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Assessing Recent Selection and Functionality at Long Non-Coding RNA Loci in the Mouse Genome

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Abstract

Long non-coding RNAs (lncRNAs) are one of the most intensively studied groups of non-coding elements. Debate continues over what proportion of lncRNAs are functional or merely represent transcriptional noise. While characterisation of individual lncRNAs has identified ~200 functional loci across the Eukarya, general surveys have found only modest or no evidence of long-term evolutionary conservation. Although this lack of conservation suggests that most lncRNAs are non-functional, the possibility remains that some represent recent evolutionary innovations. We examine recent selection pressures acting on lncRNAs in mouse populations. We compare patterns of within-species nucleotide variation at ~10,000 lncRNA loci in a cohort of the wild house mouse, \textit{M. m. castaneus}, with between-species nucleotide divergence from the rat (\textit{Rattus norvegicus}). Loci under selective constraint are expected to show reduced nucleotide diversity and divergence. We find limited evidence of sequence conservation compared to putatively neutrally evolving ancestral repeats (ARs). Comparisons of sequence diversity and divergence between ARs, protein-coding (PC) exons and lncRNAs, and the associated flanking regions, show weak, but significantly lower levels of sequence diversity and divergence at lncRNAs compared to ARs. lncRNAs conserved deep in the vertebrate phylogeny show lower within-species sequence diversity than lncRNAs in general. A set of 74 functionally characterised lncRNAs show levels of diversity and divergence comparable to PC exons, suggesting that these lncRNAs are under substantial selective constraints. Our results suggest that, in mouse populations, most lncRNA loci evolve at rates similar to ARs, while older lncRNAs tend to show signals of selection similar to PC genes.

Keywords: long non-coding RNAs, selection, genomics, functionality
Introduction

Non-coding RNAs (ncRNAs), including tRNAs and rRNAs, are well established as key functional elements of the cellular machinery (Mattick & Makunin 2006). Recently, large-scale transcriptomics studies have revealed that a high proportion of the eukaryotic genome is transcribed, and this has led to the proposal of new classes of ncRNAs and to the addition of many new members to already existing classes. One of the most interesting classes are the long non-coding RNAs (lncRNAs) (Okazaki et al. 2002; Carninci et al. 2005; Mattick & Makunin 2006; Mercer et al. 2009), which are typically defined as transcribed RNA fragments exceeding a length threshold of 200 base pairs (bp) (Amaral et al. 2011; Brown et al. 2012; Kapusta et al. 2013). Many lncRNAs have been shown to be important regulators of gene expression (Ilik and Akhtar 2009; Rinn and Chang 2012). Increasing numbers have been predicted in various genomes, including a recent estimate of ~58,000 lncRNA loci in the human genome (Iyer et al. 2015). More recently, lncRNAs have been the topic of much debate concerning their functionality.

Large sets of lncRNAs and other non-coding transcripts have been identified by high-throughput genomic methods. Thousands of lncRNAs have been identified and annotated using methods such as DNA sequence tagging that indicate 5’ and 3’ transcript boundaries (Okazaki et al. 2002; Carninci et al. 2005; Ponjavic & Ponting 2007), “chromatin state maps” that reveal regions of chromatin suggestive of active transcription (Guttman et al. 2009) and, more recently, RNA sequencing (RNA-seq) to directly investigate the transcriptome (Necsulea et al. 2014). To assess the non-coding status of a region of interest, the potential for a DNA sequence to encode a protein can be assessed by a variety of algorithms (Lin et al. 2008). One example is the codon substitution frequency (CSF) metric, which assesses protein-coding potential by examining the relative frequency of synonymous and non-synonymous codon substitutions (Lin et al. 2007).

Although little is known about the function for most identified lncRNAs, several individual loci are now known to be important in the control of gene expression, suggesting that they could
constitute an important set of regulatory elements (Rinn & Chang 2012). One example is that of the
Air locus in mice (Antisense Igf2r RNA), which is involved in silencing the paternal Igf2r allele in cis. Air also silences the paternal alleles of other, nearby genes (Sleutels et al. 2002; Nagano et al.
2008; Latos et al. 2012). The human homolog of Air, AIRN, was found to have conserved function
(Yotova et al. 2008). Another well-known example is Xist, which is involved in X-chromosome
inactivation in mammals by localising to multiple regions on the X chromosome and recruiting
other molecules that repress transcription (Nesterova et al. 2001; Engreitz et al. 2013). Several other
examples are detailed in Table 1 of Ilik and Akhtar (2009) and in the online IncRNA database
(Amaral et al. 2011; Quek et al. 2014).

The involvement of individual IncRNAs in gene regulation leads to the question of the
relative importance of IncRNAs evolution and protein-coding gene evolution to adaptation
2012; Kutter et al. 2012; Kapusta et al. 2013). An extreme view proposes that IncRNAs represent
mostly “transcriptional noise” from an inherently stochastic transcription machinery (Ponjavic et al.
2007; Doolittle 2012). Efforts to study functionality have mostly consisted either of the detailed
characterisation of individual IncRNAs or surveys of large sets of IncRNAs in order to search for
evolutionary signals indicative of function. While initial large-scale studies found little evidence of
evolutionary constraint (Okazaki et al. 2002; Wang et al. 2004; Carninci et al. 2005; Maeda et al.
2006; Ponjavic et al. 2007; Marques & Ponting 2009; Guttman et al. 2009; Derrien et al. 2012),
evidence is mounting that at least some IncRNAs are conserved (Sleutels et al. 2002; Espinoza et al.

