Integrative annotation of human large intergenic noncoding RNAs reveals global properties and specific subclasses

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<td>Publisher</td>
<td>Cold Spring Harbor Laboratory Press</td>
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<tr>
<td>Version</td>
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<tr>
<td>Accessed</td>
<td>Tue Feb 12 03:19:30 EST 2019</td>
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Access the most recent version at doi:10.1101/gad.17446611

**Supplemental Material**
http://genesdev.cshlp.org/content/suppl/2011/09/01/gad.17446611.DC1.html

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Integrative annotation of human large intergenic noncoding RNAs reveals global properties and specific subclasses

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Large intergenic noncoding RNAs (lincRNAs) are emerging as key regulators of diverse cellular processes. Determining the function of individual lincRNAs remains a challenge. Recent advances in RNA sequencing (RNA-seq) and computational methods allow for an unprecedented analysis of such transcripts. Here, we present an integrative approach to define a reference catalog of >8000 human lincRNAs. Our catalog unifies previously existing annotation sources with transcripts we assembled from RNA-seq data collected from ~4 billion RNA-seq reads across 24 tissues and cell types. We characterize each lincRNA by a panorama of >30 properties, including sequence, structural, transcriptional, and orthology features. We found that lincRNA expression is strikingly tissue-specific compared with coding genes, and that lincRNAs are typically coexpressed with their neighboring genes, albeit to an extent similar to that of pairs of neighboring protein-coding genes. We distinguish an additional subset of transcripts that have high evolutionary conservation but may include short ORFs and may serve as either lincRNAs or small peptides. Our integrated, comprehensive, yet conservative reference catalog of human lincRNAs reveals the global properties of lincRNAs and will facilitate experimental studies and further functional classification of these genes.

Keywords: long noncoding RNAs, RNA sequencing, lincRNAs

Supplemental material is available for this article.

Received July 12, 2011; revised version accepted August 11, 2011.
screen identified dozens of lincRNAs required to maintain pluripotency and suggested that these lincRNAs work in trans (Guttman et al. 2011). Another class of “enhancer RNAs” may either be by-products of transcription [De Santa et al. 2010; Kim et al. 2010] or serve to activate gene expression in cis [Orom et al. 2010; Wang et al. 2011]. Despite these intriguing studies of individual lincRNAs, generalizing these findings to thousands of lincRNAs remains a substantial challenge. Collectively, lincRNAs are likely to reflect different families with distinct roles.

A first requirement toward functional categorization is a systematic catalog of lincRNA transcripts and their expression across tissues. In practice, however, researchers studying human lincRNAs are faced with an excessive set of noncoding transcripts of varying or unknown reliability that may not be well defined [Khalil et al. 2009] and have little or no expression data [Harrow et al. 2006], or with very small sets of experimentally validated ones [Amaral et al. 2010]. Transcripts in current annotations of the human transcriptome from the GENCODE/HAVANA [Harrow et al. 2006] or the University of California at Santa Cruz (UCSC) Genome Browser [Hsu et al. 2006] are valuable resources, but it is hard to evaluate their biological characteristics in the absence of expression levels and further processing.

Recent advances in RNA sequencing (RNA-seq) [Mortazavi et al. 2008] and computational methods for transcriptome reconstruction (Guttman et al. 2010; Trapnell et al. 2010; Garber et al. 2011) now provide an opportunity to comprehensively annotate and characterize lincRNA transcripts. Indeed, an initial application of this approach in three mouse cell types characterized the gene structure of more than a thousand mouse lincRNAs, most of which were not previously identified (Guttman et al. 2010).

Here, we present an integrative approach to define a reference set of lincRNAs that unifies existing annotation sources with transcripts reconstructed from >4 billion RNA-seq reads collected across 24 human tissues and cell types. We developed a conservative, broadly applicable pipeline to identify transcripts that are sufficiently expressed and have a negligible potential to encode proteins. We identified 8195 putative lincRNAs, of which 4662 (57%) form a “stringent” set. We characterized each lincRNA in the catalog by a panorama of structural, sequence, and expression features as an initial step toward fine categorization.

We used these features to test some of the proposed roles and characteristics of lincRNAs in a global and systematic way. For example, we found that lincRNAs—at all expression levels—are expressed in a highly tissue-specific manner—much more so than protein-coding genes. We observed no significant enrichment of correlated coexpression between lincRNAs and their neighboring genes beyond that expected for any two neighboring protein-coding genes. We identified expressed orthologous transcripts in another vertebrate species for 993 (12%) human lincRNAs. An additional set of 2305 other transcripts with high evolutionary conservation but ambiguous coding potential may function as noncoding RNAs or as small peptides. Finally, we highlight 414 lincRNAs that reside within intergenic regions previously associated with specific diseases/traits by genome-wide association studies (GWAS) as candidates for future disease-focused studies. Our reference catalog will facilitate future experimental and computational studies to uncover lincRNA functions.

