Promoter Analysis Reveals Globally Differential Regulation of Human Long Non-Coding RNA and Protein-Coding Genes

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Abstract

Transcriptional regulation of protein-coding genes is increasingly well-understood on a global scale, yet no comparable information exists for long non-coding RNA (lncRNA) genes, which were recently recognized to be as numerous as protein-coding genes in mammalian genomes. We performed a genome-wide comparative analysis of the promoters of human lncRNA and protein-coding genes, finding global differences in specific genetic and epigenetic features relevant to transcriptional regulation. These two groups of genes are hence subject to separate transcriptional regulatory programs, including distinct transcription factor (TF) proteins that significantly favor lncRNA, rather than coding-gene, promoters. We report a specific signature of promoter-proximal transcriptional regulation of lncRNA genes, including several distinct transcription factor binding sites (TFBS). Experimental DNase I hypersensitive site profiles are consistent with active configurations of these lncRNA TFBS sets in diverse human cell types. TFBS ChIP-seq datasets confirm the binding events that we predicted using computational approaches for a subset of factors. For several TFs known to be directly regulated by lncRNAs, we find that their putative TFBSs are enriched at lncRNA promoters, suggesting that the TFs and the lncRNAs may participate in a bidirectional feedback loop regulatory network. Accordingly, cells may be able to modulate lncRNA expression levels independently of mRNA levels via distinct regulatory pathways. Our results also raise the possibility that, given the historical reliance on protein-coding gene catalogs to define the chromatin states of active promoters, a revision of these chromatin signature profiles to incorporate expressed lncRNA genes is warranted in the future.

Introduction

Evidence for important, including essential, cellular and organismal roles of lncRNA in mammalian systems began to emerge prior to the advent of high-throughput genome and transcriptome sequencing. These early examples included the demonstration that the lncRNA XIST [1] was necessary and sufficient for X-chromosome silencing, as well as the discovery of SRA [2], an lncRNA that directly regulates the estrogen receptor α, one of the nuclear hormone receptors. Other essential functional ncRNAs in eukaryotic cells, such as ribosomal, transfer, and spliceosomal RNAs, have been well-known for an even longer time. Although the human genome project [3] initially focused almost exclusively on protein-coding genes in the human gene count, the ubiquity, in addition to the existence and the functional significance, of mammalian lncRNAs has been a key revelation of transcriptome sequencing projects [4].

Many lncRNA transcripts, similarly to mRNAs, are 5'-capped, polyadenylated, frequently spliced with conventional GT-AG intron excision, and readily evident in cytoplasmic polyA+ RNA preparations; thousands of lncRNAs have been discovered from cDNA libraries [5], although abundant nuclear and polyA- lncRNAs have also been identified [6]. Up to one-third of polyA+...
lncRNAs encoded in the human genome may not be evolutionary conserved beyond primates [4]. In contrast, the majority of human protein-coding genes have pan-mammalian, and usually pan-vertebrate, conservation, many with homologs identifiable throughout metazoan. It has been suggested that non-conserved lncRNAs comprise a part of the molecular basis of species phenotypic uniqueness, distinguishing closely related species from one another by providing substrates for exaptation as well as adaptive evolution [7]. Despite their frequent lack of conservation, overwhelming evidence of lncRNA functions has emerged: they are characterized by diverse, positive and negative, nuclear and cytoplasmic, epigenetic and post-transcriptional regulatory modalities. Documented lncRNA functions include: positive regulation of sense mRNA translation by an antisense lncRNA [8], transcriptional repression of mRNAs by repeat-containing lncRNAs through the Staufen-1 mRNA decay pathway [9]; epigenetic regulation of protein-coding targets by lncRNAs that recruit PRC2 to gene promoters [10], and direct RNA-protein interactions between lncRNAs and TFs: the Evt-2 lncRNA directly interacts with distal less homeobox proteins to regulate mouse hippocampal development [11]. The Gas5 lncRNA contains a precise ribomimic of the genomic DNA binding site of the human glucocorticoid receptor, therefore titrating out bioavailable glucocorticoid receptor molecules and preventing them from binding their cognate sites in gene promoters along genomic DNA [12]. More generally, endogenous riboregulation of DNA-binding NRs through direct interactions with lncRNAs [2,12,13] is an emerging leitmotif of post-genomic lncRNA biology.

