Many human large intergenic noncoding RNAs associate with chromatin-modifying complexes and affect gene expression

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Detailed Terms
Many human large intergenic noncoding RNAs associate with chromatin-modifying complexes and affect gene expression

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We recently showed that the mammalian genome encodes >1,000 large intergenic noncoding (linc)RNAs that are clearly conserved across mammals and, thus, functional. Gene expression patterns have implicated these lincRNAs in diverse biological processes, including cell-cycle regulation, immune surveillance, and embryonic stem cell pluripotency. However, the mechanism by which these lincRNAs function is unknown. Here, we expand the catalog of human lincRNAs to ~3,300 by analyzing chromatin-state maps of various human cell types. Inspired by the observation that the well-characterized lincRNA HOTAIR binds the polycomb repressive complex (PRC2), we tested whether many lincRNAs are physically associated with PRC2. Remarkably, we observe that ~20% of lincRNAs expressed in various cell types are bound by PRC2, and that additional lincRNAs are bound by other chromatin-modifying complexes. Also, we show that siRNA-mediated depletion of certain lincRNAs associated with PRC2 leads to changes in gene expression, and that the up-regulated genes are enriched for those normally silenced by PRC2. We propose a model in which some lincRNAs guide chromatin-modifying complexes to specific genomic loci to regulate gene expression.

histone modifications | epigenetic regulation | polycomb

Mammalian genomes produce a wide variety of noncoding RNA transcripts (1–3). In addition to classical RNAs (e.g., ribosomal RNAs, transfer RNAs, and others) and more recently discovered classes of small noncoding RNAs (e.g., microRNAs) (3, 4), there are many large noncoding RNAs of unknown function (3). Several, such large noncoding RNAs have been biologically characterized (including XIST, TSIX, HOTAIR, and AIR) (3), but shotgun cDNA sequencing and microarray hybridization have suggested that the vast majority of the mammalian genome can produce RNA transcripts under some circumstances (2). However, the biological significance of these transcripts has been highly controversial because most occur at extremely low levels and show little evolutionary conservation (5). A previous study reported that a subset of these ncRNAs shows evidence of evolutionary conservation (27), suggesting that at least some of these ncRNAs are functional.

Recently, we developed an approach for identifying large noncoding RNAs based on a distinctive chromatin signature that marks actively transcribed genes (1). The signature consists of a short region with histone H3 lysine 4 trimethylation (H3K4me3) (corresponding to the promoter) and a longer region with histone H3 lysine 36 trimethylation (H3K36me3, corresponding to the transcribed region) (1, 6, 7). We refer to this chromatin signature as a K4-K36 domain. We generated chromatin-state maps across 4 mouse cell types, searched for K4-K36 domains, and then eliminated those corresponding to known protein-coding genes. We found 1,586 previously uncharacterized K4-K36 domains in the 4 mouse cell types, and showed that the vast majority encode large intergenic noncoding (linc)RNAs. These lincRNAs show similar expression levels as protein-coding genes, but lack any protein-coding capacity. Importantly, lincRNAs show significant evolutionary conservation relative to neutral sequences, providing strong evidence that they have been functional in the mammalian lineage (1). We note that nonconserved RNA sequences identified in other collections could be functional, but biological evidence such as loss-of-function experiments would be needed to establish their functionality (5) (Fig. S1 A). Previous studies by us and others have demonstrated that groups of lincRNAs exhibit expression patterns across cell types and tissues that correlate with patterns seen for protein-coding genes involved in cellular processes such as cell-cycle regulation, innate immune responses, and stem cell pluripotency (1, 14). Although these studies clearly demonstrate that there are many functional lincRNAs, key questions remain, including: How many lincRNAs are encoded in mammalian genomes? How do lincRNAs exert their functions? To begin to investigate the number of lincRNAs, we extended our approach of mapping K4-K36 domains to 6 human cell types. The results expand our catalog to 3,289 lincRNAs, which show clear evolutionary conservation within their transcripts. Extrapolation suggests that the total number may approach ~4,500 lincRNAs.