Results

A computational approach for comprehensive annotation of lincRNAs

To comprehensively identify human lincRNAs, we developed a computational approach that integrates RNA-seq data with available annotation resources [Fig. 1A] and consists of four key steps (see the Materials and Methods): (1) transcriptome reconstruction of each sample from RNA-seq data using two transcript assemblers: Cufflinks (Trapnell et al. 2010), and Scripture (Guttman et al. 2010); (2) compilation of all noncoding and unclassified transcripts previously annotated; (3) integration of RNA-seq reconstructions with all annotation resources, using Cuffcompare (Trapnell et al. 2010) to determine a unique set of isoforms for each transcript locus; and (4) processing of the collected transcripts to identify lincRNAs, defined as transcripts that are reliably expressed, large, multixonic, noncoding, and intergenic.

There are two main challenges in applying this integrative approach to annotate lincRNA gene loci: (1) distinguishing lowly expressed lincRNAs (Guttman et al. 2010) from the tens of thousands of lowly expressed, single-exon, unreliable fragments assembled from RNA-seq; and (2) distinguishing novel transcripts encoding proteins or short peptides from bona fide noncoding ones. To address the first challenge, we removed unreliable lowly expressed transcripts using a learned read coverage threshold (Supplemental Material) and focus only on multixonic transcripts. To address the second challenge, we evaluated the coding potential of each of the remaining putative lincRNAs using two methods. First, we removed any putative ORFs that are evolutionarily constrained to preserve synonymous amino acid content, as reflected by a positive phylogenetic codon substitution frequency (PhyloCSF) metric [Lin et al. 2011] calculated for each locus across 29 mammals (Supplemental Material). Second, we scanned each transcript in all three reading frames to exclude transcripts that encode any of the 31,912 protein domains cataloged in the protein family database Pfam [Finn et al. 2010].

An annotated human lincRNA catalog

To generate a human lincRNA catalog, we applied our pipeline to polyadenylated RNA-seq data collected from 24 human tissues and cell lines. These included both single- and paired-end reads that are 50 or 75 bases long, sequenced on Illumina platforms (~4 billion reads total; ~175 million reads per sample on average) [Materials and Methods]. We integrated those with annotations from RefSeq [Pruitt et al. 2002], the UCSC Genome Browser [Hsu et al. 2006], and GENCODE (version 4) [Harrow et al. 2006] that were processed through our pipeline. We eliminated all annotated non-lincRNA transcripts (e.g., annotated protein-coding genes, microRNAs, tRNAs, and pseudogenes).
Figure 1. lincRNA catalog generation. [A] An integrative computational pipeline to map, reconstruct, and determine the coding potential of lincRNAs based on known annotations and computational methods, and its application to human lincRNAs. The pipeline takes as input RNA-seq data (top, red) and existing annotation sources (top) (RefSeq NR, Gencode, and UCSC annotation for humans). RNA-seq data are assembled by two assemblers: Cufflinks (gold) and Scripture (blue). Transcripts from all inputs are filtered by known annotations, presence of a Pfam domain, and positive coding potential. Transcripts annotated by RefSeq NR [+] were not filtered by the Pfam domain scan and the coding potential score. Finally, only multiexonic transcripts >200 base pairs [bp] are retained. [B] The number of lincRNA loci identified and their overlap with other annotation sources. The Venn diagram shows the overlap between transcripts from RNA-seq assembly (red), GENCODE and UCSC (purple), and RefSeq (green). [C] A representative example of a noncoding transcript that was reconstructed by Cufflinks and Scripture and was also curated in GENCODE and UCSC. [Top] The human genomic locus of the human lincRNAs (red) and its protein-coding neighbors. (Black, arrowhead) Direction of transcription. [Bottom] Magnified view of the lincRNA locus showing the coverage of RNA-seq reads from the testes [red] and the transcripts identified by each source [black], [iso] Isoform.

The initial catalog consists of a provisional set of 8195 intergenic transcripts (Fig. 1B). Although many of the previously annotated transcripts are also captured from the sequencing data [1864 lincRNAs identified by both] (Fig. 1B,C), most [4819] novel lincRNAs were only identified using RNA-seq. Based on the three samples for which we had two biological replicates [brain, testes, and lung fibroblasts], the reconstructed transcripts are highly reproducible: 70%–80% of assembled transcripts in the lower coverage replicate are also assembled in the higher coverage replicate [Supplemental Table 1; Supplemental Material].

Despite the high correspondence between protein-coding transcripts reconstructed by Cufflinks and Scripture (~85% of coding genes) [Supplemental Material; Supplemental Fig. 1A], there were larger differences between the noncoding transcripts assembled by the two methods, due to the differences in how each assembler reconstructs low-abundance transcripts (~46% of the putative lincRNAs were identified by only one source) [Supplemental Fig. 1B]. This is comparable with previously observed discrepancies in reconstruction of lowly expressed protein-coding genes (Garber et al. 2011) and is handled below.