These diverse functional mechanisms summarily indicate that jointly with TFs, lncRNAs are key regulators of protein-coding genes – including those that encode TFs. A prerequisite toward understanding the biology of lncRNAs is their assignment into tractable gene regulatory networks. We previously showed [14] that TFs – in particular, Oct4 and Nanog, which are essential for stem cell pluripotency [15] – bind directly at the promoters or within gene bodies of hundreds of lncRNA genes. ChIP-qPCR validation of TF binding to lncRNA gene promoters has elucidated numerous targets of key TFs, including non-conserved lncRNAs repressed by REST/NRSE in the human DiGeorge Syndrome critical region and in mouse [16]. We have used forward and reverse genetics to validate the regulation of lncRNAs by these TFs, uncovering feedback loops in the network that also use the lncRNAs to regulate these TFs during cell lineage specification [14]. More recently, we have assigned lncRNAs into deterministic regulatory networks, using reverse genetic approaches to show that a primate-specific antisense lncRNA regulates neuronal activity-dependent epileptogenesis in the m vivo human brain [17]. However, despite this progress, a genome-wide understanding of the lncRNA regulatory network – including the characterization of TF/lncRNA interactions – has to date remained elusive.

In this study, our goal was to computationally test the hypothesis that the global transcriptional regulatory programs of lncRNA genes and protein-coding genes are different. We set this problem within the framework of machine learning classification of promoters of these two broad gene classes. Previous studies [18–21] used support vector machines to distinguish non-coding RNAs (ncRNAs) from mRNAs, whereas experimental approaches including Riboseq [22] and mass spectrometry [23] have documented that lncRNAs possess a low affinity for ribosomes and are rarely translated, but no comparable efforts have been devoted to comparing lncRNA and protein-coding gene promoters. Recently Lv et al. [24] used chromatin modification and genomic features to distinguish lncRNAs from protein-coding genes, while a statistical approach [25] singled out H3R2me1 as a distinctive histone mark between protein-coding genes and lncRNAs. Here, we interrogated multiple computational and empirical sources of regulatory information at promoters on a genome-wide scale. We found genetic and epigenetic signatures unique to protein-coding and lncRNA genes, respectively. These divergent promoter grammars may help to explain the observed differential and highly tissue- and condition-specific transcriptional regulation of lncRNA genes compared to their protein-coding counterparts in the same pathways. To our knowledge, this is the first demonstration that human lncRNA and protein-coding gene promoters contain sufficiently dissimilar information to be consistently distinguished with high accuracy. Our results sum- marily suggest the existence of distinct regulatory programs for these two gene groups.

Results

DNA sequence patterns at the promoters of protein-coding and lncRNA genes

We compared DNA sequence promoter patterns of protein-coding and lncRNA genes. We found that A/T-rich mono-, di- and tri-nucleotide patterns are enriched at the promoters of lncRNA genes, relative to the promoters of protein-coding genes (“differentially enriched at lncRNA promoters”) (Table S1). CpG-derived mono-, di- and tri-nucleotide patterns are overrepresented in promoters of protein-coding genes. This result is broadly consistent with the observation that AT-rich promoters demonstrate lower expression but higher tissue specificity [26], properties known to define lncRNA promoters [4]. CG-skew, a feature of protein-coding gene promoters, is significantly reduced in lncRNA gene promoters, while AT-skew is almost depleted (Figure 1a–b). Figure 1c shows that word commonality score (Text S1 Methods section) is decreased around the transcriptional start sites (TSS) of lncRNA genes, although this depletion is stronger around TSSs of protein-coding genes, suggesting that lncRNA gene regulation, in contrast to protein-coding genes, is less driven by unique recognition sequences.

Palindromes, widespread regulatory elements in the promoters of protein coding genes [27], are less frequent around TSSs of lncRNA genes (Figure 1d). CpG islands (CGIs) are known to overlap with about two-thirds of protein-coding gene promoters [28]. Although CGIs are also hosting numerous non-coding transcripts [29,30], an observation independent of the method of CGI detection [31], such ncRNAs are most likely short and unprocessed. On the contrary, we find that lncRNA promoters quite rarely overlap with CGIs (Figure 1e). LncRNA exons and splice junctions have been reported as enriched in repetitive elements [32]. We show that repetitive elements are also enriched at lncRNA promoters (Figure 1f, Figure S1). DNA sequence properties of non-zero similarly expressed protein-coding and lncRNA genes show feature patterns similar to those of the whole promoter sets without considering any expression levels (Figure S1).

