Exploring the stability of long intergenic non-coding RNA in K562 cells by comparative studies of RNA-Seq datasets

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Abstract

Background: The stability of long intergenic non-coding RNAs (lincRNAs) that possess tissue/cell-specific expression, might be closely related to their physiological functions. However, the mechanism associated with stability of lincRNA remains elusive. In this study, we try to study the stability of lincRNA in K562 cells, an important model cell, through comparing two K562 transcriptomes which are obtained from ENCODE Consortium and our sequenced RNA-Seq dataset (PH) respectively.

Results: By lincRNAs analysis pipeline, 1804 high-confidence lincRNAs involving 1564 annotated lincRNAs and 240 putative novel lincRNAs were identified in PH, and 1587 high-confidence lincRNAs including 1429 annotated lincRNAs and 158 putative novel lincRNAs in ENCODE. There are 1009 unique lincRNAs in PH, 792 unique lincRNAs were in ENCODE, and 795 overlapping lincRNAs in both datasets. The analysis of differences in minimum free energy distribution and lincRNA half-life showed that a large proportion of overlapping lincRNAs were more stable than the unique lincRNAs. Most lincRNAs were more unstable than protein-coding RNAs through comparing their minimum free energy.

Conclusions: Identification of overlapping and unique lincRNAs can be helpful to classify the stability of lincRNAs. Our results suggest that overlapping lincRNAs (relatively stable lincRNAs) and unique lincRNAs (relatively unstable lincRNAs) might be involved in different cellular processes.

Reviewers: This article has been reviewed by Prof. Oliviero Carugo, Dr. Alistair Forrest and Prof. Manju Bansal. 

Keywords: RNA Sequencing (RNA-Seq), Long intergenic non-coding RNAs (lincRNAs), K562, RNA stability

Open peer review

Reviewed by Prof. Oliviero Carugo, Dr. Alistair Forrest and Prof. Manju Bansal. For the full reviews, please go to the Reviewers’ comments section.

Background

The mammalian genome is extensively transcribed, giving rise to many thousands of non-coding transcripts including both short transcripts (<200 nucleotides in length) and long non-coding RNAs (lncRNAs) (>200 nucleotides in length) [1]. LncRNAs recently have caused more attention because they interact with other biological molecules to regulate diverse cellular processes [2]. According to the location and context in genome, lncRNAs can be classified into intergenic lncRNAs, intronic lncRNAs, sense lncRNAs and antisense lncRNAs [3]. Present studies are mainly focused on intergenic lncRNAs (lincRNAs) located and transcribed from intergenic genomic regions, not only because lncRNAs are more convenient for experimental manipulation and computational analysis without the interference of annotated protein-coding regions than other lncRNAs [4], but also because lincRNAs participate in many cellular processes from embryonic stem cell pluripotency to cell proliferation and cancer progression [5,6].
With large-scale transcriptome sequencing, a growing number of lincRNAs have been identified in mammals [4]. However, only a few of lincRNAs have been functionally characterized. The lincRNA HOTAIRM1 modulates the expression of several HOXA genes, which encode key transcription factors for differentiation of myeloblasts [7]. LincRNA-p21 plays a functional role in p53 response pathways through affecting the expression of numerous genes associating with p53, and recruits ribonucleoprotein K (hnRNP-K) to trigger the regulation of p53-mediated apoptosis [8]. LincRNA-EPS at the nuclear localization is up-regulated in murine erythroid terminal differentiation, which promotes survival of murine erythroblasts [9]. Overall, many lincRNAs can regulate gene expression through specific interactions with other cellular factors, DNA, and other RNA molecules, such as acting in cis on neighboring genes or acting in trans regardless of gene location [10,11]. Another proposal is that lncRNA genes (including lincRNA) act as enhancer regions and may be incidental by-products of transcription [12].