We annotated each putative lincRNA in the provisional catalog with a comprehensive “profile” listing dozens of traits, such as its chromatin state, maximal expression level, proximity to coding genes, and evolutionary conservation [Materials and Methods, Supplemental Data Sets 1, 2]. Below, we use these features to define particular criteria by which we focus our analysis. Future users may leverage the annotated catalog through criteria of their choosing. A stringent set of 4662 human lincRNAs

We defined a stringent lincRNA set that includes those loci for which at least one lincRNA isoform was reconstructed in at least two different tissues or by two assemblers in the same tissue [Supplemental Material]. This leverages the unique benefits of each assembler, while in principle removing transcripts with insufficient coverage. The stringent set includes 4662 lincRNA loci [14,353 transcripts], 2798 of which (~60%) were not identified by RefSeq, UCSC, and GENCODE. We focused on the characteristics of this stringent set.

lincRNAs are alternatively spliced and preferentially proximal to developmental regulators

We characterized the basic features of lincRNAs, comparing them with protein-coding genes when appropriate. First, the size of lincRNAs is smaller than that of protein-coding transcripts, and they have fewer exons (on average, 2.9 exons and a transcript length of ~1 kb for lincRNAs vs. 10.7 exons and ~2.9 kb for protein-coding transcripts) [Supplemental Fig. 2A,B]. Notably, we may underestimate the length and exon number of lincRNAs, since their lower abundance may result in incomplete assembly. Second, lincRNAs are alternatively spliced [on average, ~2.3 isoforms per lincRNA locus] [Supplemental Fig. 2C]. Third, lincRNA loci are located from a few bases to >3 Mb from a protein-coding locus, with 28% within 10 kb of their coding neighbor [median = ~40 kb] [Supplemental Fig. 2D]. Finally, protein-coding genes proximal [≤10 kb] to lincRNAs are enriched for those associated with development and
transcriptional regulation [e.g., GATA2, GZF1, and NEUROG2 all have lincRNA neighbors] (Supplemental Fig. 3), consistent with previous reports [Guttman et al. 2009; Ponjavic et al. 2009].

Many lincRNAs are characterized by K4–K36 domains

We next explored the chromatin features of lincRNA loci as reflected in chromatin state maps from the nine ENCODE cell lines and other cells [Khalil et al. 2009; Ernst et al. 2011]. We examined each locus for the presence of a “K4–K36 domain,” a chromatin signature of actively transcribed genes that we previously used to identify lincRNAs [Guttman et al. 2009]. This domain consists of histone 3 Lys 4 trimethylation [H3K4me3] at the promoter followed by histone 3 Lys 36 trimethylation [H3K36me3] along the transcribed region. Despite the lack of paired matched samples of histone modifications and RNA-seq, 24% of the lincRNAs in our catalog have previously defined chromatin K4–K36 domains and ~40% have such domains when using less stringent criteria (with the remaining exhibiting partial signatures) (Supplemental Fig. 4; Supplemental Material).

lincRNA genes are no more likely to overlap enhancers than protein-coding genes

Recent studies reported short transcripts derived from enhancer elements, termed eRNAs, that are most likely not polyadenylated [Kim et al. 2010]. While this suggests that eRNAs and lincRNAs come from different classes, it is possible that longer polyadenylated transcripts may also be derived from enhancer elements and hence be related to eRNAs. To test this possibility, we examined the overlap between lincRNAs’ exons and two recent annotations of human enhancers based on genome-wide chromatin state maps. Twenty-seven percent of our lincRNAs and 44% of coding genes overlap 111,362 genomic regions previously suggested to function as enhancers [Ernst et al. 2011] in nine ENCODE cell lines (each overlap, $P < 0.001$, permutation test) [Supplemental Material]. When considering a more stringent subset of regions that are more likely to function only as enhancers [Supplemental Material], ~10% and 14% of lincRNAs and coding genes, respectively, overlap such regions [both $P < 0.001$]. Both lincRNAs and protein-coding genes have even lower overlap [both <3%, $P < 0.001$] with an enhancer set from human embryonic stem (ES) cells [Rada-Iglesias et al. 2010], possibly due to the lack of biological correspondence between the cell types and the tissue-specific nature of both lincRNAs and enhancers. Notably, this low overlap persists even when comparing more closely matched samples. Thus, only 15% of lincRNAs defined in mouse ES cells [Guttman et al. 2010] overlap enhancers defined in mouse ES cells [Zentner et al. 2011] [Supplemental Methods], and <1% of lincRNA defined in mouse neuronal progenitor cells [Guttman et al. 2010] overlap enhancer elements that express eRNAs in mouse cortical neurons [Supplemental Material, Kim et al. 2010]. Taken together, these data suggest that lincRNAs and eRNAs represent different subtypes of lincRNAs.

lincRNAs are expressed in a more tissue-specific manner than protein-coding genes

The maximal expression levels of lincRNAs are lower than those of protein-coding genes across the 24 samples [Fig. 2A], with a ~10-fold lower median maximal expression level [expression estimated with Cufflinks] [Fig. 2B; Materials and Methods; Trapnell et al. 2010]. Importantly, lincRNAs identified by RefSeq annotations were similarly lowly expressed relative to coding genes (~10 fold lower) [Supplemental Fig. 5]. These lower expression levels are consistent with previous reports [Ravasi et al. 2006; Guttman et al. 2010], suggesting a general property of lincRNAs.

