aspect. Finally, the observation that the Ensembl sequences show positive expression levels in several tissues is consistent with the fact that the corresponding mouse transcript models are based on transcriptional evidences from multiple tissues and cell cultures [37]. We conclude that the analyses of reads coming from teleost fishes provides supporting evidences for the transcription and, at least for the CNS specific elements, the tissue specific expression of the predicted conserved regions. As long as new sequencing datasets will become available further light can be shed to improve our knowledge about the similarity of the transcriptional outputs among different vertebrate species.

CNS specific IncRNA

To better demonstrate the utility of our analysis we focused on 11 conserved CNS lncRNAs and selected a candidate element ideal for further functional validations. Each phastCons element was given a unique ID at the start of the analysis which is used here as reference (Table 5). The majority of elements show evidence of transcription by overlap with EST sequences in zebrafish and mouse. Many elements also overlap UTRs of protein coding genes pointing towards a putative regulatory function. The conserved element 113364 (Figure 6) belongs to the CNS dataset, it falls completely in an intergenic region in zebrafish and a small part of the lncRNA in mouse overlaps the UTR intron of the coding gene *Lmo3*. The zebrafish sequence shows a conservation of 96 base pairs with the murine lncRNA AK020962 at an e-value of 4e-21 and 88% identity. The lmo3 gene flanks the region of conservation in zebrafish too. This gene is known to be a transcriptional regulator [58] and is reported to be involved in cell proliferation and differentiation during embryonic development [59]. It is also implicated in neuroblastoma through its interaction with the neuronal transcription factor hen2 [59]. The dataset of conserved elements isolated in our analysis will be of help in focusing on specific sets of elements that might have an evolutionary conserved role in development and



differentiation which led to their sequence conservation across species.

Conclusions

Long non-coding RNAs are not characterized by the same sequence conservation properties as protein coding genes. However, in our analysis we demonstrated a systematic procedure to identify significant sequence conservation of candidate lncRNAs in vertebrates. It resulted in the selection of a set of mouse lncRNA fragments significantly conserved in fish genomes demonstrating that a proportion of up to 11% of mouse lncRNAs contains fragments conserved across vertebrates. The candidate lncRNAs reflect a sub-population of the mouse lncRNAome sharing sequence homology with zebrafish phastCons elements. The addition of annotation layers on top of sequence

ID	Genomic feature overlap in Zebrafish (Aligned region of phastCons element)						Genomic feature overlap in Mouse (Whole mouse IncRNA)			
	UTR	Exon	Intron	EST	Other	UTR	Exon	Intron	EST	Other
334146	yes	yes	-	yes	-	yes	-	-	yes	lincRNA (Ensembl)
377442	yes	-	-	yes	-	yes	-	-	yes	-
391744	yes	-	-	-	-	yes	-	-	yes	-
414089	yes	-	-	yes	-	-	-	-	yes	-
759212	-	-	yes		-	-	-	yes	yes	-
113364	-	-	-	-	-	yes	-	-	-	-
208793	-	-	-	yes	-	yes	-	-	yes	miRNA
268839	-	-	-	-	-	yes	-	yes	yes	-
460295	-	-	-	yes	-	yes	-	-	yes	-
604458	-	-	-	yes	-	-	-	yes	yes	-

Table 5 Genomic features of conserved regions overlapping mouse CNS specific IncRNAs

The genomic features for each of the 10 conserved phastCons fragments (from the CNS dataset) and their mouse lncRNA homologs. To be noted that while for zebrafish only the region of alignment is considered not the whole phastCons element, in case of mouse the complete lncRNA was used to perform the analysis. Each region is classified in terms of it being overlapped by a UTR, exon or intron of an annotated protein coding gene, or an EST.

conservation statistics provides biological significance to the results obtained. GO enrichment analyses of coding genes flanking the conserved sequences showed striking similarity at the functional level in both species. In addition, significant neural expression enrichments in both mouse and zebrafish are consistent with previous observations that lncRNAs play an important role in neural development, differentiation and functions. The presence of overlapping ESTs and the significant overlap with developmental zebrafish RNAseq reads provide further support that these fragments produce transcriptional output in fishes. Finally, the RNAseq reads from multiple tissues of the stickleback throw light on the tissue specificity of the conserved regions as well as add additional evidences in support of their expression in fishes. The dataset presented constitute a valuable starting point for future studies aimed at functional characterization of specific lncRNAs.

In summary, this work represents a proof of principle showing that a portion between 4 and 11% of lncRNAs retains short regions of conservation between mammals and fishes and the cutoffs to use to efficiently look for such elements. The analyses can result especially useful as a reference to analyze the conservation of lncRNAs in newly sequenced genomes and transcriptomes.