Previous studies have indicated that RNA stability influences the abundance of transcript and shapes the kinetics of gene induction in intricate gene networks in mammalian cells [13-16]. Therefore, lincRNA stability inevitably affects its function in post-transcriptional regulatory pathways. However, there have been only a few reports about lincRNAs stability [7-9,17,18]. Recently, Tani et al. determined the half-lives of transcribed RNAs in whole-genome of HeLa Tet-off (TO) cells by BRIC-Seq and found hundreds of short-lived noncoding transcripts (t1/2 < 4h) [19,20]. Clark et al. found that only a minority of lncRNAs were unstable by confirming the half-lives (half-life < 2h) of about 800 lncRNAs in the mouse Neuro-2a cell line with a custom non-coding RNA array [21]. These studies concluded that mRNAs and lncRNAs (including lincRNAs) have similar half-life distributions [11]. Furthermore, lincRNAs are transcriptionally activated similar to mRNAs, nearly always 5′-capped and 3′-polyadenylated, and are frequently spliced [3,11]. It has been known that short-lived mRNAs were enriched among genes with regulatory functions, whereas long-lived mRNAs with metabolism and structure [22,23]. Like that, lincRNA stability should be closely associated with its physiological function. Although different inhibitor approaches for identifying half-life have been documented in recent years, the stability of lincRNA is poorly understood. It limited functional research of lincRNAs in the complex post-transcriptional regulation.

The human leukemia K562 cells have the potentials for differentiating into erythroid, granulocytic, monocytic and megakaryocytic lineages [24]. It is an important model cell in studying the pluripotency and differentiation of hematopoiesis and leukemogenesis [25,26]. Hence, we investigated the stability of lincRNAs in K562 cells and comparatively analyzed lincRNAs in ENCODE and our dataset named as PH which was sequenced on an Illumina HiSeq™ 2000 with pair-end libraries by RNA-Seq technology. The goals were to (1) improve the lincRNAs analysis workflow to acquire stringent lincRNA set; (2) compare and analyze the lincRNAs in ENCODE and PH datasets; (3) compare the stabilities of coding-protein RNAs and lincRNAs; (4) display lincRNAs distribution in human genome. The results suggest that lincRNAs with different stability may have diverse functions in various biological processes.

**Results**

**Pipeline for lincRNA analysis**

In order to attain high-confidence lincRNA catalog, there are two main different improvements comparing to previous lincRNA predicting pipeline. (1) The more rigorous set of intergenic transcripts was acquired from the intersection of intergenic transcripts by comparing with four database annotations (Ensembl [27], UCSC [28], Gencode [29] and Refseq [30]), respectively. (2) Putative novel lincRNAs were more strictly attained by filtering coding potential RNAs of the results including iseeRNA [31], CPAT [32], CPC [33] and PhyloCSF [34] (see Methods). Through the lincRNA predicting pipeline, unreliable transcripts were removed and reliable lincRNAs were remained in downstream analysis.

**The features of lincRNAs in PH and ENCODE**

In PH, 1804 lincRNAs were identified, 1564 of which (86.7%) were annotated with the Ensembl or Gencode, and the remaining 240 (13.3%) were putative novel lincRNAs. There were 312 lincRNAs (17.3%) with FPKM (Fragments Per Kilobase of exon model per Million mapped fragments) ≥ 1, including 159 annotated lincRNAs and 153 putative novel lincRNAs. Simultaneously, 1587 lincRNAs were identified in ENCODE, 1429 of which (90.0%) were annotated with the Ensembl or Gencode, and the remaining 158 (10.0%) were putative novel lincRNAs. There were 514 lincRNAs (32.4%) with FPKM ≥ 1, including 455 annotated lincRNAs and 59 putative novel lincRNAs. LincRNAs in K562 cells, whether they were annotated or not in both datasets, were uniformly and prevalently distributed at every chromosome in human genome, even if most of the transcripts were low-abundant (FPKM <1) (Figure 1). By comparing lincRNAs of both datasets, 1009 lincRNAs were unique in PH, 792 lincRNAs were unique in ENCODE (unique lincRNAs present in only PH or ENCODE dataset), and 795 lincRNAs were overlapping in both datasets (overlapping lincRNAs present in both ENCODE and PH datasets) (Figure 2).
Comparison of minimum free energy and lincRNAs half-lives between PH and ENCODE