The vast majority of lincRNAs exhibit tissue-specific expression patterns—more so than protein-coding genes—based on unsupervised clustering of expression profiles [Fig. 2A]. We further calculated a tissue specificity score for each transcript using an entropy-based metric that relies on Jensen-Shannon (JS) divergence [Materials and Methods]. This specificity metric (ranging from 0 to 1) quantifies the similarity between a transcript’s expression pattern across tissues and another predefined pattern that represents the extreme case in which a transcript is expressed only in one tissue. Thus, a perfect tissue-specific pattern will be scored as \( JS = 1 \).

Based on this measure, the majority of lincRNAs (78%) are tissue-specific, relative to only ~19% of coding genes \( (P < 10^{-300}, \) Fisher exact test) [Fig. 2C, Supplemental Fig. 6]. These differences are not the result of the low expression levels of lincRNAs and hold true for lincRNAs and protein-coding genes expressed at similar levels [Fig. 2B,C, Supplemental Fig. 6]. This was particularly true for the 35% of more highly expressed lincRNAs [and comparably expressed protein-coding genes, each with a maximal expression level of 3–20 FPKM [fragments per kilobase of exons per million fragments mapped]]. Thus, lincRNAs exhibit more tissue specificity than protein-coding genes at different expression ranges.

Approximately a third of our lincRNAs are specific to testes. Very few (<2%) of those overlap with a previously defined set of testes-specific small piRNAs (~30 nucleotides long) [Girard et al. 2006]. Thus, testes-specific lincRNAs may define a new class of RNAs in this organ. Testes-specific lincRNAs do not bias the global transcriptional characteristics above: lincRNAs that are not testes-specific are also lowly expressed and tissue-specific [presenting a qualitatively similar distribution with only moderately reduced tissue specificity scores] [Supplemental Figs. 5, 6A].

Finally, we predicted putative functions for our lincRNAs based on the known functions of protein-coding genes with similar expression patterns. We clustered lincRNAs and protein-coding genes using k-means clustering with the tissue specificity distance measure [Supplemental Material] and annotated each cluster with enriched functions of the protein-coding gene members. Clusters of tissue-specific lincRNAs and protein-coding genes are enriched for processes specific to that tissue or its differentiation (e.g., a liver-specific cluster is enriched with functional terms.


Such as cholesterol and lipid transport and homeostasis (Supplemental Fig. 7, Supplemental Data Sets 2, 3).

**lincRNAs are coexpressed with neighboring coding genes at levels similar to those expected for any pair of chromosomal neighbors**

The enrichment of specific gene functions in protein-coding genes neighboring lincRNAs and the presence of some pairs of neighboring lincRNA:protein-coding genes within expression clusters raise the hypothesis that such organization may be important for the regulatory function of lincRNAs. In particular, recent studies suggested that some lincRNAs may act in cis and affect the gene expression of their chromosomal neighborhood (Ponjavic et al. 2009; Orom et al. 2010).

One expectation from this hypothesis is that the expression of lincRNAs and their neighboring gene loci would be correlated across our samples. To test this hypothesis, we focused on the expression patterns of 1361 (28%) of our stringent lincRNAs that are located within 10 kb from a coding gene. Indeed, these lincRNAs and their coding neighbors were more correlated to each other than random gene pairs $[P < 5 \times 10^{-46}$, Kolomogorov-Smirnov [KS] test, $P < 10^{-807}$, Student’s t-test, effect size = 0.86] (Fig. 3A; Supplemental Material).

We must interpret this result with care, since the coexpression between a lincRNA and its protein-coding neighbor may result from either 1) a true cis effect of lincRNAs on its neighbor or 2) proximal transcriptional activity in the surrounding open chromatin [Ebisuya et al. 2008], since coexpression of chromosomal protein-coding gene neighbors was previously shown across eukaryotes (Cohen et al. 2000; Hurst et al. 2004). Supporting the second possibility, pairs of neighboring protein-coding genes were also more correlated to each other than random pairs $[P < 3.4 \times 10^{-159}$, KS test] (Fig. 3A). Furthermore, the correlation between lincRNA:protein-coding gene neighbors was only modestly higher than between protein-coding gene:protein-coding gene neighbors of a similar distance [effect size = 0.23, $P < 4.3 \times 10^{-7}$, KS test; $P < 6.9 \times 10^{-7}$, Student’s t-test] (Fig. 3A).

To further distinguish between these two possibilities, we focused on those protein-coding genes that had a lincRNA neighbor on one side and a coding neighbor on the other side, and used a paired test to compare the correlation between each protein-coding gene and its lincRNA neighbor with that between the same protein-coding gene and its protein-coding gene neighbor. This paired comparison showed a weak opposite trend, where pairs of coding gene neighbors are slightly more correlated to each other than neighboring lincRNA:protein-coding gene pairs $[P < 0.001$ paired Student’s t-test, effect size = 0.23], thus favoring option 2, an effect of gene proximity.

Taken together, this analysis suggests that, overall, lincRNAs are not more correlated to their protein-coding gene neighbors than expected for a pair of neighboring protein-coding gene loci. Yet, the ultimate test of cis- or...
trans-regulatory mechanisms for lincRNAs requires experimental gain-of-function or loss-of-function data.