Known TFBSs and novel motif families distinguish the promoters of lncRNA genes

We in silico predicted the incidence of known transcription factor binding sequences (TFBSs) at the promoters of both gene types, using the HOCOMOCO [33] human TFBS models database. We found 74 TFBSs overrepresented in protein-coding gene promoters and 140 TFBSs overrepresented in lncRNA gene promoters (“differentially enriched in lncRNA promoters”) (Table S2).
Several TFs regulated by specific lncRNAs emerge as potential global regulators of lncRNA transcriptome in our analysis. A representative example is PGR (progesterone receptor), a nuclear hormone receptor (NHR), whose predicted TFBSs are differentially enriched at lncRNA promoters. The human PGR gene itself is cis-regulated by two lncRNAs: an lncRNA containing primate-specific repetitive elements provides transcriptional regulation [34,35] and another cis-antisense transcript acts post-transcriptionally [36]. Here, we show widespread genome-wide association of lncRNA promoters with the same TF families that have been previously implicated as regulatory targets of lncRNAs. The human NHR superfamily provides the most abundant evidence of preferential involvement in genome-wide lncRNA cis-regulatory programs: the TFBSs of 13 (27%) of the 48 total known human NHRs (PGR, NR1I2, NR1I3, NR2C2, NR2E3, NR5A2, RARG, ESR2, PPAR, HNF4A, RXRB, ERR1, and ERR2) were differentially enriched at lncRNA promoters.

We additionally found that 14 FOX-family TFs, 6 SOX-family TFs, 3 members of the HOXD homeobox family, 3 members of the CEBP family, 3 NKX-family TFs, and 2 PPAR TFs (Table S2) demonstrate similar patterns of differential TFBS enrichment at lncRNA promoters. Several of these same TFs have been previously reported as regulatory targets of lncRNAs as well [41]. Our in silico predicted binding site results for homeobox TFs at the promoters of lncRNA genes are consistent with a recent evolutionary study [42]. Summarily, the TF families that are characterized by TFBS enrichments at lncRNA promoters in our analysis include TFs that are known to be direct targets of lncRNAs from prior mechanistic studies.

The human proteome harbors approximately 1500 TFs [43], although TFBS models are available through HOCOMOCO for only 401 TFs. To compensate for this and to allow the detection of TFBSs whose motifs remain unknown, we applied ab initio motif discovery to genome-wide promoters, in order to complement the HOCOMOCO results. Ab initio identified motif families (MFs) generated by the Dragon Motif Finder [44], suggest multiple levels of sequence complexity specific to lncRNA promoters. These include reverse-complement motifs (palindromes) unique to lncRNA promoters, long motifs (20 bps), and polyA/polyT-rich regions (Figure S2a–d).

Condition-specific binding preferences are an important biological property of certain TFs [45]. Polymorphisms and de novo mutations may also alter a sequence of a particular binding site complicating known-TFBS discovery [46,47]. Hence, we reasoned that certain ab initio MFs might reflect condition-driven, or protein complex-dependent, deviations from known TFBS models. We therefore compared the ab initio identified MFs to those already associated with known TFs. We confirmed five

Figure 1. DNA feature distributions in the promoters of lncRNA genes and protein-coding genes. DNA feature distributions in a sliding window of 100 bp with a step of 50 bp in the promoters of protein-coding and lncRNAs. Blue line corresponds to promoters of protein-coding genes; red line corresponds to lncRNA gene promoters. Figure 1a–d shows distribution of the feature in a sliding window of 100 bp with a step of 50 bp, resulting in 39 windows on the plot. Figure 1e–f show the percentage of promoters where features were found. Transparent regions correspond to 5–95% bootstrap confidence interval of the statistics. WC: word commonality, PALIN: palindromes, CGI: CpG Islands, RE: repetitive elements, all types of repeats except "simple repeats", "low complexity regions" and "satellite repeats". The enrichment score was calculated using right-sided exact Fisher’s test (Table S3). doi:10.1371/journal.pone.0109443.g001
models and added one new TFBS model (NKX3-2) to our roster of lncRNA-promoter-enriched TFBSs (Table S2, Figure S2e–f).

**Chromatin configuration of lncRNA and protein-coding gene promoters**

To test whether lncRNA and protein-coding gene promoters possess different epigenetic signatures, we compared the genomic overlap of the two promoter types with defined chromatin states (CSs) in eight human cell lines [48].

Protein-coding gene promoters more often overlapped CSs associated with active, weak or inactive/ poised promoters, and were also more strongly enriched for Polycomb-repressed regions. Relative to protein-coding gene promoters, those of lncRNA genes more often overlapped CSs associated with insulators, regions of transcriptional transition (regions located between the initiation and elongation histone marks), elongation, weak transcription and heterochromatin (Figure S3, Table S3). After the subsets of lncRNA and protein-coding genes with similar expression levels in different cell lines were selected (see Methods), the same tendency remains but the difference between the promoters of protein-coding and lncRNA genes becomes less pronounced (Figure S5, Table S6). The role for enhancer-associated lncRNAs in regulating protein-coding genes over large genomic distances was recently reported [49]. Our data shows that in genes with non-zero similar expression levels most of the enhancer states are overrepresented in lncRNAs vs protein-coding genes (Figure S5), while in six out of eight studied cell types for all (independent of the expression levels) promoters only one out of four enhancer-associated CSs (weak enhancers) displays significant overrepresentation at lncRNA versus protein-coding gene promoters (Text S1 Results section).