Methods

Selection of the datasets used for the study

The mouse CNS (Central Nervous System specific) and NCNS (non Central Nervous System specific) constrained lncRNA datasets were obtained from a previous study [31]. Ensembl lincRNA dataset was obtained from BioMart and is based on the Ensembl version 62 [37,60]. The lncRNA sequences in each dataset were shuffled with the shuffle program (part of the SQUID C library by Sean Eddy, the executable can be found in the HMMER3 program) [61]. Each sequence in each dataset was randomized 100 times giving rise to three random sequence datasets rCNS, rNCNS and rEnsembl. PhastCons elements for zebrafish (zPHS) were obtained from the UCSC table browser [35,62] with the "most conserved" option selected for sequence retrieval. The coordinates of the phastCons elements were mapped to the zebrafish current genome build (zv9) using the UCSC liftover tool [63]. A total of 816,471 conserved elements could be mapped out of 881,975 original elements.

Identification of sequence homology between the IncRNAs and the phastCons elements

The mouse lncRNAs (CNS, NCNS, Ensembl) as well as the random datasets (rCNS, rNCNS, rENsEMBL) were searched individually against the zPHS using BLASTn from the BLAST+ software package (version 2.25) [64]. All the BLASTn parameters were kept default except for the word size. Parsing of the blast results was carried out in a pipeline using custom perl scripts. BLASTn searches for word sizes from 8 to 11 were executed for the CNS specific lncRNA and rCNS datasets against the phastCons elements. The NCNS/rNCNS and Ensembl/ rEnsembl datasets were queried against zPHS at word size 11. Four parameters from the BLASTn search results were considered in the ROC analyses: query coverage (fraction of a lncRNA which is aligned to a phast-Cons element), alignment length (the length of the alignment including the gaps inserted), percentage identity (number of identical base matches between the query and the subject sequences) and e-value (a score which defines the probability of an alignment not being random in nature). The alignments of the lncRNAs





(CNS/NCNS/Ensembl) against the zPHS were taken as the true positive dataset while those from the randomized datasets (rCNS/rNCNS/rEnsembl) were considered to be the false positive set. The ROCR package in R environment was utilized to build the receiver operating characteristic (ROC) curve of false positive against true positive values for each parameter [65]. ROC curves for the e-value parameter in the plots show the reciprocal of the e-value (1/e-value) because plotting the e-value produced curves sensibly skewed below the diagonal line. Each alignment produced from the BLASTn search of the CNS dataset against zebrafish was also considered for structural conservation analysis. SISSIz program [66] was used to randomize each alignment 100 times using a dinucleotide model (SISSIz –simulate –tstv -n 100) to generate a randomized alignment dataset to measure the structural conservation (srCNS). The alignments of the CNS and srCNS datasets were checked for structural conservation with the RNAz 2.0 software (default parameters) [41]. To build ROC curves we used the following parameters from the RNAz output: ratio of pairwise identity by sequence conservation index, Z score and P values (1/P value). The parameters from the original alignments were considered to be true positive while those from the randomized alignments were considered to be the false positive. ROC curves of the false positive against the true positive were plotted for each the parameter.

Genomic features identification and enrichment analysis

The predicted conserved mouse lncRNAs were obtained after using the e-value and query alignment length thresholds as defined by the ROC curves in order to have less than 0.05% false positives passing it. The conserved lncRNAs (named cCNS, cNCNS, cEnsembl) and their respective zPHS elements sharing sequence similarity (named zCNS, zNCNS, zEnsembl) were back mapped to the mouse and zebrafish genomes (mm9 and zv9) respectively using BLASTn with default parameters but -culling limit = 1. The mapped coordinates of each mouse lncRNA and zebrafish conserved element were used to retrieve overlapping genes, transcripts, exons, and the closest flanking protein coding genes in a window up to 1 mb using custom perl scripts which use the Ensembl core modules API [67] and programmatic access to the Ensembl databases version 62. DAVID gene annotation tool was used for the GO term enrichment and tissue expression enrichment analyses for the protein-coding genes flanking and overlapping the conserved elements using the whole transcriptome as universe [46]. An EASE score of 0.05 [48] was used as a cut-off for the enrichment analysis. Sequences of ultraconserved elements [52,56] were mapped on the mouse genome using BLASTn (-task blastn -culling_limit 1) with default parameters. The coordinates of the mapped elements on the mouse genome were checked for overlap with conserved mouse lncRNAs using overlapBed program from the BEDTools package [68] (version 2.14.2) with default parameters. In all the overlap analyses performed we have considered sufficient an overlap of at least 1 bp between the conserved element and the specific feature considered.