Divergently transcribed lincRNAs

Unstable, likely noncoding, transcripts can also be derived from divergent [bidirectional] transcription in both yeast and mammals [Core et al. 2008; Preker et al. 2008; Seila et al. 2008]. These may be either by-products of chromatin remodeling and recruitment of the transcription machinery to the neighboring gene’s promoter or functional transcripts [Kanhere et al. 2010]. Due to limited read length and computational methods, previous studies did not determine whether these transcripts are spliced. Interestingly, several functionally studied lincRNAs, including Tug1 [Young et al. 2005], HOTAIR [Rinn et al. 2007], and HOTTIP [Wang et al. 2011], are divergent transcripts. We therefore hypothesized that other divergently transcribed transcripts may be spliced and polyadenylated lincRNAs.

Indeed, 588 (~13%) of our stringent lincRNAs are spliced transcripts divergently transcribed within 10 kb of a coding gene promoter, with a majority (~65%) that initiate within 1 kb of a coding gene’s annotated transcription start site [Supplemental Fig. 8]. Furthermore, ~35% of the 588 pairs share a H3K4me3 domain [a hallmark of active promoters], based on the ENCODE chromatin state maps [Supplemental Material], although we cannot definitively determine whether these divergently encoded pairs are also divergently transcribed from the same promoter. These divergent coding gene neighbors are enriched for developmental and metabolic processes [Supplemental Fig. 3B]. Focusing on the 68% that are spliced in the tissue where they are maximally transcribed (Supplemental Fig. 3B). Furthermore, while ~49% of the divergently transcribed lincRNAs are tissue-specific, for approximately half of those, the neighboring gene is ubiquitously expressed [Fig. 3C]. Thus, although there are clearly bidirectionally transcribed, spliced lincRNAs in our catalog, we found no clear additional distinguishing features for this set.

Expressed syntenic orthologs of human lincRNAs in mammals and vertebrates

We and others have previously reported evidence for purifying selection at different sets of mammalian lincRNAs [Ponjavic et al. 2007; Guttman et al. 2009; Orom et al. 2010]. A recent study has also identified expressed orthologs of a few highly conserved and brain-expressed mouse lincRNAs in species as distant as opossums and chickens [Chodroff et al. 2010]. However, the number of human lincRNAs that have an orthologous, actively expressed, transcript in other species remains unknown.

To identify human lincRNAs with orthologous expressed transcripts in other species [supported by experimental evidence], we surveyed a catalog of mammalian and nonmammalian vertebrate transcripts that were syntenically mapped to the human genome by TransMap [Zhu et al. 2007], a cross-species mRNA alignment method. TransMap maps all known transcripts [e.g., full-length cDNAs and others in RefSeq or UCSC] and ESTs across vertebrate species using syntenic BLASTZ alignments [Schwartz et al. 2003] that use conserved gene order [synteny]. Since EST coverage varies between species [Supplemental Table 2], TransMap can only provide a lower-bound estimate of orthologous transcripts.

Of the 8195 lincRNAs, 993 are syntenically paired with an orthologous transcript [Fig. 4A–D], comprising a trans-mapped lincRNA set (~135 expected by random permutations) [Materials and Methods, Supplemental Material]. Seven-hundred-two of the trans-mapped lincRNAs are in the stringent lincRNA set (~15% of stringent lincRNAs). The majority [53%] of the trans-mapped lincRNAs was not previously annotated in the human transcriptome.
Trans-mapped lincRNAs have tissue specificity and low expression, comparable with that of all other lincRNAs (Supplemental Figs. 6A, 9B,C). Fifty-nine percent of the trans-mapped lincRNAs were mapped to annotated transcripts that had evidence beyond ESTs. Supporting our noncoding classification scheme, only 18% of the 641 lincRNAs with trans-mapped orthologous transcripts in mice were classified as coding in mice and only ~11% have a positive PhyloCSF score (Materials and Methods, Supplemental Fig. 9A). Trans-mapped lincRNAs have orthologs in species from mice to fish, with closer species that have more transcriptome data showing more orthologs than distant ones (Fig. 4D).

Orthologous lincRNAs exhibit modest sequence homology

We evaluated the degree of sequence similarity between the trans-mapped transcripts. We measured the portion of each lincRNA transcript’s length that is aligned to the orthologous transcript. The majority of trans-mapped lincRNAs are only moderately spanned by an orthologous mapped transcript (a median of 21% and 56% of their transcript or genomic locus, respectively, aligned) (Fig. 4E).
be due to either cryptic small peptides in the human transcript or the evolution of a noncoding transcript from a coding one. The available data are insufficient to distinguish between these hypotheses, which can be tested as paired cross-species RNA-seq samples are collected.

We next compared the fraction of identical bases aligned between the lincRNAs and their orthologs with that of random sequence pairs, randomly selected syntenic blocks, or orthologous coding genes. *Trans*-mapped lincRNAs and their orthologous transcripts show sequence identity similar to that of randomly selected syntenic blocks, which is lower than pairs of orthologous protein-coding genes and higher than for random pairs of genomic regions of similar size (Fig. 4F; Supplemental Fig. 10C,D; Materials and Methods). With only 34% of the human genome syntenically mapped to the mouse genome (Kent et al. 2003), the resemblance of *trans*-mapped lincRNAs to random syntenic blocks still implies evolutionary constraint to preserve sequence elements.