To examine the biochemical mechanism by which lincRNAs function, we drew inspiration from one of the few well-studied lincRNAs: HOTAIR. We previously reported HOTAIR as a lincRNA transcribed from within the HOX D cluster, and showed that it acts to repress genes in the HOXD cluster, by binding to the polycomb repressive complex (PRC2) and recruiting it to the locus (8). PRC2 is a methyltransferase that trimethylates H3K27 to repress transcription of specific genes (9, 10). Recently, several other large noncoding RNAs have been found to associate with chromatin-modifying complexes, including a large noncoding RNA encoded within the 5′ of XIST that can target PRC2 to the inactive X chromosome (11), the antisense transcript AIR that is associated with the chromatin-modifying complex G9a, an H3K9me2 meth-
yltransferase (12), and the Kcnq1ot1 transcript that binds both G9a and PRC2 (13). Some recent studies have demonstrated a few large noncoding RNAs bind chromatin proteins that add activating modifications (e.g., Trithorax) in mES cells (14). These few examples raised the possibility that many lincRNAs might be physically associated with chromatin-modifying complexes and might potentially target them to specific genomic regions. To test this hypothesis on a genomic scale, we performed RNA commounprecipitation (RIP) with antibodies directed against several proteins involved in chromatin-modifying complexes (PRC2, CoREST, and SMXC). We find that as many as 38% of the lincRNAs expressed in the cell types studied are reproducibly associated with one of these complexes. Also, we show that RNA-interference-based depletion of various PRC2-associated lincRNAs results in activation of genes known to be repressed by PRC2. Together, our results indicate that thousands of functional lincRNAs are encoded in the human genome, and a significant proportion of lincRNAs are physically associated with chromatin-modifying complexes. We propose a shared mechanism for hundreds of lincRNAs that function in regulating the epigenetic landscape at distinctive target loci.

**Results**

**Identification of Human lincRNAs.** We recently identified lincRNAs in the mouse genome by analyzing chromatin-state maps of 4 mouse cell types: mES, lung fibroblasts (mLFs), neural precursor cells (NPCs), and mouse embryonic fibroblasts (MEFs) (1). Specifically, we used a computational algorithm to identify K4-K36 domains that do not overlap known protein-coding genes, and then eliminated the small proportion (~5%) that showed any significant protein-coding capacity. We were left with 1,586 previously uncharacterized K4-K36 domains. We demonstrated that the vast majority of these regions encode lincRNAs, with 95% showing significant conservation to the human genome within their transcripts. To further extend the catalog of lincRNAs, we sought to analyze chromatin-state maps of 6 human cell types: hESC (10), 2 hematopoietic stem cells (CD133+ and CD36+) (15), T-cells (6), hLFs (1), and normal embryonic kidney (hEK). Using our previous computational approach, we identified K4-K36 domains that are well-separated from (i) the regions containing known protein-coding genes and all known classes of small noncoding RNAs in humans and (ii) the orthologous regions of known protein-coding genes in mouse, rat, and dog. We also eliminated the orthologous regions of our previously identified mouse lincRNAs. We previously showed that, for similar cell types in mouse and human, lincRNA loci show cross-species conservation not only at the level of nucleotide sequence, but also with respect to the presence of K4-K36 domains (1). We found a total of 1,833 previously uncharacterized intergenic K4-K36 domains in these 6 human cell types (Fig. 1A; Fig. S1B and table 1 in Dataset S1). We analyzed the coding potential of each such K4-K36 domain using the codon substitution frequency score (SI Methods), and found that <8% showed any evidence of protein-coding capacity (Fig. S1C) (16). After eliminating these cases, we were left with 1,703 loci encoding putative lincRNA genes.

To test whether these loci actually encode lincRNAs, we designed genomic microarrays (at 10-base resolution) across 1,147 of the 1,703 loci (1) to determine their exonic structure. We hybridized poly(A)-amplified RNA from hES, brain, breast, hEK, hFF, hLF, hK562, ovary, skin, spleen, testis, and thymus tissues. We analyzed the hybridization data using our previously reported peak-calling algorithm. This analysis revealed multixonic RNA transcripts in 74% of the K4-K36 domains examined (Fig. 1A). There was an average of 4 exons per K4-K36 domain (total of 4,860 exons).