Identification of orthologs between mouse and zebrafish and mapping of ESTs in the region of conservation

Zebrafish and mouse homology information were downloaded from BioMart [60] based on EnsEMBL version 62. We collected all the Ensembl genes mapped in intervals up to 2 Mb (1 Mb up and down-stream) around each conserved element in both the genomes. For each element we looked for genes considered evolutionary related (classified as ortholog one to one, ortholog one to many or ortholog many to many) in Ensembl Compara [69]. Conserved elements were considered syntenic if showing at least one evolutionary related gene in the given interval for the species considered. The analysis was performed individually on all lncRNAs stemming from the cCNS, cNCNS and cEnsembl datasets. The EST coordinates for mouse and zebrafish were downloaded from UCSC databases on 14th September 2011 [70,71]. The region of sequence conservation in the mouse lncRNAs (cCNS/cNCNS/cEnsEMBL) were checked for the overlap with a reported EST on the mouse genome. The same process was repeated on the zPHS conserved fragments (zCNS/zNCNS/zEnsEMBL) with respect to zebrafish ESTs. The EnsEMBL genome browser was used to generate the images for the conserved zPHS region 113364 [72] and its corresponding lncRNA in mouse as well as the region in human showing sequence homology to 113364.

Mapping of RNAseq data and read count on conserved regions

The zebrafish paired end RNAseg data from 7 developmental stages and stickleback paired end RNAseq from 9 tissues were downloaded from the European Nucleotide Archive in the fastq format (Accessions: SRP012923 and SRP009426). The raw reads were mapped to the zebrafish and stickleback genome using Tophat 2.0.4 [73] (tophat -p -o -G) and overlap associations for the conserved regions were calculated using custom perl scripts and the coverageBed (coverageBed -split -aBam -b) program from the BEDTools package [68] (version 2.14.2). Zebrafish sequences of the conserved elements were mapped on the stickleback genome using BLASTn (-task blastn -culling_ limit 1) with default parameters and all the regions were mapped with a minimum percentage identity of 70%. Random regions (~1,200) on the zebrafish genome were selected using the shuffleBed (shuffleBed -i -g) program from the BEDTools package. Overlap associations for the random regions were calculated in the same way as that for conserved regions.

Additional material

Additional File 1: ROC curves of CNS dataset at word size 8-10. ROC curves for query coverage (QCoverage), percentage identity (Pldentity), query alignment length (QAlength) and e-value (EValue) for the CNS dataset at word size A) 8 B) 9, C) 10. The cut-off for a parameter is defined as the point of steep incline in the true positive rate as compared to the false positive rate. The significant cut-off defined in the present analysis are indicated by arrows.

Additional File 2: Conserved IncRNAs. Results of BLASTn search and annotations concerning the genomic locations and proximal proteincoding genes of predicted conserved elements in mouse and zebrafish. The excel file contains three data sheets. The data is from CNS, NCNS and Ensembl datasets. The data includes BLASTn search results of the mouse IncRNAs against zebrafish phastCons elements. For each predicted conserved element in zebrafish and mouse information pertaining to its genomic location and overlapping/flanking gene features is provided as obtained from the Ensembl databases. If an element overlaps more than one feature each overlap is reported in a separate row.

Additional File 3: ROC curve for structural conservation of CNS IncRNAs dataset. A) Pairwise identity/Sequence conservation index (AUC 0.74), B) Z score (AUC 0.47) and C) inverse *P*-value (AUC 0.57) for the **mouse CNS constrained IncRNAs** against the zebrafish phastcons elements.

Additional File 4: Function and expression of proteins flanking the conserved elements of the CNS and NCNS dataset. GO biological process term (level 5) enrichment of **A**) flanking proteins of conserved elements in zebrafish **B**) flanking proteins of conserved elements in mouse for the CNS and NCNS dataset. Tissue enrichment of **C**) flanking proteins of putative conserved elements in zebrafish **D**) flanking proteins of conserved elements in conserved elements in mouse for the CNS and NCNS dataset. A, **B**, **C**, **D**: GO terms and tissue of expression are listed only if they are significantly over-represented according to the EASE score. Grey bars indicate the percentages of genes associated to the respective functional classes from the group of genes flanking the identified conserved elements. Black bars indicate the percentages from the entire transcriptome of the given species.

Additional File 5: RNAseq data overlap on conserved zebrafish elements. The figure depicts the percentage of conserved elements in the zebrafish genome which show overlap with > 1, > 25 and > 1000 short reads (coming from RNAseq of zebrafish development stages) as compared against a set of random elements in the fish genome.