**Novel transcripts with potential coding capacity**

While our stringent lincRNA classification strategy focused on noncoding transcripts, we also characterized 2305 transcripts that were excluded by our coding potential criteria [a Pfam domain, a positive PhyloCSF score, or previously annotated as pseudogenes] and termed them the transcripts of uncertain coding potential [TUCP] set [Supplemental Material]. These may include lincRNAs as well as other transcripts. The majority (1533; ~66%) was previously annotated as pseudogenes that, due to our focus on multiexonic transcripts, are probably not retrotransposed, spliced mRNAs that were integrated back to the genome (Fig. 5A). Similar to the stringent set, TUCP transcripts are expressed at lower and more tissue-specific patterns than protein-coding genes (Fig. 5B,C).

The coding potential of most of these transcripts was very low compared with known coding genes, and only 32% (757) exceeded our PhyloCSF score criteria (Fig. 5A,D; Materials and Methods). The evolutionarily constrained ORFs in these transcripts are mostly short [51% are <70 amino acids long] and cover a small portion of the transcript [53% cover <25%] (Fig. 5E,F). Thus, some of these transcripts may encode small functional peptides (Kondo et al. 2010), whereas others may function as noncoding RNA.

TUCP transcripts are under stronger purifying selection than stringent lincRNAs. First, the exonice sequence in TUCP transcripts is more highly conserved than that of stringent lincRNAs \( P < 10^{-168}, \) effect size = 0.77 [Supplemental Fig. 11; Supplemental Material], even when excluding pseudogenes [Supplemental Fig. 11]. Second, a larger fraction of them has a *trans*-mapped syntenic ortholog \( \sim 36\% \) [838], or \( \sim 34\% \) when excluding pseudogenes, compared with \( \sim 15\% \) [702] of stringent lincRNAs, and the syntenic alignments cover a slightly larger portion of the transcript [Supplemental Fig. 12]. Third, 74% of the *trans*-mapped transcripts have an ortholog in a species more distant than mice [vs. 37% of the *trans*-mapped lincRNAs; \( \sim 67\% \) when excluding TUCP pseudogenes] (Fig. 5G).

**lincRNAs in disease-associated regions**

Although GWAS have identified thousands of common genetic variants related to specific traits or disease phenotypes, many of these variants \( \sim 43\% \) [Hindorff et al. 2009] lie in intergenic regions and hence remain largely unexplained. We identified 414 lincRNAs from our comprehensive catalog [215 of the stringent set] that are located within 1146 disease- or trait-associated regions from the published GWAS catalog [Hindorff et al. 2009] that do not contain annotated coding genes [Supplemental Material; Supplemental Data Set 2]. Notably, 30 and 81 of those lincRNAs overlap a common variant that was associated with a disease phenotype within their exon or their intron, respectively [both tag and imputed SNPs]. Another 76 intergenic disease/trait regions overlap 84 TUCP transcripts [Supplemental Data Set 6].

The 215 stringent lincRNAs in these regions are typically expressed in a tissue-specific manner, which in a few cases directly corresponds to the tissue relevant to the associated disease [Supplemental Table 3]. For example, a lincRNA positioned \( \sim 3 \) kb downstream from a thyroid cancer-associated SNP in chromosome 14q13.3 \([r s 944289, \) odds ratio [OR] = 1.37, \( P = 2.0 \times 10^{-7} \) [Gudmundsson et al. 2009]] is strongly expressed specifically in the thyroid \([\sim 5.4 \log 2 \text{FPKM}] \). The “tag SNP” and the proximal lincRNA are within a 249-kb linkage disequilibrium (LD) region that does not contain any known genes. \( r s 944289 \) is \( \sim 3.5 \) kb upstream of the transcription start site of the thyroid-specific lincRNA. \( r s 944289[T] \) is predicted to be part of a binding motif for C/EBP-\( \alpha \) [Supplemental Material; Supplemental Fig. 13] and may affect the lincRNA’s expression. The LD region is \( \sim 250 \) kb upstream of the gene *NKX2-1* (TTF1), a transcription factor with a prominent role in thyroid development and a previously suggested candidate gene for this SNP association. The lincRNA may be an additional candidate, playing a role in thyroid-specific processes [possibly in coordination with the neighboring *NKX2-1*] and in thyroid cancer.

**Discussion**

We generated a reference catalog of 8195 human lincRNAs based on integrating RNA-seq data from 24 tissues and cell types with publicly available transcript annotations. Fifty-eight percent of the transcripts in our catalog are novel and are now identified for the first time using RNA-seq. We annotated each lincRNA with a broad range of structural, expression, and evolutionary features, shedding new light on their global properties and testing or generalizing previous hypotheses.

lincRNAs are remarkably tissue-specific compared with protein-coding genes. This possibility was previously raised [Mercer et al. 2009; Ponting et al. 2009] based on differential expression patterns in specific biological systems and has several implications. First, researchers studying a particular system may benefit from RNA-seq profiling followed by de novo assembly in that system. Second, it is consistent with the hypothesis that some lincRNAs interact with chromatin modulators and provide
their target specificity. Third, it may indicate that lincRNAs could serve as specific fine-tuners. Fourth, the low level of lincRNA expression in a complex tissue such as the brain may in fact be a by-product of their expression in only a few specific cells. Future targeted perturbations of tissue-specific lincRNAs defined in our study may elucidate their role in tissue-specific processes.