![Fig. 1. Intergenic K4-K36 domains in the human genome produce multiexonic noncoding RNAs. (A) Representative example of an intergenic K4-K36 domain for lincFOXF1. For each histone modification (K4me3, green; K36me3, blue), the results of Chip-sequence experiments are plotted as the number of DNA fragments obtained by Chip-sequence at each position divided by the average number across the genome. Intergenic K4-K36 domains were interrogated for presence of transcription by hybridizing RNA to DNA tiling arrays. The resulting RNA hybridization intensity (red) within each K4-K36 domain is plotted with respect to its genomic location. The start and stop of each exon, as determined by our RNA peak calling algorithm (SI Methods), is indicated by a black bar. Arrowheads indicate the orientation of transcription. (B) Sequence conservation scores (SI Methods) across 21 mammalian species indicates lincRNAs (blue) are much more conserved than neutrally evolving intronic sequences (red), although less so than protein-coding genes (green). For each lincRNA exon, protein-coding gene exon, and protein-coding gene intron, a conservation score was calculated and plotted along the x axis as a log-odd enrichment score (compared with random genomic regions of equivalent size). The cumulative number of exons with a given score or lower is represented on the y axis.

Combing these data, we identified lincRNA loci that encode protein-coding genes. We found that the vast majority of the full set of 1,703 loci encode bona fide lincRNAs. We then studied the evolutionary conservation of the lincRNA loci. For each exon, we calculated the extent of sequence conservation across 21 mammalian species as previously described (SI Methods) (1). Human lincRNAs showed evolutionary conservation at levels similar to those seen for the protein-coding loci in our previous study (Fig. 1B; table 2 in Dataset S1) (1).

**Dataset S1.** This dataset contains all the accession numbers for the lincRNA loci, the exons detected in the exome array data, and the conservation scores for the exons across the 21 mammalian species.

**Dataset S2.** This dataset contains all the accession numbers for the protein-coding loci, the exons detected in the exome array data, and the conservation scores for the exons across the 21 mammalian species.
have expression patterns that are more tissue- or condition-specific, the total number could be considerably higher. Obtaining a complete catalog will require generating chromatin-state maps across many more tissues.

**Many lincRNAs Are Associated with PRC2.** We next explored the mechanism by which lincRNAs function. As noted above, the lincRNA HOTAIR has been shown to physically associate with PRC2 (8). This physical association was shown by RIP-PCR assay: total (non-cross-linked) nuclear extract was incubated with an antibody against the SUZ12 protein, a component of PRC2; the extract was precipitated with Protein-A-coupled beads; and the coprecipitated RNA was hybridized to the hybridization data with a peak-calling algorithm that finds regions in which the signal from the RIP assay is significantly enriched over the signal from the mock controls (SI Methods). Regions were defined based on a maximum familywise error rate (FWER) < 0.05 (SI Methods and table 3 in Dataset S1) (1). Given that RIP assays are known to show considerable variability (with typical reproducibility of ~60%) (8), we performed several biological replicates for each cell type. We observed that ~76% of the genes detected in one replicate are also detected in a second replicate (hLF, 70%; hFF, 75%; HeLa, 83%; see table 3 in Dataset S1). As a positive control, we checked whether HOTAIR and XIST were detectably coprecipitated in our RIP-Chip data. Consistent with previous reports, HOTAIR coprecipitated with PRC2 in both HeLa and hFFs, but not in hLFs. Similarly, XIST, which is expressed only in female cells, was detectably coprecipitated in the hLF cells (which came from a female source), but not the hFF cells (which came from a male source) (Fig. 2). These results were consistent across all replicates.