List of abbreviations

IncRNA: long noncoding RNA; EST: expressed sequence tag; UCE: ultraconserved element; GO: gene ontology; CNS: central nervous system specific; NCNS non-central nervous system specific; ROC: receiver operating characteristic; AUC: area under curve; FDR: false discovery rate.

Authors' contributions

RS and SB conceived the study. SB designed and executed the analysis, interpreted the results and drafted the manuscript. RS supervised the study, participated in the design and execution of the analysis, the interpretation of results and helped to draft the manuscript. FM participated in the design of the analysis and interpretation of results and helped to draft the manuscript. All authors read and approved the final manuscript.

Competing interests

The authors declare that they have no competing interests.

Acknowledgements

This work is part of the PhD project of SB funded by the Stazione Zoologica Anton Dohrn (SZN) under the affiliated research centre PhD program of The Open University, Milton Keynes, UK. The work was also supported by the Seventh Framework Program of the European Commission (DOPAMINET project to RS and FM). The authors would like to acknowledge Stefano Gustincich, Piero Carninci and Elia Stupka for helpful discussions, Graziano Fiorito for support and three anonymous reviewers for their valuable comments and suggestions. This article has been published as part of *BMC Bioinformatics* Volume 14 Supplement 7, 2013: **Italian Society of Bioinformatics (BITS): Annual Meeting 2012.** The full contents of the supplement are available online at http://www.biomedcentral.com/bmcbioinformatics/supplements/14/S7

Declarations

The publication costs for this article were funded by the corresponding author's institution.

Author details

¹Laboratory of Animal Physiology and Evolution, Stazione Zoologica Anton Dohrn, Villa Comunale, 80121, Naples, Italy. ²Centre for Rare Diseases and Personalized Medicine, School of Clinical and Experimental Medicine, College of Medical and Dental Sciences, University of Birmingham, Birmingham, UK.

Published: 22 April 2013

References

 Ota T, Suzuki Y, Nishikawa T, et al: Complete sequencing and characterization of 21,243 full-length human cDNAs. Nat Genet 2004, 36:40-45. Page 15 of 17