To understand the biological context of the heterochromatin CS enrichment at lncRNA gene promoters, we analyzed histone modification marks (HMs) in the ENCODE Tier 1 cell line GM12878. LncRNA gene promoters were significantly depleted of almost all histone modification marks, except for H3K27me3 and H3K9me3 (Figure 2, Figure S4). H3K27me3 contributes to maintenance of 'bivalent domains', transcriptionally-poised regions combining activating and repressing histone marks [50,51], suggesting that lncRNA promoters are not permanently repressed and could be subject to activation under specific conditions. H3K9me3 marks transcriptional repression [52] but is also found in certain transcribed regions [53], and may be involved in elongation [54]. After the subsets of lncRNA and protein-coding genes with non-zero and similar expression levels in different cell lines were selected, lncRNA gene promoters demonstrated enrichment for H3K9me3 and surprisingly for H3K36me3 in all tested cell types. H3K36me3 is a mark of transcriptional elongation [55,56]. Interestingly, lncRNA gene promoters demonstrate a decreased level of H3K27me3 and, in H1-hESC, an increased level of H3K27ac, a mark of active promoters and enhancers [57]. Taken together, these results support active chromatin organization of lncRNA promoters, yet distinct from the one of protein-coding genes.

**Distinguishing promoters of protein-coding and lncRNA genes through an ensemble of decision trees model**

Several lines of evidence indicate that the transcriptional regulation of lncRNAs may differ substantially from that of protein-coding genes. To computationally test for any evidence of this phenomenon, we leveraged recent advances in machine learning to fit an integrative model based on the information from all analyzed data types to distinguish the promoters of protein-coding genes from those of lncRNAs. Our fitted ensemble model correctly classified the promoters (lncRNA or protein-coding) with more than 80% accuracy. Hence, across the majority of the genome sequence space, genetic and epigenetic information is sufficient to confidently separate these two classes of promoters (Table 1, Table S4). Interrogation of our fitted models revealed that the strongest effects accounting for this predictive power are DNA k-mers and CSs. These were more discriminative than TFBSs, although most feature types, including TFBSs, had significant discrimination power (Figure 3).

Since we had originally considered the regions of [-1000, +1000] bp around the TSS (Dataset S2) as a putative promoter region, for protein-coding genes we might have included some coding exonic sequences, therefore introducing coding sequence bias. To avoid this, we also performed the analysis (Text S1 Methods section) using only upstream promoter regions ([−1000, 0] bp upstream of the TSS). Using this promoter set, we were able to distinguish between lncRNA and protein-coding gene promoters with 77% accuracy (Table 1, Table S3). Moreover, to avoid a bias caused by the more abundant presence of CGIs at protein-coding gene promoters, we built another model for the upstream promoter regions ([−1000, 0] bp, Dataset S2) having no overlap with CGIs (Text S1 Methods section). Although the performance of the model decreased, we were still able to distinguish between lncRNA and protein coding gene promoters with 71% accuracy (Table S3).

**Figure 2. Distribution of histone modification marks in the GM12878 cell line across lncRNA and protein-coding gene promoters.**  
Figure demonstrates fraction of all promoters covered by chromatin a particular mark. Blue line corresponds to promoters of protein-coding genes; red line corresponds to lncRNA gene promoters. Transparent regions correspond to 5–95% bootstrap confidence interval of the statistics.  
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Distinguishing promoters of protein-coding and IncRNA genes with similar expression levels

IncRNAs show lower expression in almost all cell types as compared to mRNAs [49]. Low and highly expressed promoters tend to have distinct epigenetic features [58,59]. Sequence specific differences of high and low expressed genes have been discussed for years [60]. To avoid a potential bias of differentiating low and highly expressed promoters rather than IncRNA and protein-coding promoters, we compared only the IncRNAs and protein-coding genes with similar expression level in several cell types. Our model achieves over 81%, 80%, 81% and 80% accuracy for GM12878, H1-hESC, K562 and HUVEC, respectively (Table 2) when expression levels were controlled for. If we exclude the CGIs and downstream regions from the consideration, the models still demonstrate over 76% accuracy. Importantly, the performance of the model has been increased as compared to the model with the same set of features, but without controlling for the expression levels (71%, see previous section, vs 76%). These results suggest that expression bias is very unlikely to contribute to the accuracy of the models.