In addition to the RIP assay, we also assayed expression patterns of lincRNAs and protein-coding genes on the custom exon-tiling array. We extracted total RNA from the same 3 human cell types (HeLa, hLF, and hFF), performed a poly(A+) amplification reaction, and hybridized the product to the exon-tiling array. Of the lincRNA genes on the array, we found that 47% were detectably expressed in at least 1 of the 3 cell types (HeLa, 25%; hLF, 37%; and hFF, 33%; see table 4 in Dataset S1). Consistent with the design of the tiling array, essentially all of the protein-coding genes were detectably expressed in the relevant cell type. Analysis of the RIP-Chip results, in conjunction with the expression analysis, suggests that a significant proportion of all lincRNAs expressed in 1 of these 3 cell types are physically associated with PRC2. Specifically, we find that ~30% of expressed lincRNAs are detected in at least 1 of the replicates. As a conservative estimate, we only considered lincRNAs detected in at least 2 replicates. Using this criterion, we observe that 24% of lincRNAs (114 of 469) expressed in 1 of the 3 cell types is detected as physically associated with PRC2 (Fig. 2; Fig. S2).

As an independent validation of the association with PRC2, we selected 5 lincRNAs that were detected in our RIP-Chip data as associated with PRC2 in both HeLa and hFF, and performed RIP-quantitative (q)PCR assays for these transcripts, using qRTPCR. In all 10 tests (5 lincRNAs in 2 cell types), the results were confirmed (Fig. S3 and table 5 in Dataset S1). Notably, the RIP-qPCR assays showed a higher degree of enrichment than the RIP-Chip assays, consistent with the fact that arrays have a narrower dynamic range. As a validation that the associations of lincRNAs with PRC2 are specific, we tested whether the enrichment in the RIP-Chip experiment was simply a reflection of transcript abundance, which would suggest nonspecific interactions. We found no significant correlation between transcripts levels of the lincRNAs and their level of PRC2-enrichment ($r = -0.109, P > 0.99$; Fig. S4).

As a second approach to assess the specificity of PRC2 binding to lincRNAs, we examined the proportion of mRNAs bound to PRC2. In sharp contrast to the lincRNAs, very few of the mRNAs assayed in the RIP-Chip experiments showed physical association with PRC2. Of the 1,000 mRNAs represented on the array, only 16 (~2%) were detected in 2 replicates (Fig. 3A). We suspect that
many of these 16 cases are artifacts, because only a small proportion (<1% of expressed mRNAs) are detected in 3 replicates. Thus, the proportion of transcripts associated with PRC2 is much higher for lincRNAs than for mRNAs. To demonstrate that this result is not simply due to a low concentration of mRNAs in the nucleus, we find that the abundance of lincRNAs in the nucleus (compared the concentration of lincRNAs and mRNAs in the cytoplasm) is much higher for lincRNAs than for mRNAs. To demonstrate that this result is not due to a lower concentration of lincRNAs in the nucleus, we have shown that depletion of HOTAIR causes up-regulation of chromatin proteins themselves (rather than chromatin-modifying proteins) are associated with lincRNAs. Specifically, we performed RIP-Chip with antibodies against the modified histones H3K27me3 and H3K4me2. We found no significant enrichment of lincRNAs (Fig. 2). These findings are consistent with other studies that identified XIST to coprecipitate with PRC2, but not H3K27me3, despite their immediate nuclear proximity (11).

Functional Evidence That lincRNAs Act Through the PRC2 Pathway. Having found that a substantial fraction of lincRNAs is physically associated with PRC2, we sought evidence that they have a functional role in polycomb-mediated repression. Previously, we have shown that depletion of HOTAIR causes up-regulation of genes normally repressed by PRC2 (8). To test whether other lincRNAs have a similar effect, we studied HOTAIR and 5 additional lincRNAs found to be associated with PRC2. For each of these 6 lincRNAs, we designed pools consisting of 4 siRNAs targeting each lincRNA (SI Methods). We also used standard control siRNA pools that do not correspond to any human sequence. We transfected the siRNA pools into hLF (3 pools), hFF (3 pools), or both (1 pool), with each experiment performed in duplicate. We measured the level of lincRNA knock-down by qRT-PCR, and compared the results to the control siRNA pool; we only used experiments in which we achieved >2-fold depletion (Fig. S7).