Could many lincRNAs act as enhancer elements, promoting the transcription of their neighboring coding genes? Recent studies have demonstrated that several lincRNAs have enhancer-like functions (Orom et al. 2010; Wang et al. 2011). While our coexpression analysis is consistent with this notion, it is insufficient to suggest a global trend in which lincRNAs act as enhancers of their neighbors, since neighboring coding genes exhibit similar coexpression patterns. Further systematic perturbation studies in individual systems (as in Orom et al. 2010) may help assess the scope of this function. Notably, a very recent study that systematically perturbed 150 lincRNAs expressed in mouse ES cells suggested that lincRNAs primarily affect gene expression in cis (Guttman et al. 2011). Collectively, this suggests that some lincRNAs can work in cis, while others work in trans.

Nine-hundred-ninety-three lincRNAs have an orthologous transcript expressed from a syntenic region in another species, 50% of which were identified for the first time in this study. These lincRNAs had only moderate sequence identity and alignment to their orthologs. This moderate...
conservation may indicate the importance of transcription from a specific genomic location, the reduced selective pressure on the primary sequence of noncoding RNAs [Brown et al. 1992; Zhao et al. 2008], or the rapid evolution of new functions. It may also be due to alignment to orthologous ESTs that are incomplete transcripts. Our analysis was limited by available transcript data in other species, and will be enhanced as more transcriptomes are sequenced in other organisms.

TUCP intergenic transcripts did not pass our stringent classification criteria as lincRNA due to evidence of possible protein-coding potential. These transcripts have expression levels and tissue specificity similar to the stringent lincRNA set, but a significantly higher level of sequence conservation. Many could encode small peptides, similar to those that function in Drosophila melanogaster embryogenesis [Kondo et al. 2010]. Another 1533 TUCP transcripts are classified as pseudogenes, and may represent pseudogenes that have evolved to function as noncoding regulatory agents. Ribosome profiling [Ingolia et al. 2009] and mass spectrometry of small peptides will help to resolve which of the TUCP transcripts are more likely to be coding.

Substantial progress has been recently made toward the essential goal of annotating long noncoding RNA loci. Our study presents an integrative yet conservative computational approach to mapping lincRNA transcripts that can be used for mapping new transcripts in other species. This is critical to overcome major barriers for future experiments (e.g., cloning, expression profiling, gain of function, and loss of function), as well as for the interpretation of genetic association studies. Indeed, 414 lincRNAs in our catalog stand out as located within intergenic regions associated with common disease. Future work will be necessary to identify RNA sequence domains that relate to function [Zhao et al. 2008; Kanhere et al. 2010], and to further classify lincRNAs into families. Our panorama of lincRNA properties will greatly advance these goals.

Materials and methods

RNA-seq data sets

We used two data sets of RNA-seq for transcriptome reconstruction. The first includes polyadenylated RNA samples from 16 tissues that were sequenced using Illumina HiSeq 2000 as part of the Human Body Map 2 project (235 million reads per sample on average) [Supplemental Table 4]. The second data set included eight additional tissues and cell lines, each sequenced by the Illumina Genome Analyzer II (GAII) (54 million reads per sample on average) [Supplemental Table 4]. The Human Body Map 2 data are accessible from ArrayExpress [accession no. E-MTAB-513; http://www.ebi.ac.uk/arrayexpress/browse.html?keywords=E-MTAB-513&expand=on].

The eight additional tissues and cell lines are available at Gene Expression Omnibus (GEO) [accession no. GSE30554] [see the Supplemental Material].

Publicly available annotations

All known annotations that were used for the analysis of this study are specified in Supplemental Table 5.

lincRNA classification pipeline

Once the transcriptome of each tissue sample was assembled (Supplemental Material), we further processed the assemblies and used Cuffcompare [Trapnell et al. 2010] to eliminate intron and polymerase run-on fragments surrounding all transcripts annotated by GENCODE 4. We then used Cuffcompare to generate a unique set of assembled isoforms from all processed tissue assemblies. Next, we ran the unique transcript set through the following filters: (1) size selection, (2) minimal read coverage threshold, (3) filter of known non-lincRNAs annotations, (4) positive coding potential threshold, (5) known protein domain filter, and (6) intergenic classification (see the Supplemental Material).

To derive a unique set of lincRNAs that includes previous annotations, we used Cuffcompare to integrate the RNA-seq-derived lincRNAs with the predetermined set of lincRNAs previously annotated by RefSeq, UCSC, or GENCODE 4. The publicly available lincRNA sets were derived by running specific steps of our lincRNA classification pipeline on the transcripts annotated in the public data sets [Fig. 1A; see Supplemental Table 5 for specific details].

lincRNAs catalog and annotation

The complete lincRNA catalog (including the TUCP transcripts) as well as all RNA-seq alignments and transcriptome reconstructions are available at http://www.broadinstitute.org/genome_ bio/human_lincrnas. Specific descriptions of all characterization fields are provided on the site. The catalog is also provided as Supplemental Data Sets 1–6.