- Okazaki Y, Furuno M, Kasukawa T, et al: Analysis of the mouse transcriptome based on functional annotation of 60,770 full-length cDNAs. Nature 2002, 420:563-573.
- 3. Carninci P, Kasukawa T, Katayama S, *et al*: **The transcriptional landscape of the mammalian genome**. *Science* 2005, **309**:1559-1563.
- Brown CJ, Hendrich BD, Rupert JL, Lafrenière RG, Xing Y, Lawrence J, Willard HF: The human XIST gene: analysis of a 17 kb inactive X-specific RNA that contains conserved repeats and is highly localized within the nucleus. *Cell* 1992, 71:527-542.
- Zearfoss NR, Chan AP, Kloc M, Allen LH, Etkin LD: Identification of new Xlsirt family members in the Xenopus laevis oocyte. *Mech Dev* 2003, 120:503-509.
- Dinger ME, Amaral PP, Mercer TR, Pang KC, Bruce SJ, Gardiner BB, Askarian-Amiri ME, Ru K, Soldà G, Simons C, Sunkin SM, Crowe ML, Grimmond SM, Perkins AC, Mattick JS: Long noncoding RNAs in mouse embryonic stem cell pluripotency and differentiation. *Genome Res* 2008, 18:1433-1445.
- Rinn JL, Kertesz M, Wang JK, Squazzo SL, Xu X, Brugmann SA, Goodnough LH, Helms JA, Farnham PJ, Segal E, Chang HY: Functional demarcation of active and silent chromatin domains in human HOX loci by noncoding RNAs. *Cell* 2007, 129:1311-1323.
- Kino T, Hurt DE, Ichijo T, Nader N, Chrousos GP: Noncoding RNA gas5 is a growth arrest- and starvation-associated repressor of the glucocorticoid receptor. *Sci Signal* 2010, 3:ra8.
- Hutchinson JN, Ensminger AW, Clemson CM, Lynch CR, Lawrence JB, Chess A: A screen for nuclear transcripts identifies two linked noncoding RNAs associated with SC35 splicing domains. *BMC Genomics* 2007, 8:39.
- Zhou Y, Zhong Y, Wang Y, Zhang X, Batista DL, Gejman R, Ansell PJ, Zhao J, Weng C, Klibanski A: Activation of p53 by MEG3 non-coding RNA. J Biol Chem 2007, 282:24731-24742.
- Pang KC, Dinger ME, Mercer TR, Malquori L, Grimmond SM, Chen W, Mattick JS: Genome-wide identification of long noncoding RNAs in CD8+ T cells. J Immunol 2009, 182:7738-7748.
- Silva JM, Perez DS, Pritchett JR, Halling ML, Tang H, Smith DI: Identification of long stress-induced non-coding transcripts that have altered expression in cancer. *Genomics* 2010, 95:355-362.
- Ørom UA, Derrien T, Beringer M, Gumireddy K, Gardini A, Bussotti G, Lai F, Zytnicki M, Notredame C, Huang Q, Guigo R, Shiekhattar R: Long noncoding RNAs with enhancer-like function in human cells. *Cell* 2010, 143:46-58.
- Mercer TR, Qureshi IA, Gokhan S, Dinger ME, Li G, Mattick JS, Mehler MF: Long noncoding RNAs in neuronal-glial fate specification and oligodendrocyte lineage maturation. *BMC Neurosci* 2010, 11:14.
- Wang Z, Gerstein M, Snyder M: RNA-Seq: a revolutionary tool for transcriptomics. Nat Rev Genet 2009, 10:57-63.
- Lin M, Pedrosa E, Shah A, Hrabovsky A, Maqbool S, Zheng D, Lachman HM: RNA-Seq of human neurons derived from iPS cells reveals candidate long non-coding RNAs involved in neurogenesis and neuropsychiatric disorders. *PLoS ONE* 2011, 6:e23356.
- Pauli A, Valen E, Lin MF, Garber M, Vastenhouw NL, Levin JZ, Fan L, Sandelin A, Rinn JL, Regev A, Schier AF: Systematic identification of long non-coding RNAs expressed during zebrafish embryogenesis. *Genome Research* 2011.
- Ulitsky I, Shkumatava A, Jan CH, Sive H, Bartel DP: Conserved Function of lincRNAs in Vertebrate Embryonic Development despite Rapid Sequence Evolution. *Cell* 2011, 147:1537-1550.
- Nam J-W, Bartel D: Long non-coding RNAs in C. elegans. Genome research 2012.
- Amaral PP, Clark MB, Gascoigne DK, Dinger ME, Mattick JS: IncRNAdb: a reference database for long noncoding RNAs. Nucleic Acids Res 2011, 39: D146-151.
- 21. Khachane AN, Harrison PM: Mining mammalian transcript data for functional long non-coding RNAs. *PLoS ONE* 2010, **5**:e10316.
- Jia H, Osak M, Bogu GK, Stanton LW, Johnson R, Lipovich L: Genome-wide computational identification and manual annotation of human long noncoding RNA genes. *RNA* 2010, 16:1478-1487.
- 23. Arrial R, Togawa R, Brigido M: Screening non-coding RNAs in transcriptomes from neglected species using PORTRAIT: case study of the pathogenic fungus Paracoccidioides brasiliensis. *BMC Bioinformatics* 2009, **10**:239.
- Kong L, Zhang Y, Ye Z-Q, Liu X-Q, Zhao S-Q, Wei L, Gao G: CPC: assess the protein-coding potential of transcripts using sequence features and support vector machine. *Nucleic Acids Res* 2007, 35:W345-349.