Open chromatin and specific regulatory programs at IncRNA-enriched TFBSs

We aimed to assess the extent of experimental support for transcriptionally permissive chromatin configurations across all TFBSs enriched at IncRNA promoters. We reasoned that binding sites should have an open chromatin configuration in the cell or tissue types where binding occurs. We leveraged the empirical genome-wide catalog of DNase I hypersensitive sites, indicative of open chromatin, derived for 125 human cell types by the ENCODE Consortium [61]. We found that 67 of the 140 IncRNA-TSS-enriched TFBSs were significantly associated with hypersensitive sites in one or more cell types. This presence of DNase I hypersensitivity sites in IncRNA promoters supports the regulatory potential of such regions in at least one of the 125 studied cell types, despite the repressed chromatin conditions at their promoters in the eight cell types with available data in the CS analysis above.

In parallel, we overlapped IncRNA-promoter-enriched TFBSs with ENCODE ChIP-seq experimental evidence for the corresponding TFs across all ENCODE ChIP-seq datasets [62]. A moderate FDR approach (Benjamini-Hochberg procedure) identified three TFs – GATA3, ARID3A, and MEF2A – as being dually supported by HOOCOMOCO computational evidence of their TFBS overrepresentation at IncRNA promoters and by ENCODE ChIP-seq experimental evidence for their binding at IncRNA promoters genome-wide (Table S2). This intersection of TFBS overrepresentation at IncRNA promoters and empirical ChIP-seq support for the binding of these same TFs at those promoters provides important evidence that these three TFs may direct genome-wide IncRNA transcriptional programs in the ENCODE ChIP-seq-profiled cell types.

Discussion

We present the first genome-wide demonstration of a significant difference in sequence characteristics between the promoters of human IncRNA and protein-coding genes, suggesting distinct regulation of the two gene groups. In view of the frequent bidirectionality of human promoters that simultaneously give rise to protein-coding and IncRNA genes [63], the distinctions we find are all the more remarkable, since bidirectional promoters are counted by our approach as both protein-coding and IncRNA. We speculate that specific TFs may function as network nodes that not only accept directional edges from regulatory IncRNAs, but also serve as network hubs that extend multiple new directional edges toward other IncRNA genes whose promoters contain their cognate TFBSs. In particular, our study, for the first time, suggests that specific NRHs - members of the nuclear receptor family, which are already known to be targeted by IncRNA-protein interactions that join IncRNAs and NRHs in ribonucleoprotein complexes – in turn, may regulate IncRNA transcription through promoter binding. Among the other TFs we considered, GATA3, ARID3A, and MEF2A have the singular distinction of being significantly supported by all three lines of evidence: TFBS motif enrichment at our 18,000 IncRNA promoters, DNase I hypersensitive site overrepresentation at their TFBS-containing IncRNA promoters, and ChIP-seq experimental evidence of enriched binding at these promoters genome-wide, across the ENCODE DNase I and ChIP-seq-profiled cell and tissue types. GATA3, one of our most-enriched TFs at IncRNA promoters and an essential regulator of type 2 helper T-cell (Th2) cytokine production, is itself cis-regulated by an antisense IncRNA (GATA3-AS1), which is increased in patients with allergic rhinitis, a Th2-associated disease [64]. More recently, evidence for large-scale GATA3 regulation of IncRNAs associated with Th2 functions has emerged, and an IncRNA was assigned into a GATA3-containing regulatory network in Th2 cells [65]. Our results support large-scale regulation of IncRNA transcription programs by GATA3, and enhance the list of IncRNAs whose promoters may comprise GATA3 targets.

Our observation that IncRNAs might be selectively regulated by a distinct set of TFs has substantial implications for systems biology: cells are potentially capable of harnessing a defined subset of regulatory switches to toggle the expression levels of IncRNAs without altering mRNA levels. Most of the disease-associated sequence variants in the human genome are non-coding [66], necessitating an integration of IncRNA TSS and exon locations with the increasingly abundant common-variant Genome Wide Association Studies (GWAS), as well as throughout whole-exome and whole-genome resequencing datasets designed to capture rare, large-effect disease-associated variants. Our results empower the GWAS community to re-annotate cryptic disease-associated variants at in silico predicted TFBSs that we have linked to global catalogs of IncRNA promoters and to IncRNA regulatory programs modulated by specific TFs. By virtue of their TFBS localization, such variants may emerge as direct functional candidates.