Early work on ~15,800 FANTOM 2 ncRNA transcripts identified in mice found they did not
differ in level of sequence conservation compared to a putatively neutrally evolving set of control
sequences (Wang et al. 2004). In further analysis of the most stringently annotated set of IncRNAs
from the FANTOM 2 and 3 project datasets (Okazaki et al. 2002; Carninci et al. 2005; Maeda et al.
2006; Ponjavic et al. 2007) substitution rates were found to be depressed by ~10% between mouse and rat, relative to putatively neutrally evolving AR sequences (Ponjavic et al. 2007). Ponjavic et al. (2007) also found conservation in both the sequence of the ncRNAs themselves and in their promoter regions (taken to be the 400 nt immediately upstream of the ncRNA transcriptional start site). Similarly, analysis of a novel set of lncRNAs identified by Guttman and colleagues (Guttman et al. 2009) revealed that both the sequence of the lncRNAs and promoter regions were conserved compared to random genomic fragments. Taking advantage of both the FANTOM and Guttman et al. datasets, Marques and Ponting (2009) found that substitution rates were depressed by ~10-12% relative to ARs. More recently, a study of ~9,000 human lncRNA loci from the GENCODE consortium showed higher PhastCons scores than ARs, but conservation was still substantially lower than protein-coding (PC) loci (Derrien et al. 2012). Furthermore, derived allele frequencies in a human polymorphism dataset were significantly lower at lncRNA loci than for random intergenic regions, though differences were small (Necsulea et al. 2014). However, results appear to vary across lineages: ~35.8% of mutations within Drosophila melanogaster lncRNAs have been found to be weakly deleterious, while in Homo sapiens all mutations have been estimated to be effectively neutral, an effect that could be attributed to differences in effective population sizes (Haerty & Ponting 2013).

Here, we test for evidence of recent selection on lncRNAs using a genome-wide polymorphism dataset from the wild house mouse species Mus musculus castaneus and a newly described, large dataset of lncRNAs identified in the mouse genome (Necsulea et al. 2014). The house mouse represents an excellent study system for evolutionary analysis, due to its large effective population size ($N_e$) in the wild, which is expected to reduce the effects of genetic drift and increase the efficacy of selection compared to species with lower $N_e$, such as humans (Charlesworth 2009; Phifer-Rixey et al. 2012; Haerty & Ponting 2013). Furthermore, extensive functional genomics and genetic resources are available for the mouse, which has become an important model
species for functional investigation of lncRNAs.

We assess the evidence for recent purifying selection at lncRNA loci and compare the extent of selection at lncRNAs compares with functional loci known to be under evolutionary constraints, such as PC genes. We examine patterns of diversity in regions flanking the focal loci to investigate selection at these loci. We also examine the extent of recent purifying selection within modern wild mouse populations at lncRNA loci of varying evolutionary ages and at lncRNAs expressed in different tissue types.

Material and Methods

lncRNA dataset

In this study, we focus on a set of 10,088 mouse lncRNAs from a recent large-scale evolutionary study (Necsulea et al. 2014). This dataset provides an estimate of the evolutionary age of lncRNA loci, allowing us to test for a relationship between within-species diversity and long-term evolutionary conservation. The evolutionary ages used here were determined by the phylogenetic distribution in species in which homologous sequence was detected and for which there was evidence of transcription at orthologous lncRNA loci (Necsulea et al. 2014). If homologous sequences and transcription evidence could be found across all amniotes, but not in the non-amniote species, then the age of the locus was set as the approximate age of the amniote lineage. This therefore represents a minimum estimate of the evolutionary age, because the reliable detection of transcription becomes difficult for genes that are not highly expressed and because homologous sequence identification is difficult for fast-evolving loci (Necsulea et al. 2014). Additionally, in the current dataset it is possible to divide lncRNAs depending on the tissue in which they are maximally expressed, as measured by reads per kilobase per million mapped reads (RPKM). Such a division might uncover selection acting on biologically important groups of lncRNAs. Finally, to partly control for potentially higher levels of conservation at lncRNAs near
protein coding genes, lncRNAs were also grouped into intronic or intergenic lncRNAs. Intergenic
lncRNAs are defined here as any lncRNA that did not overlap with an Ensembl annotation protein
coding gene.

We analysed sequence diversity in this set of lncRNAs using the genome sequences of 10
wild *M. m. castaneus* aligned to the NCBIM37 mm9 mouse reference genome (Halligan et al.
2013). Additionally, we examine nucleotide divergence between mouse and rat using the alignment
of the mm9 mouse reference genome to the rat (rn4, *Rattus norvegicus*) reference genome. These
alignments are available from the UCSC Genome Browser (Kent et al. 2002). We compare levels of
diversity and divergence at IncRNA loci with levels at putatively neutrally evolving ancestral repeat
(AR) loci and evolutionarily conserved protein-coding (PC) loci. We also investigate the patterns of
diversity and divergence in the genomic regions flanking these loci to determine to what extent
characteristic signals of selection are present at these loci.

To ensure that the sequences analysed are correctly assigned to the above groups, a number
of filtering steps were applied prior to analysis. The regions corresponding to the most divergent
lncRNAs from the mm9 – rn4 alignments were visually inspected to ensure that extreme divergence
was not due to unreliable alignments, and none were rejected. We note that the use of a single rat
genome as an outgroup does not allow for discrimination between diversity within rat and
divergence between mouse and rat, thus we likely over-estimate the extent of between-species
divergence. Some IncRNAs overlap with known protein-coding (PC) exons. To avoid confounding
diversity at IncRNA loci with diversity at neighboring protein-coding loci, a procedure for
removing known elements from the datasets was carried out. Overlapping elements were always
completely removed regardless of annotated strand. If neutrally evolving sites are considered in
flanking regions it is possible to determine the effect of selection at a focal locus on diversity at
linked neutral sites, either through genetic hitchhiking (Maynard-Smith & Haigh 1974) or through
background selection (Charlesworth et al. 1993). Both of these mechanisms are expected to lead to
diversity reductions at linked sites resulting in a dip around the selected site. Examining diversity in sequences surrounding a focal class of sites has previously been employed in other studies to identify signals of selection (Sattath et al. 2011; Hernandez et al. 2011; Halligan et al. 2013). To this end, sites that could potentially be under selection in the flanking regions of all focal loci (PC exons, lncRNAs and ARs) were removed as follows. At lncRNA loci, PC loci and their associated untranslated regions (UTRs) were removed, resulting in the complete removal of 51 lncRNA loci. Conserved non-coding elements (CNEs), PC exons and UTRs were removed from flanking regions of lncRNA loci. At AR loci, PC loci, associated UTR, and lncRNA loci were removed to ensure that ARs analysed were as far as possible neutrally evolving. CNE loci, lncRNAs, PC exons and UTRs were removed from the flanking regions of AR loci. At PC loci and their UTRs, overlapping lncRNA loci were removed. CNEs and lncRNA loci were removed from the flanking regions of PC loci. Loci that mapped to sex chromosomes (191 loci) were excluded from the analyses, because such loci are likely to have had different selective and mutational pressures (Charlesworth et al. 1987; Baines & Harr 2007; Kousathanas et al. 2014). This resulted in a dataset comprising 10,088 lncRNA loci from an initial set of 10,330.