Whether lincRNAs expressed with FPKM $\geq 1$ or FPKM $< 1$, the minimum free energy of most overlapping lincRNAs were higher than unique lincRNAs (Figure 3). Although the expressed abundances of the majority of overlapping lincRNAs were higher than unique lincRNAs in the cases of FPKM $< 1$ (Figure 4), the minimum free energy and expression level (FPKM) are not correlated (Pearson's correlation coefficient $< 0.2$ in both datasets). Comparing the determined ncRNAs half-lives in Hela cells [19], 12 overlapping lincRNAs were found with half-life $> 4$ in both datasets, 2 unique lincRNAs in PH and 3 unique lincRNAs in ENCODE. In this study, random 10 overlapping lincRNAs half-lives (including FPKM $\geq 1$ and FPKM $< 1$, Figure 5) and random 9 unique lincRNAs half-lives (including FPKM $\geq 1$ and FPKM $< 1$ from PH or ENCODE, Figure 6) were determined by qPCR after ActD treatment [21]. The results proved that 10 overlapping lincRNAs half-lives are more than 4 hours (3 of which more than 30 hours), while 7 unique lincRNAs half-lives are less than 4 hours.
Comparison of the minimum free energy of the common protein-coding RNAs and lincRNAs

2914 protein-coding RNAs for the intersection of Cuffcompare’s results with four public database annotations (Ensembl, UCSC, GENCODE and RefSeq) were identified in PH, 2546 (87.4%) of which presented in ENCODE. Likewise, 795 lincRNAs (44.1%) in PH appeared in both datasets (Figure 7). By comparing the minimum free energy of the common protein-coding RNAs and lincRNAs of the both datasets, the result showed that the minimum free energy of a large proportion of protein-coding RNAs was significantly higher than lincRNAs (Figure 8).

Discussion

In this study, the lincRNA analysis workflow was improved to acquire stringent lincRNA set. The method to acquire the intersection of intergenic transcripts is simpler and more convenient than integrating several database annotations by their scripts [4,35]. Moreover, possible novel lincRNAs with uncertain coding potential (TUCPs) which possibly possess protein-coding or small peptides potential, were filtered by four softwares (iseeRNA, CPAT, CPC and PhyloCSF), respectively. Thereby more reliable putative novel lincRNAs were attained.

The genome encodes far more lincRNAs than previously known, and pervasively transcriptional lincRNAs might play widespread roles in gene regulation and other cellular processes [36]. In the study, thousands of lincRNAs including annotated lincRNAs and putative novel lincRNAs were identified in both datasets through our lincRNA predicting pipeline. And, those lincRNAs were transcribed from thousands of locations at every chromosome in human genome.

The sequence of nucleotides of an RNA molecule carries the functional information. As more lincRNAs are identified, it will become important to study lincRNA sequences and their secondary structures to reveal mechanisms of lincRNA functions through establishing structure-function relations [11,37-40]. These roles including sensory, guiding, scaffolding and allosteric...
capacities derive from folded modular domains in lincRNAs [41]. Minimum free energy secondary structure, which is predicted with minimizing the free energy of a conformation according to a thermodynamic mode, is functionally important [42]. The lower the value of minimum free energy is, the more energy unfolding RNA requires, so that RNA can be unfolded to be bound by some factors (including miRNA) to trigger degradation processes. Hence, minimum free energy can partly reflect lincRNAs' stability. Previous studies showed that protein-coding RNAs are more stable than lncRNAs (including lincRNAs) [43], which agree with our result that the secondary structures of a large proportion of protein-coding RNAs are more stable than lincRNAs by the analysis of minimum free energy.

The large variation in lncRNA stability is associated with their functional diversity [4,25,44]. Furthermore, lncRNAs can cluster into the same decay profiles which may be regulated by the same post-transcriptional regulatory pathways and/or contain similar regulatory sequences [21]. Previous studies propose that lncRNAs with more structural elements are more stable [21]. Through the analysis of Gene Ontology (GO) terms with differences in mRNA half-life distribution, Tani et al. found that short-lived mRNAs \( (t_{1/2} < 4 \text{ h}) \) were significantly enriched in implicated regulatory functions, and long-lived mRNAs \( (t_{1/2} \geq 4 \text{ h}) \) were disproportionately represented implicated housekeeping functions [19]. Furthermore, some lncRNAs with short half-lives \( (t_{1/2} < 4 \text{ h}) \) were well-known regulatory lincRNAs, and several ncRNAs with long half-lives \( (t_{1/2} \geq 4 \text{ h}) \) were involved in housekeeping functions [19], as is the case with mRNAs. The median lncRNA (including lincRNA) half-life is 3.5 h (mean 4.8 h) [21], so unstable lincRNA is determined with half-life \( < 4 \text{ h} \), and stable lincRNA is determined with half-life \( \geq 4 \text{ h} \).