Our IncRNA gene collection is a composite of three previously published IncRNA sources – Gencode [4], the Broad Institute [67], and our own [5] – and three additional public IncRNA collections (see Methods). The methods used in the compilation of these IncRNA collections rely on a combination of full-length cDNAs, deep-coverage RNAseq, targeted RT-PCR validation, and extensive manual curation. Therefore, the IncRNA genes that we used are largely as reliable in terms of their underlying evidence and annotation quality as protein-coding genes, and the differences we have uncovered relative to their protein-coding counterparts are not likely to be due to annotation disparities.

Until recently, only protein-coding gene sets were used in characterizing general promoter features. Therefore, some widely accepted promoter features and chromatin state signatures may be biased as a consequence of having been inferred from protein-coding genes. In this context, it is hardly surprising that certain sequence and epigenetic features, more specific for protein-coding genes, are less pronounced at IncRNA promoters, while the chromatin states associated with IncRNA promoters are predominantly labeled as inactive promoters. However, these labels were...
Table 1. Summary of the results for separation of promoters of protein-coding and lncRNA gene promoters using different combinations of features.

<table>
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<th>Considering [-1000,0] bp</th>
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<td></td>
<td>Sensitivity (%)</td>
<td>Specificity (%)</td>
<td>Accuracy (%)</td>
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doi:10.1371/journal.pone.0109443.t001
A non-redundant set of promoters for human protein-coding and lncRNA genes

We used RefSeq transcripts from the UCSC Genome Database (http://hgdownload.cse.ucsc.edu/goldenPath/hg19/database/refGene.txt.gz, download date: 14 January, 2013) for the human genome (version hg19). Out of the 44,140 transcripts, we considered only the 34,475 that were clearly protein-coding (i.e. having an NM RefSeq ID) and that were located on chromosomes 1–22, X, and Y. To construct a non-redundant (i.e. a single reference transcript per gene) set, we considered at least 1 bp overlap in the entire genomic span (including exons and introns along hg19 coordinates) among all transcripts located on the same strand in the same locus, and we randomly selected one transcript per locus. Through these filtering steps, we ultimately arrived at 18,789 protein-coding non-redundant representative transcripts conforming to our one-transcript-per-gene data structure (Dataset S1a).

We also assembled 18,490 (Dataset S1b) experimentally supported (with full-length cDNA, Broad Institute RNAseq, or Gencode-curated cDNA and/or expressed sequence tag, i.e. EST, evidence), non-redundant (with respect to genomic position and orientation) lncRNA genes from six published sources: 1) our manually annotated list of human lncRNAs that are supported by full-length cDNA clones from 5' cap-trapped, dT-primed libraries [5]; 2) the Broad Institute lincRNA resource consisting of transcript assemblies inferred from exhaustive RNAseq of a human tissue collection [67]; 3) the ENCODE Consortium’s official Gencode catalog [70] of human genes (www.gencodegenes.org), a manually curated list of coding and non-coding genes that applies a unified set of manual annotation and targeted validation standards to uniformly assign biotypes to all transcripts and genes throughout its coding and non-coding gene collections, precluding lncRNA-specific quality control bias.

For each gene’s representative transcript, we considered the [–1000, +1000] bp around the TSS as the putative promoter region, except in the specific analyses listed under Results where an alternate [–1000, 0] bp TSS set was used. We chose relatively large promoter regions with the purpose of incorporating alternative TSS, which in turn allowed us focus on gene-based rather than transcript-based analysis, since alternative promoter usage is a widespread phenomenon in human transcriptome [74, 75]. Although such promoters may incorporate some exonic sequence, it was shown that downstream elements also regulate transcription [76], and therefore including the first kilobase of gene bodies – provided that protein-coding gene properties such as codon bias are controlled for – can provide valuable regulatory
information in addition to that residing in the region upstream of the TSS. We obtained the promoter sequences using Galaxy (www.galaxyproject.org/).

Computational model to discriminate the promoters of protein-coding and lncRNA genes

To identify the regulatory patterns which may facilitate the computational discrimination between the promoters of protein-coding genes and lncRNA genes, we extracted features from several broad categories. These include various frequency-based properties of the promoters such as k-mers, word commonality, skew, palindromes; regulatory elements such as CpG islands, repetitive elements, TFBS found within the promoter regions; epigenetic features such as chromatin states and separate histone modification marks (see Text S1 Methods section). We used an ensemble of decision trees [77] to generate a classification model and estimate its accuracy with 20-fold cross-validation.