To establish the level of evolutionary signal that can be expected from functional lncRNAs, a set of 108 well-characterised lncRNAs from the mouse was downloaded from the lncRNA database (Amaral et al. 2011; Quek et al. 2014). Since some of these loci may have been selected for further experiments based on their evolutionary conservation, we note that they are potentially biased towards more conserved sequence. We used BLAT (Kent 2002) with default settings to determine the genomic positions of exons in these loci. A single best hit for each locus was chosen on the basis of highest bit score and sequence identity. Many of the entries in the database represent alternative isoforms of the same locus. If the best hit locations of two or more isoforms overlapped the same region in the mouse genome, the longest isoform was kept and the others discarded from further analysis. The final set was manually checked and ambiguously defined and overlapping loci
removed. A total of 74 autosomal loci were recovered (Table S1, Supplementary Materials) and analysed as described below. Of these 74 loci of known function, the positions of 23 loci overlap with counterparts in the larger dataset of 10,088. The remaining 51 are not present in our selected dataset, generally due to the lack of detectable expression in the transcriptome dataset used by Necsulea et al. (2014) or because they could not be unambiguously assigned to one-to-one orthologous gene families across species. In some cases, these lncRNAs were filtered out of the original dataset due to overlap with annotated PC genes, or because they were classified as protein-coding based on their CSF score or based on the presence of sequence similarity with known protein and protein domains (Necsulea et al., 2014).

**Signatures of Selection**

As a putatively neutrally evolving standard against which to compare the lncRNA loci, a set of ARs was used. ARs are transposable elements inserted in the genome before the split between two lineages (in this case mouse and rat) that have remained at that location. While there are some ARs that show evidence of functionality, the available evidence suggests that the majority evolve at similar rates to that expected under neutrality (Mouse Genome Sequencing Consortium 2002; Chiaromonte et al. 2003; Lunter et al. 2006; Marques & Ponting 2009; Eöry et al. 2010; Kutter et al. 2012). Because the mutation rate, and therefore the rate of neutral evolution, varies across the genome, we examined closely linked ARs and focal loci. To this end, only AR loci within 10,000 base pairs (bp) up or downstream of a lncRNA locus were used in this study. Additionally, the set of all known PC exons and UTRs from the Ensembl 62 database was used as an example of loci believed to be under both negative and positive selection (Halligan et al. 2013).

Nucleotide diversity around the different genomic elements (lncRNAs, PC exons and ARs) was estimated with set of custom Python scripts from pooled, folded site frequency spectra (SFSs) across the loci. Estimating diversity and divergence from a pooled SFS across loci should reduce
the effect of variation in estimates that is due to short alignments. Nucleotide diversity ($\pi$),
divergence to rat ($d$), $\pi/d$, an estimate of diversity corrected for regional variation in the mutation
rate, and Tajima’s D statistics were calculated for all focal loci individually and for a series of 50
non-overlapping 1,000 nt windows upstream and downstream flanking regions of these loci
(IncRNAs, PC exons and ARs). CpG sites are known to be hypermutable in mammals and can thus
lead to biases in the estimates of substitution rates if some regions are more enriched for CpG sites.
Analyses were therefore carried out excluding CpG-prone sites (defined as any site either preceded
by a C or followed by a G). Examining diversity in sequences surrounding a focal class of sites has
previously been employed in other studies to identify signals of selection (Sattath et al. 2011;
Hernandez et al. 2011; Halligan et al. 2013), because both genetic hitchhiking (Maynard-Smith &
Haigh 1974) and background selection (Charlesworth et al. 1993) are expected to cause diversity
reductions at linked sites resulting in a characteristic “dip” around the selected site.

**Statistical Tests of Diversity and Divergence Estimates**

Formal tests of differences in conservation were performed using the estimates of
divergence and diversity calculated as described above. Divergence ratios ($d_{\text{locus}}/d_{\text{neutral}}$) measure the
extent of depression in nucleotide divergence at a locus compared to a putatively neutrally evolving
reference. These depressions were assessed by taking the ratio of divergence at IncRNA or PC loci
to the median value for a set of ARs, $d_{\text{AR}}$, i.e., $d_{\text{locus}}/d_{\text{AR}}$, where $d_{\text{locus}}$ is the median ($d$) from the
distribution of $d$ across all the focal loci (all other numbers reported below are estimates from a
pooled SFS across loci). Regression models were used to investigate the relationship between
evolutionary age and diversity. Two models were built for each of the diversity related statistics
(Tajima’s D, $\pi$ and $\pi/d$), one including a quadratic term and a linear term, and one including only a
linear term. The best-fitting models were chosen by F-tests comparing nested models. Non-
parametric 95% confidence intervals were obtained by 1,000 rounds of bootstrap sampling of the
loci, allowing an unpaired 2-tailed test of the degree of overlap between distributions of divergence
and diversity statistics between different groups of lncRNAs and ARs. Patterns of diversity in flanking regions around focal loci can be described by non-linear least squares models. Parameters were estimated for the equation \( \pi/d \approx A \times (1 - B \times \exp(-\text{abs}(x)/q)) \) where \( x \) is the distance from the focal locus. The term \( A \) estimates the neutral level of \( \pi/d \) as \( x \) tends towards infinity. The term \( B \) estimates the reduction in \( \pi/d \) when \( x = 0 \). Finally, \( q \) estimates the distance over which neutral levels of \( \pi/d \) is recovered (Hernandez et al. 2011; Halligan et al. 2013). This model was compared to a nested model including only the term \( A \) (i.e. where \( B = 0 \)) by an F-test which tests whether a model with a reduction in diversity closer to the focal loci is better than a model without such a reduction. All statistical tests and calculations were performed in R (R Development Core Team 2008).