Estimating expression abundance and normalization

We estimated the expression abundance of all lincRNAs and protein-coding genes by running Cufflinks in its expression abundance estimation mode across our 24 samples [Trapnell et al. 2010]. We used the complete noncoding transcripts catalog and all coding transcripts annotated in UCSC for a comprehensive representation of transcripts along the genome while performing abundance estimation. FPKM calls were log2-normalized (after addition of $\varepsilon = 0.05$). The HeLa and liver samples from the eight-sample set were eliminated from further expression analysis due to low coverage and a lower expression range in comparison with other samples.

Tissue specificity score

To evaluate the tissue specificity of a transcript, we relied on Trapnell et al. [2010] and devised an entropy-based measure that quantifies the similarity between a transcript’s expression pattern and another predefined pattern that represent an extreme case in which a transcript is expressed in only one tissue. This specificity measure relies on the JS divergence. The JS divergence of two discrete probability distributions, $p', p''$, is defined to be

$$JS(p', p'') = H\left(\frac{p' + p''}{2}\right) - \frac{H(p')} + H(p'')}{2},$$

where $H$ is the entropy of a discrete probability distribution:

$$H(p) = - \sum_{i=1}^{n} p_i \log(p_i).$$

Relying on the theorem that the square root of the JS divergence is a metric [Fuglede and Topsoe 2004], we define the distance...
between two tissue expression patterns, $e^1$ and $e^2$, $e^i = (e^i_1, \ldots, e^i_n)$, as

$$JS_{dibis}(e^1, e^2) = \sqrt{JS(e^1, e^2)}.$$  \hspace{1cm} (3)

The tissue specificity of a transcript’s expression pattern, $e$, across $n$ tissues with respect to tissue $t$ can then be defined as

$$JS_{sp}(e) = 1 - JS_{dibis}(e^t, e'),$$  \hspace{1cm} (4)

where $e'$ is a predefined expression pattern that represents the extreme case in which a transcript is expressed in only one tissue. Formally, $e' = (e'_1, \ldots, e'_n)$, s.t. $e'_t = \begin{cases} 1 & \text{if } t = t \\ 0 & \text{otherwise} \end{cases}$.

Finally, we define the tissue specificity score of a transcript as the maximal tissue specificity score across all $n$ tissues of the transcripts expression pattern $e$:

$$JS_{sp}(e) = \arg\max_t JS_{sp}(e^t), \quad t = 1 \ldots n.$$  \hspace{1cm} (5)

Refer to the Supplemental Material for further details on the normalization of expression vector for tissue specificity calculation.

Identification of trans-mapped syntenic orthologs of human lincRNAs

We downloaded all available TransMap mappings of expressed transcripts to the human genome [NCBI39/Hg19] from the UCSC Genome Browser [http://genome.ucsc.edu, Zhu et al. 2007]. The TransMap methodology maps all annotated transcripts of one species to the other by using the syntenic BLASTZ alignments between two species [Schwartz et al. 2003]. First, it aligns all mRNA sequences of species $a$ to its own genome. Then, it uses the syntenic alignment between species $a$ and $b$ to project the mRNA sequence of $a$ to the genome of $b$ and finally refines this mapping. We crossed all UCSC, RefSeq, mRNA, and EST transcripts trans-mapped to humans with our lincRNA set and included every lincRNA that had an exon overlap with a trans-mapped transcript in the trans-mapped lincRNA set. We used the UCSC classification of coding and noncoding transcripts applied to human and mouse transcripts known to UCSC (and downloaded from the UCSC Genome Browser [Hsu et al. 2006]).

Refined alignment of human lincRNAs and their mouse orthologs

To assess the alignment quality of the trans-mapped lincRNAs and their syntenic orthologs, we realigned the transcript sequence of all human lincRNAs and their mouse orthologs using the fast statistical alignment algorithm with default parameters (see the Supplemental Material, Bradley et al. 2009).

Acknowledgments

We gratefully thank Manuel Garber, Mitchell Guttman, Or Zuk, and Noam Shoresh for insightful discussions and code. C.T. is a Damon Runyon Cancer Foundation Fellow. L.G. is an NSF Post-doctoral Fellow. M.K. is a Human Frontier Science Program Fellow. A.R. is supported by an NIH Pioneer Award, the Burroughs Wellcome Fund Career Award at the Scientific Interface, and HHMI. J.L.R. is a Damon Runyon-Rachleff, Searle, Smith Family Scholar, and is supported by Director’s New Innovator Awards (1DP2OD00667-01). This work was funded in part by a Center of Excellence in Genome Science from the NHGRI (A.R. and J.L.R.). A.R. and J.L.R. are investigators of the Merkin Foundation for Stem Cell Research at the Broad Institute. The body map data were kindly provided by the Gene Expression Applications research group at Illumina.

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