- Cabili MN, Trapnell C, Goff L, Koziol M, Tazon-Vega B, Regev A, Rinn JL: Integrative annotation of human large intergenic noncoding RNAs reveals global properties and specific subclasses. *Genes & Development* 2011, 25:1915-1927.
- Chodroff RA, Goodstadt L, Sirey TM, Oliver PL, Davies KE, Green ED, Molnár Z, Ponting CP: Long noncoding RNA genes: conservation of sequence and brain expression among diverse amniotes. *Genome Biol* 2010, 11:R72.
- Hubbard SJ, Grafham DV, Beattie KJ, Overton IM, McLaren SR, Croning MDR, Boardman PE, Bonfield JK, Burnside J, Davies RM, Farrell ER, Francis MD, Griffiths-Jones S, Humphray SJ, Hyland C, Scott CE, Tang H, Taylor RG, Tickle C, Brown WRA, Birney E, Rogers J, Wilson SA: Transcriptome analysis for the chicken based on 19,626 finished cDNA sequences and 485,337 expressed sequence tags. *Genome Res* 2005, 15:174-183.
- Duret L, Chureau C, Samain S, Weissenbach J, Avner P: The Xist RNA gene evolved in eutherians by pseudogenization of a protein-coding gene. *Science* 2006, 312:1653-1655.
- Pollard KS, Salama SR, Lambert N, Lambot M-A, Coppens S, Pedersen JS, Katzman S, King B, Onodera C, Siepel A, Kern AD, Dehay C, Igel H, Ares M Jr, Vanderhaeghen P, Haussler D: An RNA gene expressed during cortical development evolved rapidly in humans. *Nature* 2006, 443:167-172.
- Amaral PP, Neyt C, Wilkins SJ, Askarian-Amiri ME, Sunkin SM, Perkins AC, Mattick JS: Complex architecture and regulated expression of the Sox2ot locus during vertebrate development. *RNA* 2009, 15:2013-2027.
- 31. Ponjavic J, Oliver PL, Lunter G, Ponting CP: Genomic and transcriptional co-localization of protein-coding and long non-coding RNA pairs in the developing brain. *PLoS Genet* 2009, **5**:e1000617.
- Dinger ME, Gascoigne DK, Mattick JS: The evolution of RNAs with multiple functions. *Biochimie* 2011, 93:2013-2018.
- Ingolia NT, Lareau LF, Weissman JS: Ribosome profiling of mouse embryonic stem cells reveals the complexity and dynamics of mammalian proteomes. *Cell* 2011, 147:789-802.
- 34. Bánfai B, Jia H, Khatun J, Wood E, Risk B, Gundling WE Jr, Kundaje A, Gunawardena HP, Yu Y, Xie L, Krajewski K, Strahl BD, Chen X, Bickel P, Giddings MC, Brown JB, Lipovich L: Long noncoding RNAs are rarely translated in two human cell lines. *Genome Res* 2012, 22:1646-1657.
- Pollard KS, Hubisz MJ, Rosenbloom KR, Siepel A: Detection of nonneutral substitution rates on mammalian phylogenies. *Genome Res* 2010, 20:110-121.
 Instruction of the construction of the construction
- 36. [http://www.ensembl.org/info/docs/genebuild/ncrna.html].
- 37. Flicek P, Amode MR, Barrell D, Beal K, Brent S, Chen Y, Clapham P, Coates G, Fairley S, Fitzgerald S, Gordon L, Hendrix M, Hourlier T, Johnson N, Kähäri A, Keefe D, Keenan S, Kinsella R, Kokocinski F, Kulesha E, Larsson P, Longden I, McLaren W, Overduin B, Pritchard B, Riat HS, Rios D, Ritchie GRS, Ruffier M, Schuster M, Sobral D, Spudich G, Tang YA, Trevanion S, Vandrovcova J, Vilella AJ, White S, Wilder SP, Zadissa A, Zamora J, Aken BL, Birney E, Cunningham F, Dunham I, Durbin R, Fernández-Suarez XM, Herrero J, Hubbard TJP, Parker A, Proctor G, Vogel J, Searle SMJ: Ensembl 2011. Nucleic Acids Res 2011, 39:D800-806.
- Fawcett T: ROC graphs: Notes and practical considerations for researchers. Machine Learning 2004, 31:1-38.
- 39. Kelley DR, Rinn JL: Transposable elements reveal a stem cell specific class of long noncoding RNAs. *Genome Biology* 2012, **13**:R107.
- Carrieri C, Cimatti L, Biagioli M, Beugnet A, Zucchelli S, Fedele S, Pesce E, Ferrer I, Collavin L, Santoro C, Forrest ARR, Carninci P, Biffo S, Stupka E, Gustincich S: Long non-coding antisense RNA controls Uchl1 translation through an embedded SINEB2 repeat. *Nature* 2012, 491:454-457.
- 41. Washietl S, Hofacker IL, Stadler PF: Fast and reliable prediction of noncoding RNAs. Proc Natl Acad Sci USA 2005, 102:2454-2459.
- Seemann SE, Gilchrist MJ, Hofacker IL, Stadler PF, Gorodkin J: Detection of RNA structures in porcine EST data and related mammals. *BMC Genomics* 2007, 8:316.
- McCutcheon JP, Eddy SR: Computational identification of non-coding RNAs in Saccharomyces cerevisiae by comparative genomics. *Nucleic Acids Res* 2003, 31:4119-4128.
- 44. Katayama S, Tomaru Y, Kasukawa T, Waki K, Nakanishi M, Nakamura M, Nishida H, Yap CC, Suzuki M, Kawai J, Suzuki H, Carninci P, Hayashizaki Y, Wells C, Frith M, Ravasi T, Pang KC, Hallinan J, Mattick J, Hume DA, Lipovich L, Batalov S, Engström PG, Mizuno Y, Faghihi MA, Sandelin A, Chalk AM, Mottagui-Tabar S, Liang Z, Lenhard B, Wahlestedt C: Antisense transcription in the mammalian transcriptome. *Science* 2005, 309:1564-1566.