We hybridized the total RNA from these experiments to standard gene-expression arrays to measure the resulting changes in gene expression. Specifically, for each of the 6 lincRNAs, we determined the gene sets \( S_1, S_2, \ldots, S_6 \) that were up-regulated relative to the control siRNA pools [at a false discovery rate (FDR) \( < 0.1 \)]. These sets contained between 103 and 352 genes (Fig. 4A; Fig. S8A). The sets of genes affected by each lincRNA did not show significant overlap, suggesting that each lincRNA has distinct target sets. We searched for, but found no common motifs enriched among the up-regulated genes for each lincRNA. However, given the small number of target genes and the inability to...
Results are shown from knockdown experiments in hLFs (gray) and in hFFs (black).

An Example: TUG1 Represses p53-Dependant Cell-Cycle Regulation.

Last, we decided to focus on a specific PRC2-associated lincRNA, TUG1. TUG1 was originally identified as a transcript up-regulated by taurine, and siRNA-based depletion of TUG1 in the developing mouse eye was found to block retinal development (20); the mechanism by which TUG1 depletion produces this phenotype is unknown. In our study, we found that TUG1 is ubiquitously expressed in human and mouse cell types and tissues, and is bound to PRC2 in all 3 of the cell types examined. Previously, we studied regulation of lincRNAs in response to DNA damage, and found that TUG1 was among the 39 lincRNAs specifically induced in p53-wild type, but not p53-mutant cells (Fig. 4B) (1). Also, TUG1 promoter contains many highly conserved binding sites for p53 (Fig. S8). We selected TUG1 as one of the 6 lincRNAs above that we depleted with siRNA pools. Depletion of TUG1 led to significant up-regulation of 120 genes, which were strongly enriched for those involved in cell-cycle regulation (regulation of mitosis, spindle formation, and cell-cycle phasing) (Fig. 4C; table 7 in Dataset S1). Thus, TUG1 is induced by p53, binds to PRC2, and has a role in repressing specific genes involved in cell-cycle regulation. Interestingly, p53 is well known to cause both activation and repression of many genes. Although p53 has been shown to be a direct activator of many genes, the mechanism of p53-induced repression remains unknown. Our results suggest the intriguing hypothesis that TUG1, and perhaps other lincRNAs, may function as downstream repressors in transcriptional pathways.

Discussion

It is becoming clear that the mammalian genome encodes thousands of lincRNAs that are highly conserved and, thus, biologically functional (1, 27). The results of our previous article (1) and this study together identify 3,289 lincRNAs, and suggest that the total may be in the range of ~4,500. Expression patterns suggest that these lincRNAs are involved in diverse biological processes, including cell-cycle regulation, innate immunity, and ES pluripotency, but the mechanisms by which they have their roles were completely unknown. Inspired by studies of the lincRNAs HOTAIR (8) and XIST (11), we investigated the idea that many lincRNAs are involved in the establishment of chromatin states. In this study, we report that a substantial proportion (24%) of lincRNAs expressed in a cell type are physically associated with the repressive chromatin-modifying complex PRC2, and the proportion is even larger (38%) when additional chromatin-modifying proteins (CoREST and SCMX) are included. Thus, it seems likely that a significant fraction of lincRNAs will be associated with chromatin-modifying proteins. Beyond the physical association, our functional analysis demonstrates that siRNA-mediated depletion of these lincRNAs results in preferential derepression of PRC2 regulated genes at distant loci, consistent with a trans-acting mechanism. Collectively, these results suggest that many lincRNAs collaborate with chromatin-modifying proteins to repress gene expression at specific loci.