Results

Summary statistics for the datasets (and subsets) analysed in this study are shown in Tables S2, S3 and S4. In total, 10,088 lncRNA loci (and 74 loci of known function) are considered in this study. These loci have been split on the basis of the estimated minimum evolutionary age, ranging from 370 million years (MY) to 12 MY, and on the basis of expression in different tissue types (Adkins et al. 2001; Springer et al. 2003; Necsulea et al. 2014). The “mouse” group is given an age of 12 MY, since evidence of transcription of these lncRNAs comes only from contemporary mouse populations and 12 MY is the earliest estimated split between mouse and rat (Adkins et al. 2001; Springer et al. 2003). However, RNA-seq data for these loci are not available for rat and these loci may not all be mouse-specific. The age groups differ in the number of lncRNAs they contain, ranging from 7,306 to 62 loci, but neither the average nor the median lengths of lncRNA loci vary substantially across the different age groups (Table S3). 423 loci in the dataset show evidence of transcription in primates only, although homologous sequences could be identified in the mouse. These loci are included in the group of all lncRNAs, but have been excluded from the analyses of different age groups in order to include only the loci from different age groups that are most likely to be active in the contemporary mouse genome. Of the 22 lncRNA loci of known function that also
occur in the larger dataset, 12 are described as mouse-specific in the annotation of the larger dataset. A further six loci are found across the Eutherian lineage and the remaining two loci across Tetrapods and across Mammals.

**Patterns of Diversity at Focal loci**

ARs show the highest level of diversity ($\pi = 0.897\%$), PC exons show the lowest level ($\pi = 0.351\%$), and the group of all lncRNAs shows a level of diversity slightly, though significantly, lower than ARs ($\pi = 0.800\%$) (Table 1). Similar patterns (highest in ARs, lowest in PC exons and an intermediate level in lncRNAs) hold for all measures of diversity and divergence ($\pi$, $d$, $\pi/d$ and Tajima’s D) (Table 1). For all lncRNA loci, unpaired 2-tailed tests indicate that there is a significant reduction for all diversity measures ($\pi$, $d$, $\pi/d$ and Tajima’s D) of lncRNAs compared to distributions for ARs (Table 1, Table S5, all $p < 0.01$). The divergence ratio (measured as $d_{\text{locus}}/d_{\text{AR}}$) at lncRNAs is $\sim 0.95$ (Table 1). For comparison, the divergence ratio for PC exons is 0.540 (Table 1). The divergence ratio for the 74 lncRNAs of known function is also lower than that for set of all lncRNAs ($0.797$, Table 1). Differences in diversity and divergence measures between intergenic and intronic lncRNAs are small although intronic lncRNAs did show consistently lower levels of diversity and divergence (Table 1, Figures 1 and 2).

Patterns of variation in the different lncRNA age groups suggest a trend of decreasing within-species diversity with increasing age (Table 2 and Figure 1). Linear regression models describing the relationship between the diversity estimates ($\pi$, $d$, $\pi/d$ and Tajima’s D) and evolutionary age support this trend of reduction of $\pi$, $d$ and Tajima’s D with age (Figures S1 – S4, Tables S6 – S9, Supplementary Materials). However, this trend is marginally non-significant for $\pi/d$ (diversity controlled for substitution rates) (Table S9, Figure S3, Supplementary Materials). Similarly, pairwise comparisons between the bootstrap distributions of diversity and divergence measures ($\pi$, $d$, $\pi/d$ and Tajima’s D) for each age group and the distributions for ARs give statistical
support for most differences, except in the cases of $\pi/d$ and Tajima’s D (Table 2). The greatest reductions in diversity ($\pi/d = 5.26$) are seen in lncRNAs common to all Tetrapods, representing about 0.6% of the total number of lncRNAs in the dataset. The value of $\pi/d$ is significantly different between lncRNAs and ARs in the case of only one age group (Eutherians = 5.42 vs. ARs = 5.82, $p < 0.0001$; Table 2) which represents about 16% of the total number of lncRNAs in the dataset.

These patterns all hold even when lncRNAs are grouped into fewer categories based on age (Tables S10 and S11, Supplementary Materials). Figure 1 also shows that confidence intervals of some diversity measures for older age groups, while overlapping with confidence intervals for ARs, also overlap the point estimates obtained for PC exons (e.g. Tetrapod lncRNAs $\pi$, $d$, $\pi/d$ and Tajima’s D; Figure 1). Point estimates of $\pi$, $d$, Tajima’s D and $\pi/d$ for lncRNA loci of known function show substantial and statistically significant reductions when compared to ARs (Table 2, Figure 1). These estimates for known lncRNAs are comparable to very old (~370 MY) lncRNAs in the larger dataset and also comparable to PC exons (Table 2, Figure 1).

Across lncRNA loci expressed in different tissues, there are similar levels of variation in locus and exon lengths (Table S4). The number of loci reaching highest expression in each tissue type varies substantially from 234 (placenta) to 6,157 (testes). There is some variation in diversity among lncRNAs expressed in different tissue types but variation in diversity with evolutionary age is greater (Table 2 and 3, Figure 1 and 2). Statistically significant differences for all measures of diversity and divergence are found only for the numerous testis-specific lncRNAs (Table 3, Figure 2). The other tissue types show mixed results with some diversity measures showing significant reductions compared to ARs. In particular, lncRNAs expressed in neural tissue show robust and statistically significant reductions in Tajima’s D, $\pi$ and $d$. However, none show significant differences in $\pi/d$ (Table 3, Figure 2).