Transcription factor binding sites (TFBSs) enrichment

We predicted TFBSs using 426 position weight matrices (PWMs) for 401 human TFs from the HOCOMOCO [33] database (http://www.cbrc.kaust.edu.sa/hocomoco/Download.php) in the promoters of both protein-coding and lncRNA genes. Since the extent to which the original nucleotide composition of promoters is a cause or a consequence of the possible TFBS repertoires present in these promoters is unclear, we used the same strategy for both protein-coding and lncRNA promoters. For each PWM the threshold was set in the following way: for a random word generated by a background model (independent nucleotide distribution with nucleotide frequency of hg19) there was a fixed probability of 0.0005 to obtain the PWM score no less than the threshold. We generated 426 features using the binary value 0 or 1 (zero or non-zero hits above the threshold in a given promoter sequence in both strands). We selected significantly overrepresented TFBSs in promoters of protein-coding vs. promoters of lncRNA (and vice versa) gene sets (p-value \( \leq 0.05 \), right sided Fisher’s exact test with Benjamini-Hochberg multiple testing for controlling false discovery rate (FDR) [78]) (See Text S1 Methods section).

Expression analysis using RNA-seq data

We used RNA-seq data from Gm12878, H1-hESC, K562 and HUVEC cell lines to check the model performance, when expression levels of lncRNAs and protein-coding genes are similar. We used the mappings, provided by ENCODE (http://hgdownload.cse.ucsc.edu/goldenPath/hg19/encodeDCC/wgEncodeCshlLongRnaSeq/) and we quantified the expression levels as RPKM (read per kilobase of exon per million mapped reads) [79] using FluxCapacitor [80]. We excluded all the transcripts having RPKM = 0. To identify the lncRNA and protein-coding genes with similar expression distribution, for each lncRNA we selected a protein-coding gene with the nearest expression value (but not differing more than 1% of its expression level) (Text S1 Methods section). In this way we secured a one-to-one correspondence between lncRNA genes and protein-coding genes matching based on their expression level, thus avoiding any kind of possible expression bias between lncRNA and protein-coding genes (Figure S6, Dataset S3).

Compilation of a uniform list of synonymous human transcription factor names

We used the UniProt database (www.uniprot.org) and the GeneCards resource (www.genecards.org) to compile a comprehensive list of human transcription factors that accounts for all
name multiplicity, synonymy, and redundancy between the abbreviated transcription factor names used by HOCOMOCO (since Uniprot naming is in one-to-one relationship with the HOCOMOCO naming system) and ENCODE. We manually curated this list. We identified 106 (Dataset S1i) transcription factors common to both lists, and all searches for overlaps between HOCOMOCO computational TFBSs and ChIP-seq empirical TFBSs were performed using this list.

See Text S1 Methods section for additional information.

Supporting Information

Figure S1 DNA feature distributions in a sliding window of 100 bp with a step of 50 bp in the promoters of protein-coding and lncRNAs for complete promoter set (CPS). Green line corresponds to promoters of protein-coding genes; black line corresponds to lncRNA gene promoters. Sub-figure. a-d show distribution of the feature in a sliding window of 100 bp with a step of 50 bp, resulted in 39 windows on the plot. Sub-figure, e-f show the percentage of promoters where features were found. Transparent regions correspond to 5–95% bootstrap confidence interval of the statistics. WC: word commonality, PALIN: palindromes, CGI: CpG Islands, RE: repetitive elements. The enrichment score was calculated using right-sided exact Fisher’s test (Table S3). Figure I considers all protein-coding and lncRNA genes in CPS and Figure II–V shows the distribution for non-zero similarly expressed genes in cell specific manner.

Figure S2 Logos for over-represented ab initio identified motif families (MFs) from promoters of a) protein-coding genes in CPS, b) lncRNA genes in CPS, c) protein-coding genes in REFPS and d) lncRNA genes in REFPS. Logos for ab initio motif families (MFs), corresponding reverse complement (RC) MF and known TFBS match by TOMTOM system from promoters of c) lncRNA genes in CPS, f) lncRNA genes in REFPS.

Figure S3 Distribution of chromatin states in cell lines with normal karyotypes across promoters of protein-coding and lncRNA genes. Blue bar corresponds to promoters of coding genes from repeat-filtered promoter set (REFPS), green bar corresponds to promoters of coding genes from complete promoter set (CPS), red bar corresponds to promoters of lncRNAs from REFPS, and black bar corresponds to promoters of lncRNAs from CPS. This figure demonstrates fraction of all promoters overlapping with chromatin states. At the end of each bar 5–95% bootstrap confidence interval of the statistic is shown. AP: Active Promoter, WP: Weak Promoter, IP: Inactive Promoter, SE: Strong Enhancer, WE: Weak Enhancer, I: Insulator, TT: Transcriptional Transition, TE: Transcriptional Elongation, WT: Weakly Transcribed, PR: Polycrom Repressed, HC: Heterochromatin low signal, RP: Repetitive/Copy number variation.