There is a growing body of literature from yeast to mammals suggesting the noncoding RNAs have an important role in chromatin-state formation (4, 21). In Schizosaccharomyces pombe, a
process known as RNA induced transcriptional silencing (RTS) has been shown to have an important role in heterochromatin formation over centromeric repeats (22). Similarly, short RNAs have been shown to have an important role in the establishment of heterochromatin in plants. In Caenorhabditis elegans, genetic screens have identified polycomb homologs to be required for proper gene silencing in an RNA-dependent manner (4). In mammals, only a few specific RNAs (such as HOTAIR and XIST) have been implicated in directing chromatin modification. However, there is evidence that RNA has a key role in shaping mammalian epigenetic landscapes. For example, depletion of single-stranded (ss)RNA in mouse fibroblasts inhibits global heterochromatin formation (23). Similarly, ssRNA, but not ssDNA, is required for the maintenance of the histone modifications H3K27me3 and H3K9me3 (24).

Our results suggest an intriguing hypothesis that lincRNAs bind to chromatin-modifying complexes to guide them to specific locations in the genome. Whereas chromatin-modifying proteins are often ubiquitously expressed, they establish epigenetic states that differ markedly among cell types and conditions. Under our model, differentially expressed lincRNAs could bind to these complexes and help establish cell type specific epigenetic states. In particular, PRC2 is involved in establishing repressive chromatin states involving H3K27me3. Together, PRC2 and a lincRNA might have the role of a transcriptional repressor by directing silencing to specific loci. Such a mechanism could function within a larger regulatory program. Specifically, a newly induced transcription factor might establish a particular cellular state by (i) directly activating some downstream genes, and (ii) activating lincRNAs that (with PRC2) repress genes involved in a previous or competing cellular state. Our observations concerning the lincRNA TUG1 suggest that it may function in such a program. On DNA damage, TUG1 is induced in a p53-dependent manner, likely through direct binding of p53, in view of many p53-binding sites in its promoter. It then binds PRC2 (based on our RIP-Chip data) and is involved in repressing important cell-cycle related genes (based on siRNA-based depletion of TUG1). Thus, we speculate that TUG1 may serve as a downstream transcriptional repressor in the p53 pathway to repress cell-cycle progression in response to DNA damage. Similarly, we have recently shown that HOTAIR serves as a transcriptional repressor of HOXD genes. We now know that HOXA13, the key distal regulator, directly transcribes HOTAIR to establish positional identity by repressing the appropriate HOX clusters. Thus, HOTAIR serves as a downstream repressor in the HOXA13 transcriptional network. This model raises many mechanistic questions, including (i) whether most lincRNAs associated with chromatin-modifying complexes directly guide the complexes to specific loci, and (ii) if so, how the guidance is accomplished (e.g., by direct binding at specific sequence motifs).

Our experiments have focused on chromatin-modifying complexes that add repressive chromatin marks. It is possible that many additional lincRNAs are associated with chromatin-modifying complexes that confer activating modifications, as has been recently reported in a few cases (14). These questions can be addressed by performing RIP experiments with a wide range of antibodies across a wide range of cell types, to create a catalog of lincRNA-protein interactions. Last, although we have found that a substantial proportion of lincRNAs are associated with repressive chromatin-modifying complexes, we do not mean to suggest that all lincRNAs necessarily function in this manner. There may be classes of lincRNAs that function in entirely different ways. For example, the lincRNAs NEAT1 and NEAT2 have been recently shown to be important in the formation of paraspeckles (25), and the lincRNA NRON has a role in repressing nuclear import (26). It is possible that additional lincRNAs have roles in these and numerous other cellular pathways. The full range of biological diversity of lincRNAs and their mechanisms clearly remains to be explored.

Materials and Methods

Identification of K4-K36 domains was performed as previously described (1). RIP was performed as previously described (8) with some modifications. Hybridization to tiling arrays was performed as previously described (1). For detailed methods, see SI Methods. The data concerning the lincRNAs and the experiments here are available in Dataset S1 and public databases. All microarray data including RNA hybridization to tiling arrays, RIP-Chip experiments, and gene expression profiling of lincRNA knockdowns is deposited at the Gene Expression Omnibus (GEO) under accession no. GSE16226.

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