**Patterns of Diversity in Regions Flanking Focal Loci**
The patterns of diversity and divergence in the regions flanking a locus can reveal the signature of selection by its effect on closely linked neutral sites. In the flanks of focal loci (lncRNA exons, PC exons and UTRs and AR loci), there is an obvious reduction only for PC exons, whereas lncRNAs and ARs show very similar patterns (Figure 3). These results hold when controlling for variation in mutation rate across the genome by normalizing by the between-species divergence ($\pi/d$ Figure 3). Patterns of diversity in regions flanking the loci are described by non-linear least squares models. In all cases, the model describing a curve is a better description, explaining significantly more of the variation in the data, than a model including only a y intercept (i.e. a straight line) (See Methods; lncRNAs $F_{(98, 199)} = 193.16$; ARs $F_{(98, 199)} = 39.43$; PC exons $F_{(98, 199)} = 439.93$; all $p < 0.01$). The parameter estimates in Table 4 show that patterns of $\pi/d$ in the regions flanking lncRNAs are both quantitatively and qualitatively more similar to ARs than to PC genes. The term $A$, measuring the “background” genomic level of $\pi/d$, is, as expected, comparable across lncRNAs, ARs and PC exons. The term $B$, measuring the reduction in $\pi/d$ as $x$ approaches 0 (i.e. closer to the focal locus), is negative and of similar magnitude in both lncRNAs and in ARs, whereas it is positive and greatly reduced in PC exons (Figure 3, Table 4). However, when mouse specific lncRNAs are removed from the dataset, the non-linear model is no longer the best-fitting model and the best prediction is a straight line with no slope and a $y$ intercept of $\pi/d$ similar to the background values of $\pi/d$ for ARs and PC exons (Figure 4, Table 4). This suggests that mouse specific lncRNAs drive up the diversity in the overall set of lncRNAs and older lncRNAs show lower levels of diversity. Additionally Tajima’s D is only markedly reduced at and around PC exons, indicating the presence of an excess of rare variants at these loci. Confidence intervals around estimates of $\pi$, $d$, $\pi/d$ and Tajima’s D in the regions immediately flanking the AR and lncRNA loci mostly overlap, suggesting that there is at most a small difference in the extent of reductions in diversity and divergence at sites linked to lncRNAs and ARs (Figure 3 and 4) but that these are not significant.

Discussion
We find a weak positive relationship between the estimated evolutionary age of lncRNAs and the level of conservation as measured by within-species sequence diversity and between-species divergence (Figure 1, Table 2). These observations suggest that lncRNAs that show homology and evidence of transcription across the greatest number lineages also show reduced diversity within mice and reduced divergence from rat at the sequence level, indicative of recent selection in mouse populations. These evolutionarily most ancient loci are therefore more likely to be important for fitness. Moreover, lncRNAs that have well characterized functions also tend to be more conserved than lncRNAs in general. This is not completely unexpected, because substantial sequence conservation is often a criterion by which researchers choose targets for further study. However, considering all 10,088 lncRNAs together, there is only limited evidence for constraint at the sequence level. Although nucleotide divergence at lncRNA loci is depressed by ~5% compared to AR loci, the patterns of diversity around lncRNA loci seen in Figures 3 and 4 do not suggest that strong selection affects variation at linked sites. There is some evidence for variation in the levels of diversity between lncRNAs expressed in different tissue types (Figure 2, Table 3). While lncRNAs specific to some tissue types show significantly lower levels of diversity in comparison to ARs, none show levels comparable to PC exons or lncRNAs of known function. Interestingly, neural tissue-specific lncRNAs seem to be shorter, on average, than lncRNAs in general. Similar results have been found in mouse embryonic brain tissue, where lncRNAs are shorter, on average, than known lncRNAs (Lv et al.2013). However, the shorter sequences among brain tissue lncRNAs are not likely to affect the reliability of our estimates. In general, confidence intervals are narrow indicating that the sample sizes used here are sufficient and that the limit on accuracy and sources of variation are mainly due to differing numbers of lncRNAs across groups in the analyses.

Similar findings to those detailed above have been reported for lncRNAs across the greater Tetrapod lineage, where conservation was significantly lower for lncRNA loci than random intergenic regions. Derived allele frequencies in a human polymorphism dataset are significantly
lower in lncRNAs than among random intergenic regions (Necsulea et al. 2014). Kutter et al. (2012) found that depressions in sequence divergence between mouse and rat was higher for lncRNA loci that show evidence of transcription in both species (~18%) than for orthologous lncRNA loci in general (~10%). Our estimates of depressions in sequence divergence, measured as divergence ratios \(d_{\text{locus}}/d_{\text{AR}}\), for lncRNAs compared to putatively neutrally evolving ARs are lower but comparable to the ~10-12% depressions found in other studies (Pang et al. 2006; Marques & Ponting 2009; Guttman et al. 2009; Kutter et al. 2012). The depressions in divergence such as those reported here and elsewhere have previously been taken as evidence of selection and functionality.

When considered in combination with the patterns of diversity around the focal loci (Figure 3 and 4), and the extent of depressions in divergence around known functional lncRNA loci (~20%, Table 1), the observed depressions in substitution rates of 5-12% across lncRNA loci in general are modest, suggesting that selection acting on lncRNAs is generally weak. However, some of the loci in the current dataset of ~10,000 lncRNAs show greater evidence of reduced within-species diversity.

The most striking patterns are seen in lncRNAs that are conserved deep in the Tetrapod lineage (about 0.6% of the total set of lncRNAs). In general, lncRNAs seem to show decreasing levels of within-species sequence diversity conservation with increasing evolutionary age. Thus, our estimates of the recent selective pressures acting on lncRNAs in mouse populations are compatible with their levels of long-term evolutionary conservation. There are several potential causes of this. One explanation is that the set of older lncRNAs contains a higher proportion of functional lncRNAs. Evolutionarily old lncRNAs could be important early in tetrapod development and therefore highly conserved at the sequence level. On the other hand, these ancient loci have to be detectable across more lineages in order to be included in the older groupings and it is possible that these higher levels of conservation are partly due to an overrepresentation in these groups of more conserved sequences that can be robustly aligned across such a wide phylogenetic range. The fact
that lncRNAs of known function show levels of sequence conservation comparable to older lncRNAs (e.g., tetrapod lncRNAs) suggests that the oldest groups are good candidates for further work to identify functional lncRNAs. The greater levels of sequence conservation seen at known lncRNA or functionally characterised loci also suggest that datasets of lncRNA loci identified by high-throughput sequencing and automated gene prediction, are dominated by “transcriptional noise”.