Although the functions of a large proportion of both overlapping and unique lincRNAs in PH and ENCODE datasets were mysterious, we suggested their functions by correlating the stability of lincRNA to the functional categories. In this study, most overlapping lincRNAs were more stable than unique lincRNAs regardless of FPKM by the analysis of differences in minimum free energy distribution and lincRNA half-lives. Moreover, there was no correlation between lincRNA (including lincRNA) expression and half-life [21], similar to the result that lincRNA expression and minimum free energy were not correlated. Highly stable mRNAs often encode highly stable proteins with “housekeeping” functions [45]. Similarly, relatively stable lincRNAs with long half-lives may serve “housekeeping” roles. Stable lincRNAs may avoid

![Image](http://www.biologydirect.com/content/9/1/15)

**Figure 5** Stabilities of overlapping lincRNAs in both datasets. LincRNA decay curves after blocking transcription in K562 with actinomycin D and measuring relative levels of lincRNA remaining relative to a control gene (GAPDH) by qPCR. The relative quantitative values at time 0 h were arbitrarily adjusted to 100%. The fitted curve was modeled using locally weighted polynomial regression (LOESS method). Results are from three biological replicates, and Error bars show standard deviation \([**] P < 0.01; (*) P < 0.05, \text{ Student's t-test}\).
Figure 7 The number of the protein-coding RNAs and lincRNAs in PH. 2914 protein-coding RNAs exist in PH, 2546 (87.4%) of which present in both datasets. 1804 lincRNAs exist in PH, 795 (44.1%) appear in both datasets.

Figure 8 Comparison of the minimum free energy between the common protein-coding RNAs and lincRNAs. Box-and-whisker plot. (Whiskers) 1st–99th percentile, without individual transcripts outside this. (Box) 25th–75th percentile. Difference calculated using a nonparametric Mann-Whitney t-test (*** P < 0.000001).
degradation for undertaking some functions through various mechanisms, such as the stable secondary structure and interactions with RNA-binding proteins. The overlapping lincRNA XIST (half-life ~ 7 h) mediates X-chromosome inactivation in cis by recruiting a chromatin-modifying complex to specific sites [46]. It is a potent suppressor of hematologic cancer in mice, which may account for that human malignancies sometimes show X chromosome aneuploidies [17]. Additionally, MALAT-1, intergenic ~7 kb single exon transcript, has a highly conserved tRNA-like sequence at the 3’ end and process to generated a short tRNA-like ncRNA mascRNA [47]. It is stable in human B cells and Hela (half-life > 7 h) and has been found to regulate alternative splicing of endogenous target genes in the light of the information of Incrnadb [48].

On the other hand, relatively unstable lincRNAs with short half-lives may be expressed in narrow time windows in response to external stimuli, which is vital for reflecting regulatory functions [19,21]. Low stabilities are characterized in many transcription factor mRNAs, and transcription factors can regulate gene expression in response to environmental signals through activating or repressing target genes [23,49]. Unstable lincRNAs such as H19 [18], HOTAIR [50] and GAS5 [51], could act almost immediately after transcription without producing a functional gene product in the nucleus, so they might not require too long half-lives. Furthermore, unstable RNAs (including lincRNAs) associating with chromatin binding proteins suggested unstable lincRNA would be suitable for regulating gene expression [10,21]. The unique lincRNA H19 (half-life ~ 1 h) is abundantly expressed during embryonic development and down-regulated after birth and can regulate processing of miR-675 [18,52]. Additionally, HOTAIR with a half-life < 4 h in human Hela cells [19], has active role in modulating the cancer epigenome by binding to the polycomb repressive complex 2 (PRC2) [50]. GAS5 with a half-life < 4 h in human Hela cells [19], is a ribo-repressor by binding DNA domain of the glucocorticoid receptor to influence cell survival and metabolic activities during starvation [51].

Conclusions
We improved lincRNAs predicting pipeline to attain high-confidence lincRNAs and showed that lincRNAs are ubiquitous transcribed in K562 cells. By comparing both RNA-Seq datasets of K562 cells to explore the stability of lincRNAs, a large proportion of overlapping lincRNAs were more stable than unique lincRNAs by the analysis of differences in minimum free energy distribution and lincRNA half-lives. These results implied that overlapping lincRNAs (relatively stable lincRNAs) and unique lincRNAs (relatively unstable lincRNAs) could be associated with different functions, which will facilitate future experimental and computational investigations about lincRNA for leukemia disease.