- Kim T-K, Hemberg M, Gray JM, Costa AM, Bear DM, Wu J, Harmin DA, Laptewicz M, Barbara-Haley K, Kuersten S, Markenscoff-Papadimitriou E, Kuhl D, Bito H, Worley PF, Kreiman G, Greenberg ME: Widespread transcription at neuronal activity-regulated enhancers. *Nature* 2010, 465:182-187.
- Huang DW, Sherman BT, Lempicki RA: Bioinformatics enrichment tools: paths toward the comprehensive functional analysis of large gene lists. *Nucleic Acids Res* 2009, 37:1-13.
- Huang DW, Sherman BT, Lempicki RA: Systematic and integrative analysis of large gene lists using DAVID bioinformatics resources. *Nat Protoc* 2009, 4:44-57.
- Hosack DA, Dennis G Jr, Sherman BT, Lane HC, Lempicki RA: Identifying biological themes within lists of genes with EASE. *Genome Biol* 2003, 4: R70.
- Mercer TR, Dinger ME, Sunkin SM, Mehler MF, Mattick JS: Specific expression of long noncoding RNAs in the mouse brain. Proc Natl Acad Sci USA 2008, 105:716-721.
- 50. Qureshi IA, Mattick JS, Mehler MF: Long non-coding RNAs in nervous system function and disease. *Brain Res* 2010, 1338:20-35.
- Sheik Mohamed J, Gaughwin PM, Lim B, Robson P, Lipovich L: Conserved long noncoding RNAs transcriptionally regulated by Oct4 and Nanog modulate pluripotency in mouse embryonic stem cells. *RNA* 2010, 16:324-337.
- Bejerano G, Pheasant M, Makunin I, Stephen S, Kent WJ, Mattick JS, Haussler D: Ultraconserved elements in the human genome. *Science* 2004, 304:1321-1325.
- 53. Calin GA, Liu C, Ferracin M, Hyslop T, Spizzo R, Sevignani C, Fabbri M, Cimmino A, Lee EJ, Wojcik SE, Shimizu M, Tili E, Rossi S, Taccioli C, Pichiorri F, Liu X, Zupo S, Herlea V, Gramantieri L, Lanza G, Alder H, Rassenti L, Volinia S, Schmittgen TD, Kipps TJ, Negrini M, Croce CM: Ultraconserved regions encoding ncRNAs are altered in human leukemias and carcinomas. *Cancer Cell* 2007, 12:215-229.
- Licastro D, Gennarino VA, Petrera F, Sanges R, Banfi S, Stupka E: Promiscuity of enhancer, coding and non-coding transcription functions in ultraconserved elements. *BMC Genomics* 2010, 11:151.
- Pennacchio LA, Ahituv N, Moses AM, Prabhakar S, Nobrega MA, Shoukry M, Minovitsky S, Dubchak I, Holt A, Lewis KD, Plajzer-Frick I, Akiyama J, De Val S, Afzal V, Black BL, Couronne O, Eisen MB, Visel A, Rubin EM: In vivo enhancer analysis of human conserved non-coding sequences. *Nature* 2006, 444:499-502.
- Sakuraba Y, Kimura T, Masuya H, Noguchi H, Sezutsu H, Takahasi KR, Toyoda A, Fukumura R, Murata T, Sakaki Y, Yamamura M, Wakana S, Noda T, Shiroishi T, Gondo Y: Identification and characterization of new long conserved noncoding sequences in vertebrates. *Mamm Genome* 2008, 19:703-712.
- Guttman M, Garber M, Levin JZ, Donaghey J, Robinson J, Adiconis X, Fan L, Koziol MJ, Gnirke A, Nusbaum C, Rinn JL, Lander ES, Regev A: Ab initio reconstruction of cell type-specific transcriptomes in mouse reveals the conserved multi-exonic structure of lincRNAs. *Nat Biotech* 2010, 28:503-510.
- Hui L, Ji C, Hui B, Lv T, Ha X, Yang J, Cai W: The oncoprotein LMO3 interacts with calcium- and integrin-binding protein CIB. Brain Res 2009, 1265:24-29.
- Aoyama M, Ozaki T, Inuzuka H, Tomotsune D, Hirato J, Okamoto Y, Tokita H, Ohira M, Nakagawara A: LMO3 interacts with neuronal transcription factor, HEN2, and acts as an oncogene in neuroblastoma. *Cancer Res* 2005, 65:4587-4597.
- Haider S, Ballester B, Smedley D, Zhang J, Rice P, Kasprzyk A: BioMart Central Portal–unified access to biological data. Nucleic Acids Res 2009, 37:W23-27.
- 61. Eddy SR: Accelerated Profile HMM Searches. PLoS Comput Biol 2011, 7: e1002195.
- Dreszer TR, Karolchik D, Zweig AS, Hinrichs AS, Raney BJ, Kuhn RM, Meyer LR, Wong M, Sloan CA, Rosenbloom KR, Roe G, Rhead B, Pohl A, Malladi VS, Li CH, Learned K, Kirkup V, Hsu F, Harte RA, Guruvadoo L, Goldman M, Giardine BM, Fujita PA, Diekhans M, Cline MS, Clawson H, Barber GP, Haussler D, James Kent W: The UCSC Genome Browser database: extensions and updates 2011. Nucleic Acids Res 2012, 40: D918-923.
- 63. [http://www.genome.ucsc.edu/cgi-bin/hgLiftOver].