The above interpretations are subject to caveats. Firstly, previous work has shown that, in humans, lncRNA loci seem to have higher recombination rates than protein-coding genes (Necsulea et al. 2014). If this is also the case in mice, then this could lead to increased diversity, which might degrade the signal of selection at lncRNA loci and potentially explain some of the high levels of diversity seen in the current study. Second, ARs may not be evolving perfectly neutrally. If these loci are subject to some selection, then constraint will be underestimated, although the currently available evidence suggests that the neutrality of ARs is generally a reasonable assumption (Lunter et al. 2006). Similarly, constraint will also be underestimated if there is lineage specific positive selection at a locus. Third, one view holds that a lack of sequence conservation does not necessarily indicate a lack of function (Pang et al. 2006). Empirical examples of this can be found in the experimentally characterised lncRNAs Air and Xist. These loci show relatively low levels of conservation, despite having well established functions. At the Xist locus, significant sequence conservation across voles, mice and human lineages is only apparent in a few regions of the full sequence of the transcribed locus, whereas most of the locus evolves at a rate similar to that of neutrally evolving loci (Nesterova et al. 2001). Additionally, a survey of conserved regions across five vertebrate genomes (human, mouse, rat, chicken and the pufferfish), show that only 42% of sites in a core set of ncRNA loci, some of which are lncRNAs, are part of “highly conserved elements” (Siepel et al. 2005). Controversy over the use of conservation as an indicator of function recently arose when the ENCODE project claimed to have assigned function to ~80% of the human
genome, despite only ~10% being detectably conserved at the sequence level (Doolittle 2012; Stamatoyannopoulos 2012; The ENCODE Project Consortium et al. 2012; Graur et al. 2013). The conflict arises because it is unclear how a sequence can be functional in any meaningful way while not being affected by mutation (Doolittle 2012; Graur et al. 2013, Haerty & Ponting 2014).

Although it seems meaningless to discuss biological function of genomic elements in the complete absence of conservation at the sequence level (Doolittle 2012; Graur et al. 2013), conservation may not be detectable in averages of diversity and divergence across entire loci. For example, conservation could be apparent only at a few nucleotides at particular positions along the length of the locus that are important for secondary structure formation, which is the case in some lncRNA loci (Diederichs 2014). IncRNAs have been found to be more folded in secondary structure than predicted by chance, albeit to a lesser extent than mRNAs, suggesting that secondary structure, rather than simple nucleotide sequence, could be under constraint (Yang & Zhang 2015). It is also possible that lncRNA loci are not constrained by nucleotide sequence directly but for sequence length or continued transcription. One interesting proposition is that simply the continued transcription of lncRNA loci is required in order to keep chromatin open and thus facilitate the transcription of other nearby genes (Gribnau et al. 2000; Schmitt & Paro 2004; Ponjavic et al. 2007). For example, the Airn mediated silencing of Igfr2 in embryonic stem cells seems to depend only on overlap of the Airn locus with the Igfr2 locus and not the transcribed product of Airn (Latos et al. 2012). Furthermore, evidence suggests that nucleotide composition within IncRNAs mirrors that of PC loci and is under selection, over very short distances, for efficient transcription and splicing (Haerty & Ponting 2015). If high primary sequence constraint is not a requirement of function then selection for maintained function might not result in detectable signals of reduced diversity and divergence at and around functional lncRNA loci. Nevertheless, the observation reported here that IncRNAs of known function do show these signals of selection (reduced diversity and divergence) that are comparable to PC loci suggests that these signals are indeed good
indicators of functionality for at least some proportion of non-coding loci.

Another caveat is that there is a potential problem of alignment reliability especially for non-coding regions. Orthologous sequences are often inferred across organisms by alignment of two or more sequences and the nucleotide sequence similarity between them. However, a high sequence similarity between the identified best matches does not necessarily imply that these regions correspond to actively transcribed sequences across the organisms being compared (Ulitsky & Bartel 2013). Previous results suggest caution in this regard. For example, of ~3,000 mouse and ~2,600 human IncRNAs only 9% and 16.1%, respectively, aligned to any zebrafish sequence and only 7 and 9 loci, respectively, mapped to known zebrafish IncRNAs. Meanwhile, of those loci that mapped to any zebrafish sequence, 100 mouse and 286 human IncRNAs mapped to zebrafish coding regions (Ulitsky et al. 2011). The lncRNA locus HOTAIR can be aligned between human and mouse, but sequence conservation varies across the locus and is as low as 75% in certain regions (He et al. 2011; Schorderet & Duboule 2011). Moreover, the mouse homolog of HOTAIR is shorter with only two exons, while the human locus has six exons and deletion of the locus in mice does not show the expected effect on predicted regulatory targets (Schorderet & Duboule 2011).

Most of the HOTAIR locus may not be subject to selective constraint. Finally, of 160 lncRNA orthologs across rodents (M. m. musculus, M. m. castaneus and R. norvegicus) only 59.7% were found to be transcriptionally conserved, dropping to 28% if the data were adjusted to account for annotation biases (Kutter et al. 2012).

**Conclusions**

This study focuses on a recently published, large dataset of ~10,000 long non-coding RNAs (IncRNAs) in the mouse genome. We assessed the levels of within-species sequence diversity at these loci, in order to evaluate the recent selective pressures acting on IncRNAs and to compare them with estimates of long-term evolutionary conservation. We compared measures of sequence
diversity within mouse populations and divergence from the rat at these loci and in regions flanking them to the patterns around ARs and PC exons. It was possible to group lncRNA loci by their estimated evolutionary age (as inferred by evidence of homologous sequences and transcription across different evolutionary lineages) to determine the relationship between conservation and age.

In addition, a set of 74 lncRNA loci of known and characterised function in the mouse were investigated for comparison. The results suggest only modest conservation at the loci in the larger dataset when compared to PC exons. There is a trend of increasing conservation with increasing age of a locus, but this trend was not strong and was not consistent across all measures of diversity.

Finally, lncRNAs of known function show substantially and significantly reduced diversity and divergence when compared to neutrally evolving ARs that are comparable to PC exons. In summary, the data suggest that the majority of the lncRNAs in our dataset are evolving at rates comparable to ARs at the sequence level, but that a subset show signals of selection similar to PC genes.

Acknowledgements

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References.