Methods
Cell culture
K562 cell line was cultured in RPMI 1640 medium (GIBCO, Life Technologies) supplemented with 10% (v/v) fetal bovine serum, 100 units/ml Penicillin, and 100 μg/ml Streptomycin (P/S) (GIBCO, Life Technologies) in incubator (5% CO₂, at 37°C).

RNA extraction, illumina library construction and sequencing
Total RNA from K562 cells was prepared using Trizol reagent (Invitrogen.USA). Subsequently, they were used for mRNA purification and library construction with the Truseq™ RNA Sample Preparation Kit v2 (Illumina, San Diego, CA, USA) following the manufacturer’s instructions. Our sample was named PH that was sequenced on an Illumina HiSeq™ 2000 (Illumina) with pair-end libraries in Encode Genomics Bio-Technology Co. (Suzhou, China).

Preprocessing RNA-Seq datasets
Transcriptome reconstruction of PH and ENCODE datasets of K562 cells by RNA-Seq was performed respectively using rigorous read set through a sliding window filtering the average quality values within the window less than 20 and the length of reads less than 35 bp by Trimmomatic [53]. After quality control, we obtained 90.7 million 2*100-base paired-end reads generated by Illumina Hiseq2000 sequencing on polyadenylated selected (Poly-A+) RNAs. On the other hand, ENCODE RNA-Seq dataset was incorporated 112.3 million 2*76-base paired-end reads generated by Illumina HiSeq2000 sequencing on polyadenylated selected (Poly-A+) RNAs from NCBI Gene Expression Omnibus (GEO) database with accession number GSM765405.

Obtaining annotated and putative novel lincRNAs
Quality-control reads were aligned by TopHat (v2.0.7) [54], and transcripts of PH and ENCODE datasets were reconstructed by Cufflinks (v2.0.2) [55] with the Ensembl annotation, respectively. Because of strand-specific of ENCODE by RSeQC script [56], fr-firststrand library type was performed for ENCODE by Cufflinks. To eliminate all annotated non-lincRNA transcripts, the intersection of transcripts of the ‘u’ category (unknown, intergenic transcript in Cufflinks) was attained using cuffcompare script with four public databases annotations (annotated protein-coding genes, microRNAs, rRNAs, tRNAs and pseudo-genes), including Ensembl (Homo_sapiens.GRCh37.70.gtf), UCSC (hg19), Gencode (gencode.v15.annotation.gtf.gz) and Refseq (ref_GRCh37.p10_top_level.gtf.gz) respectively. That is, the intersection of intergenic transcripts was acquired apart from all annotated non-lincRNA annotations of four
databases. Annotated lincRNAs were acquired through the intersection of intergenic transcripts to run Cuffcompare script again with Gencode lncRNAs annotation (gencode.v18.long_noncoding_RNAs.gtf.gz). The remaining transcripts were possible novel lincRNAs. Then, possible novel lincRNAs were filtered on the basis of some characteristics including FPKM value (Due to the low expression of lincRNAs, we considered to acquire more lincRNAs based on the density distribution of lincRNAs. ENCODE, FPKM ≥ 0.1; PH, FPKM ≥ 0.01), length (≥ 200 nt), ORF (< 100 codons) and exonic number (≥ 2). After that, putative novel lincRNAs were acquired based on non-coding potential by integrating the results of four softwares including iseeRNA (noncoding), CPAT (no), CPC (noncoding) and PhyloCSF (score < 100) (Figure 9).

iseeRNA, a lightweight SVM-based program, is designed for computational identification of lincRNAs from high-throughput transcriptome sequencing data. CPAT, which overcomes several intrinsic pairwise and multiple alignments limitations, uses logistic regression model based on ORF size, ORF coverage, Fickett TESTCODE and Hexamer bias. CPC relies on pairwise alignment to assess the protein-coding potential of a transcript based on six biologically meaningful sequence features. PhyloCSF uses a multi-species nucleotide sequence alignment to calculate the phylogenetic conservation score, which is likely to represent a protein-coding region.

Minimum free energy

Minimum free energy secondary structure is assessed by minimum free energy (thermodynamic free energy). RNAfold predicts minimum free energy and minimum energy secondary structures using a loop-based energy model and dynamic programming algorithm [57,58].