Figure S4 Distribution of histone modification marks, modified histone H2AZ, CTCF, and the Polycomb-group protein (PBC2 complex component) EZH2 in cell lines across lncRNA and protein-coding gene promoters.

Figure S5 Distribution of chromatin states in cell lines with normal karyotypes across promoters of protein-coding and lncRNA genes with similar expression. Green bar corresponds to promoters of coding genes from complete promoter set (CPS), black bar corresponds to promoters of lncRNAs from CPS. This figure demonstrates percentage of all promoters overlapping with chromatin states. At the end of each bar 5–95% bootstrap confidence interval of the statistic is shown. AP: Active Promoter, WP: Weak Promoter, IP: Inactive Promoter, SE: Strong Enhancer, WE: Weak Enhancer, I: Insulator, TT: Transcriptional Transition, TE: Transcriptional Elongation, WT: Weakly Transcribed, PR: Polycrom Repressed, HC: Heterochromatin low signal, RP: Repetitive/Copy number variation.

Figure S6 Boxplot and Quartile-Quartile plot for expression value of protein-coding genes and lncRNA genes from complete promoter set (CPS) in different cell lines.

Table S1 Mono-, di-, and tri-nucleotides frequency and observed/expected ratio for both complete promoter set (CPS) and repeat-filtered promoter set (REFPS).

Table S2 Transcription factor binding sites overrepresented in promoters of protein-coding and lncRNA genes for complete promoter set (CPS) and repeat-filtered promoter set (REFPS) and support provided by DNAseI and ChIP-seq peaks.

Table S3 P-values of overrepresentation for chromatin states, CpG islands, repetitive elements and palindromes for complete promoter set (CPS) and repeat-filtered promoter set (REFPS).

Table S4 a. Summary of the results for separation of promoters of protein-coding and lncRNA genes using different combinations of features for the complete promoter set (CPS) and repeat-filtered promoter set (REFPS). For REFPS, we used all types of repeats except “simple repeats”, “low complexity regions” and “satellite repeats”. k-mer: mono-, di-,tri-nucleotide frequencies, CS: chromatin states, WC: word commonality, TFBS: transcription factor binding sites, CGI: CpG Islands, SKEW: A/T and C/G skews, PALIN: palindromes, RE: repetitive elements, COMBINE: combination of all types of features. b. Summary of the cross validation (CV) results for separation of promoters of protein-coding from lncRNA genes using all features (COMBINE) for complete promoter set (CPS) and repeat-filtered promoter set (REFPS).

Table S5 Results from execution of the computational model from promoters considering only upstream (−1000, 0] of TSS, as well as from promoters considering only upstream (−1000, 0] of TSS having no overlap with CGI.

Table S6 P-values of overrepresentation for chromatin states for similarly expressed genes promoter in complete promoter set (CPS).

Text S1 Supporting information for the methods applied and results obtained. The details of methods are described under Methods section. The details of results are described under Results section.
Dataset S1 Set of a) RefSeq and b) lncRNA transcripts with hg19 human genome assembly coordinates in BED format for complete promoter set (CPS). Set of c) RefSeq and d) lncRNA transcripts with coordinates from hg19 in bed format for repeat-filtered promoter set (REFPS). Set of e) RefSeq and f) lncRNA promoters ([-2500…+250]) with co-ordinates from hg19 in bed format for complete promoter set (CPS). Set of g) RefSeq and h) lncRNA promoters ([-2500…+250]) with co-ordinates from hg19 in bed format for repeat-filtered promoter set (REFPS). i) “ENCODEROCO_HOCOMOCO” mapping – excel sheet contains mapping of ENCODE transcription factor name and HOCOMOCO V.8 motif name. Excel sheet “track_MEF2A_chip”, “track_AR13A_chip” contains the track information for ENCODE Chip-seq supported TFBS in CPS for MEF2A,GATA3 and ARI3A respectively. (ZIP)

Dataset S2 Promoters, considering only upstream ([-1000, 0] bp), of a) RefSeq and b) lncRNA transcripts with hg19 coordinates in BED format for complete promoter set (CPS). Promoters, considering only upstream ([-1000, 0] bp), of c) RefSeq and d) lncRNA transcripts, having no overlap with CpG islands, with hg19 coordinates in BED format for CPS. (ZIP)

Author Contributions
Conceived and designed the experiments: VBB LL. Performed the experiments: TA. Analyzed the data: TA YAM HJ JBB LL. VBB. Contributed reagents/materials/analysis tools: HJ. Wrote the paper: TA YAM JBB LL. VBB. Developed one of the lncRNA datasets: HJ.

References

Regulation of IncRNA and Protein-Coding Genes