Diederichs S 2014 The four dimensions of noncoding RNA conservation. Trends Genet. 30:121-123.


Table 1. Point estimates and number of loci (N) for divergence and diversity (%) measures across groups of genomic elements and divergence ratios (ratio of median divergence across loci to median divergence at ancestral repeats (ARs)). Bootstrapped 95% confidence intervals are given in brackets. Age group specific estimates of diversity measures with significantly different distribution of values, after a Bonferroni correction, when compared to the bootstrap distribution of ARs are denoted with * (actual p-values are reported in Table S5).

<table>
<thead>
<tr>
<th>Genomic Element (N loci)</th>
<th>Tajima’s D</th>
<th>d</th>
<th>π (%)</th>
<th>π/d</th>
<th>d/d_AR</th>
</tr>
</thead>
<tbody>
<tr>
<td>ARs (243,140)</td>
<td>-0.494</td>
<td>0.154</td>
<td>0.897</td>
<td>5.82</td>
<td>-</td>
</tr>
<tr>
<td></td>
<td>(-0.500, 0.487)</td>
<td>(0.154, 0.154)</td>
<td>(0.893, 0.900)</td>
<td>(5.80, 5.84)</td>
<td></td>
</tr>
<tr>
<td>PC Exons (49,470)</td>
<td>-0.734</td>
<td>0.0800</td>
<td>0.351</td>
<td>4.39</td>
<td>0.540</td>
</tr>
<tr>
<td>Known lncRNAs (74)</td>
<td>-0.859 *</td>
<td>0.122 *</td>
<td>0.532 *</td>
<td>4.38 *</td>
<td>0.797</td>
</tr>
<tr>
<td></td>
<td>(-1.03, -0.717)</td>
<td>(0.109, 0.130)</td>
<td>(0.436, 0.600)</td>
<td>(3.82, 4.95)</td>
<td></td>
</tr>
<tr>
<td>All lncRNAs (10,088)</td>
<td>-0.542 *</td>
<td>0.141 *</td>
<td>0.800 *</td>
<td>5.68 *</td>
<td>0.951</td>
</tr>
<tr>
<td></td>
<td>(-0.559, -0.526)</td>
<td>(0.140, 0.142)</td>
<td>(0.790, 0.808)</td>
<td>(5.61, 5.74)</td>
<td></td>
</tr>
<tr>
<td>Intergenic lncRNAs (8,433)</td>
<td>-0.541</td>
<td>0.142</td>
<td>0.815</td>
<td>5.75</td>
<td>0.958</td>
</tr>
<tr>
<td></td>
<td>(-0.558, -0.522)*</td>
<td>(0.141, 0.143)*</td>
<td>(0.805, 0.827)*</td>
<td>(5.68, 5.83)</td>
<td></td>
</tr>
<tr>
<td>Intronic lncRNAs (940)</td>
<td>-0.507</td>
<td>0.140</td>
<td>0.779</td>
<td>5.61</td>
<td>0.937</td>
</tr>
<tr>
<td></td>
<td>(-0.565, -0.443)</td>
<td>(0.137, 0.141)*</td>
<td>(0.750, 0.806)*</td>
<td>(5.41, 5.82)</td>
<td></td>
</tr>
</tbody>
</table>
Table 2. Point estimates and number of loci (N) for divergence and diversity (%) measures across groups of lncRNAs of different evolutionary ages and divergence ratios (ratio of median divergence across loci to median divergence at ancestral repeats (ARs)). Bootstrapped 95% confidence intervals are given in brackets. Age group specific estimates of diversity measures with significantly different distribution of values, after a Bonferroni correction, when compared to the bootstrap distribution of ARs are denoted with * (actual p-values reported in Table S5).

<table>
<thead>
<tr>
<th>Genomic Element (N loci)</th>
<th>Tajima’s D</th>
<th>d</th>
<th>π (%)</th>
<th>π/d</th>
<th>d/d_{AR}</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tetrapods (62)</td>
<td>-0.806</td>
<td>0.0893 *</td>
<td>0.469 *</td>
<td>5.26 (4.53, 5.99)</td>
<td>0.598</td>
</tr>
<tr>
<td>Amniotes (286)</td>
<td>-0.752 *</td>
<td>0.0860 *</td>
<td>0.470 *</td>
<td>5.46 (4.98, 5.96)</td>
<td>0.579</td>
</tr>
<tr>
<td>Mammals (165)</td>
<td>-0.769 *</td>
<td>0.0939 *</td>
<td>0.518 *</td>
<td>5.52 (4.88, 6.18)</td>
<td>0.631</td>
</tr>
<tr>
<td>Therians (255)</td>
<td>-0.669</td>
<td>0.105 *</td>
<td>0.583 *</td>
<td>5.55 (5.14, 5.97)</td>
<td>0.709</td>
</tr>
<tr>
<td>Eutherians (1,582)</td>
<td>-0.629</td>
<td>0.123 *</td>
<td>0.665 *</td>
<td>5.42 (5.27, 5.57)</td>
<td>0.828</td>
</tr>
<tr>
<td>Mouse (7,306)</td>
<td>-0.513</td>
<td>0.151 *</td>
<td>0.861 *</td>
<td>5.70 (5.63, 5.77)</td>
<td>1.02</td>
</tr>
</tbody>
</table>
Table 3. Point estimates and number of loci ($N$) for divergence and diversity measures and divergence ratios (ratio of median divergence across loci to median divergence at ancestral repeats (ARs)) for the long noncoding RNAs expressed in different tissues. Bootstrapped 95% confidence intervals are given in brackets. Age group specific estimates of diversity measures with significantly different distribution of values, after a Bonferroni correction, when compared to the bootstrap distribution of ARs are denoted with * (actual p-values reported in Table S12).