Half-life verification using quantitative real-time PCR (qRT-PCR)

For experiment of half-life, cells were grown to ~50% confluency before RNA polymerase activity was blocked by 10μg ml⁻¹ actinomycin D (sigma) in DMSO. Transcription inhibition of K562 was carried out for 30 hours, and cells were harvested at time 0 h, 0.5 h, 2 h, 4 h, 8 h, 24 h and 30 h, respectively.

Total RNA samples at different time were prepared using Trizol reagent (Invitrogen, Carlsbad, CA, USA). One microgram of RNA was reverse transcribed into cDNA using Reverse Transcriptase M-MLV (TAKARA, Japan), and diluted by 1:5 for RT-PCR assay. The assay was performed by using SYBR Premix Ex Taq (TAKARA, Japan) on ABI 7500 Real-Time PCR System. GAPDH was used as an internal control. 2^ΔΔCt values were calculated for each gene to show the fold change. All qRT-PCR reactions were performed in three biological replicates.

We determined the lincRNA half-life by calculating the time when the fold change of expression abundance reached half of the initial expression abundance (0 h time point). Decay profile was modeled using locally weighted polynomial regression (LOESS method) (www.r-project.org).

Reviewers’ comments

We appreciate the reviewer’s comments from Prof Oliviero Carugo, Dr Alistair Forrest and Prof Manju Bansal. We have revised the manuscript accordingly.

Reviewer #1 (First Round): Prof Oliviero Carugo, University of Vienna, Austria

Major comments

The main assumption of this manuscript is that overlapping lincRNAs (those that are found both in the ENCODE dataset and in the PH dataset) have functions different from unique lincRNAs (those that are found only in one dataset). In fact, on the basis of the data shown in the manuscript, this is an epistemological axiom (if not a postulate) that might be questioned. The experimental observation that five (5) unique lincRNAs are less stable than seven (7) overlapping lincRNAs is in fact insufficient to conclude that there is a functional difference.

Authors’ response: We have revised the whole paper to demonstrate our ideas. We showed that a large proportion of overlapping lincRNAs were more stable than the unique lincRNAs through the analysis of differences in minimum
free energy distribution. Furthermore, the experiments of lincRNA half-lives also supply the result. Through not only the half-lives of overlapping and unique lincRNAs but also the well-studied lincRNA with known half-life and previous studies about RNA stability, we suggested that overlapping lincRNAs (relatively stable lincRNAs) and unique lincRNAs (relatively unstable lincRNAs) could be associated with different functions.

As a consequence, the entire manuscript is highly speculative and this must be clearly pointed out in the discussion.

Authors’ response: We have added some evidences to supply our ideas and moderated some speculations which were insufficient evidences in the discussion.

In the chapter entitled “Obtaining annotated and putative novel lincRNAs”, in the “Methods” section, a more detailed description must be provided to allow one to reproduce the algorithm. If this is too long, a supplementary file may be submitted (with some examples).

Authors’ response: We have re-written the lincRNA analysis workflow in the “Methods” section.

Also the introductory section should be modified. The first part of the “Background” section is a good introduction about the stability of non-coding RNAs and its functional implications. Some additional emphasis should be put on the relationships with cancer and diseases. Otherwise, the mention to medical issues is inappropriate. The last part of the “Background” section, which is a short summary of the experiments and of the results of the manuscript, should be slightly expanded to be more easily understood by the reader. In its present form, it is too short and it becomes, inevitably cryptic.

Authors’ response: Thanks for the comments of reviewer. We have re-written the introduction and revised the whole article.

I cannot inspect Figure 8 (resolution is too small).

Authors’ response: We have adjusted it in the revised version.

Minor comments

Section “Background” line 43 – I am expecting that the Authors describe briefly which cancers and which diseases are related to lincRNAs. The first sentence in this section is otherwise too vague for a scientific publication.

Authors’ response: We have revised the introduction and deleted the first sentence.

Section “Background” line 46 – Please, check if the expression “... that are proved the importance ...” is correct. Perhaps it should be “... that are proven to be important ...

Authors’ response: We have revised the introduction and deleted the sentence.

Section “Background” line 62 – Please, check if the expression “... Clark et al found only a minority ...” is correct. Perhaps it should be “…Clark et al found that only a minority ...

Authors’ response: Thanks, we have corrected it in the revised version.

Section “Background” line 68 – Please, check if the expression “… we improve lincRNA ...” is correct. Perhaps it should be “… we improve the lincRNA ...