- Camacho C, Coulouris G, Avagyan V, Ma N, Papadopoulos J, Bealer K, Madden TL: BLAST+: architecture and applications. *BMC Bioinformatics* 2009, 10:421.
- Sing T, Sander O, Beerenwinkel N, Lengauer T: ROCR: visualizing classifier performance in R. Bioinformatics 2005, 21:3940-3941.
- 66. Gesell T, Washietl S: Dinucleotide controlled null models for comparative RNA gene prediction. *BMC Bioinformatics* 2008, **9**:248.
- 67. Flicek P, Aken BL, Ballester B, Beal K, Bragin E, Brent S, Chen Y, Clapham P, Coates G, Fairley S, Fitzgerald S, Fernandez-Banet J, Gordon L, Gräf S, Haider S, Hammond M, Howe K, Jenkinson A, Johnson N, Kähäri A, Keefe D, Keenan S, Kinsella R, Kokocinski F, Koscielny G, Kulesha E, Lawson D, Longden I, Massingham T, McLaren W, Megy K, Overduin B, Pritchard B, Rios D, Ruffier M, Schuster M, Slater G, Smedley D, Spudich G, Tang YA, Trevanion S, Vilella A, Vogel J, White S, Wilder SP, Zadissa A, Birney E, Cunningham F, Dunham I, Durbin R, Fernández-Suarez XM, Herrero J, Hubbard TJP, Parker A, Proctor G, Smith J, Searle SMJ: Ensembl's 10th year. Nucleic Acids Res 2010, 38:D557-562.
- Quinlan AR, Hall IM: BEDTools: a flexible suite of utilities for comparing genomic features. *Bioinformatics* 2010, 26:841-842.
- Vilella AJ, Severin J, Ureta-Vidal A, Heng L, Durbin R, Birney E: EnsemblCompara GeneTrees: Complete, duplication-aware phylogenetic trees in vertebrates. *Genome Res* 2009, 19:327-335.
- 70. [http://hgdownload.cse.ucsc.edu/goldenPath/mm10/database/all_est.txt.gz].
- [http://hgdownload.cse.ucsc.edu/goldenPath/danRer7/database/all_est.txt.gz].
 Flicek P, Amode MR, Barrell D, Beal K, Brent S, Carvalho-Silva D, Clapham P, Coates G, Fairley S, Fitzgerald S, Gil L, Gordon L, Hendrix M, Hourlier T, Johnson N, Kähäri AK, Keefe D, Keenan S, Kinsella R, Komorowska M, Koscielny G, Kulesha E, Larsson P, Longden I, McLaren W, Muffato M, Overduin B, Pignatelli M, Pritchard B, Riat HS, Ritchie GRS, Ruffier M, Schuster M, Sobral D, Tang YA, Taylor K, Trevanion S, Vandrovcova J, White S, Wilson M, Wilder SP, Aken BL, Birney E, Cunningham F, Dunham I, Durbin R, Fernández-Suarez XM, Harrow J, Herrero J, Hubbard TJP, Parker A, Proctor G, Spudich G, Vogel J, Yates A, Zadissa A, Searle SMJ: Ensembl 2012. Nucleic Acids Res 2012, 40:D84-90.
- 73. Trapnell C, Pachter L, Salzberg SL: **TopHat: discovering splice junctions** with RNA-Seq. *Bioinformatics* 2009, **25**:1105-1111.

doi:10.1186/1471-2105-14-S7-S14

Cite this article as: Basu *et al.*: Examples of sequence conservation analyses capture a subset of mouse long non-coding RNAs sharing homology with fish conserved genomic elements. *BMC Bioinformatics* 2013 14(Suppl 7):S14.

Submit your next manuscript to BioMed Central and take full advantage of:

- Convenient online submission
- Thorough peer review
- No space constraints or color figure charges
- Immediate publication on acceptance
- Inclusion in PubMed, CAS, Scopus and Google Scholar
- Research which is freely available for redistribution

() BioMed Central

Submit your manuscript at www.biomedcentral.com/submit