<table>
<thead>
<tr>
<th>Tissue type (N loci)</th>
<th>Tajima's D</th>
<th>$d$</th>
<th>$\pi$</th>
<th>$\pi/d$</th>
<th>$d/d_{AR}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Neural Tissue (1,345)</td>
<td>-0.595 * (-0.657, -0.537)</td>
<td>0.120 * (0.116, 0.123)</td>
<td>0.665 * (0.632, 0.696)</td>
<td>5.55 (5.29, 5.80)</td>
<td>0.809</td>
</tr>
<tr>
<td>Heart (358)</td>
<td>-0.626 * (-0.700, -0.552)</td>
<td>0.133 * (0.129, 0.136)</td>
<td>0.731 * (0.689, 0.776)</td>
<td>5.51 (5.23, 5.81)</td>
<td>0.895</td>
</tr>
<tr>
<td>Testes (6,157)</td>
<td>-0.539 * (-0.558, -0.520)</td>
<td>0.146 * (0.145, 0.147)</td>
<td>0.829 * (0.818, 0.841)</td>
<td>5.68 * (5.61, 5.76)</td>
<td>0.985</td>
</tr>
<tr>
<td>Ovary (479)</td>
<td>-0.503 (-0.569, -0.431)</td>
<td>0.138 * (0.135, 0.141)</td>
<td>0.800 * (0.761, 0.842)</td>
<td>5.80 (5.55, 6.09)</td>
<td>0.930</td>
</tr>
<tr>
<td>Placenta (234)</td>
<td>-0.451 (-0.564, -0.350)</td>
<td>0.132 * (0.127, 0.137)</td>
<td>0.736 * (0.684, 0.784)</td>
<td>5.57 (5.18, 5.95)</td>
<td>0.891</td>
</tr>
<tr>
<td>Kidney (396)</td>
<td>-0.575 (-0.644, -0.499)</td>
<td>0.135 * (0.131, 0.138)</td>
<td>0.765 * (0.725, 0.804)</td>
<td>5.67 (5.40, 5.93)</td>
<td>0.909</td>
</tr>
<tr>
<td>Liver (375)</td>
<td>-0.497 (-0.574, -0.415)</td>
<td>0.143 * (0.140, 0.146)</td>
<td>0.827 (0.782, 0.872)</td>
<td>5.76 (5.46, 6.07)</td>
<td>0.968</td>
</tr>
</tbody>
</table>
Table 4. Parameter estimates for the best-fitting non-linear least squares model applied to the patterns of diversity in the flanking regions of lncRNA, ancestral repeat (AR) and protein-coding (PC) loci. The equation applied was: $\pi/d \approx A x (1 - B x \exp(-\text{abs}(x)/q))$. Where $x$ is the distance from the focal locus (in kb).

<table>
<thead>
<tr>
<th>Genomic Element</th>
<th>A</th>
<th>B</th>
<th>q</th>
</tr>
</thead>
<tbody>
<tr>
<td>All lncRNAs</td>
<td>5.38</td>
<td>-0.08</td>
<td>42.22</td>
</tr>
<tr>
<td>All lncRNAs (excluding mouse-specific)</td>
<td>5.52</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>ARs</td>
<td>5.56</td>
<td>-0.06</td>
<td>2.25</td>
</tr>
<tr>
<td>PC Exons</td>
<td>5.66</td>
<td>0.15</td>
<td>7.97</td>
</tr>
</tbody>
</table>
Figure Legends

FIG. 1. Estimates of various diversity measures for each long non-coding RNA (lncRNA) age group. Groups are arranged along the x axis in descending order by evolutionary age from left to right. Error bars are bootstrapped 95% confidence intervals around point estimates. Estimates for lncRNAs of known function, Intronic lncRNAs and Intergenic lncRNAs, ancestral repeats (AR) and protein-coding (PC) exons are also shown for comparison.

FIG. 2. Estimates of various diversity estimates for lncRNAs expressed in different tissue types. Error bars are bootstrapped 95% confidence intervals around point estimates. Estimates for lncRNAs of known function, Intronic lncRNAs and Intergenic lncRNAs, ancestral repeats (AR) and protein-coding (PC) exons are also shown for comparison.

FIG. 3. Divergence ($d$), nucleotide diversity ($\pi$), divergence corrected nucleotide diversity ($\pi/d$) and Tajima’s D around focal loci for long non-coding RNAs (lncRNAs), known lncRNAs, ancestral repeats (AR) and protein-coding (PC) exons. Shaded areas indicate bootstrapped 95% confidence intervals. The panel “nls $\pi/d$” shows fitted values from a non-linear least squares function applied with the parameters described in Table 4. The x axis has been limited to 20kb down and upstream of the focal loci. The number of sites sampled as a function of distance in is shown in Figure S5.

FIG. 4. Divergence ($d$), nucleotide diversity ($\pi$), divergence corrected nucleotide diversity ($\pi/d$) and Tajima’s D around focal loci for long non-coding RNAs (lncRNAs), excluding “mouse-specific” lncRNAs, known lncRNAs, ancestral repeats (AR) and protein-coding (PC) exons. Shaded areas indicate bootstrapped 95% confidence intervals. The panel “nls $\pi/d$” shows fitted values from a non-linear least squares function applied with the parameters described in Table 4. The x axis has been limited to 20kb down and upstream of the focal loci.
Figure 1

- **Tajima's D**
  - Tetrapod lncRNAs
  - Amniote lncRNAs
  - Mammal lncRNAs
  - Therian lncRNAs
  - Eutherian lncRNAs
  - Mouse IncRNAs
  - All IncRNAs
  - Intergenic IncRNAs
  - Intronic IncRNAs
  - Known IncRNAs
  - Ancestral Repeats
  - Protein Coding Exons

- **π/σ**

- **π (%)**

Element:
- Ancestral Repeats
- Known IncRNAs
- IncRNAs
- Protein Coding Exons
Figure 2

[Bar graph showing the distribution of Tajima's D across different tissues and elements.]

Legend:
- Ancestral Repeats
- Known IncRNAs
- IncRNAs
- Known IncRNAs
- Ancestral Repeats
- Protein Coding Exons

Tissues:
- Neural Tissue
- Ovary
- Testes
- Heart
- Placenta
- Kidneys
- Liver
- All IncRNAs
- Intergenic IncRNAs
- Intronic IncRNAs

Elements:
- Protein Coding Exons
- Ancestral Repeats
- Known IncRNAs
- IncRNAs

Note: This graph was downloaded from http://gbe.oxfordjournals.org/ on August 26, 2015.
Figure 3

Genome Biology and Evolution

Distance from exon (kb)