Authors’ response: We have corrected it in the revised version.

Section “Background” lines 68-71 – These two sentences, which are a sort of summary of the experiments described in this manuscript, are a bit confusing. First, the Authors write that they compared “different RNA-Seq datasets. Then, they write that they compared “both RNA-Seq datasets”. The question is: are they “several” or just “two”? Moreover, the Authors should describe briefly which are these datasets, how they were assembled, validated, compared, etc. Just few sentences should be enough to improve the readability of the manuscript.

Authors’ response: We revised introduction and described the improved lincRNAs workflow in the “Methods” section.

Section “Background” lines 72-73 – The expression “through randomly testifying ... unique lincRNAs” is not clear. The Authors should re-write it.

Authors’ response: The sentence has been re-written according to the reviewer’s suggestion.

Section “Background” line 74 – The expression “coinciding” might be “in agreement with”.

Authors’ response: We have deleted the sentence in the revised manuscript.

Section “Background” line 70 – The Authors should justify why they selected K562 cells.

Authors’ response: It has been revised according to the reviewer’s suggestion.

Section “Background” lines 76-77 – The sentence “Therefore, we suggest ... lincRNA stability” is unclear and should be re-written.

Authors’ response: We rewrote it in the revised manuscript.

Section “Results and discussion” line 85 – A reference to Cufflinks is mandatory. And it is also necessary to mention what is that (briefly).

Authors’ response: We have added the cited paper in the revised manuscript.

Section “Results and discussion” line 86 – Probably it would be better to write simply “by considering the annotations present in XXX” or “by using the software XXX” and “(see Methods for details)”. In its present form and without references, this list of resources is not really readable.

Authors’ response: We added the cited paper in the revised manuscript.

Section “Results and discussion” lines 90-91 – The Authors should justify why their method is more effective...
than alternative methods based on the integration of several databases. In fact, I understand that it is simpler. But I am not sure it is more effective.

Authors’ response: We have revised it.

Section “Results and discussion” lines 103-104 – The Authors mention “four public database annotations”. They should also indicate their names and where they can be found.

Authors’ response: It has been revised according to the reviewer’s suggestion.

Section “Results and discussion” line 104 – The Authors mention “PH” here for the first time: it is necessary to define it explicitly. If I am not wrong, it is defined only in the Abstract and this is not sufficient.

Authors’ response: PH was defined in the “Methods” section.

Section “Results and discussion” lines 109-113 – The Authors describes the minimum free energy”. This should be described better and it is necessary to cite the computational procedure used to compute this thermodynamic quantity.

Authors’ response: We have revised it in the “Methods” section according to the reviewer’s suggestion.

Section “Results and discussion” lines 116-120 – These sentences repeat the “Background” section and should be removed (or moved to the introductory section).

Authors’ response: These sentences been deleted according to the reviewer’s suggestion in the revised manuscript.

Section “Results and discussion” lines 126-127 – The expression “Comparing the both datasets, …” might be inappropriate and might be even removed.

Authors’ response: It has been revised according to the reviewer’s suggestion.

Section “Results and discussion” line 128 – RNAfold can be used to compute something and it cannot be calculated.

Authors’ response: We have deleted the sentence in the revised manuscript.

Section “Results and discussion” line 132 – The verb “testified” might be inappropriate. Why not “determined experimentally”?

Authors’ response: It has been revised according to the reviewer’s suggestion.

Section “Results and discussion” line 134 – The “5 unique lincRNA” were taken from PH or from ENCODE?

Authors’ response: It has been revised according to the reviewer’s suggestion.

Section “Results and discussion” line 136 – A reference to “qPCR after ActD treatment” is necessary.

Authors’ response: We have added the cited paper in the revised manuscript.

Section “Results and discussion” lines 137-138 – A expression “… which is coordinated …” might be inappropriate. Perhaps it might be changed into “… in agreement with …”.

Authors’ response: We have deleted the sentence in the revised manuscript.

Section “Results and discussion” line 143 – The verb “express” might be “may be expressed”. Section “Results and discussion” line 151 – The verb “is” might be inserted between “It” and “highly”.

Authors’ response: It has been revised according to the reviewer’s suggestion.

Section “Results and discussion” lines 150-157 – Caution: this discussion is highly speculative and it should be re-written. It is necessary to clearly indicate that these are mere suppositions of the Authors that are not based on any new results.

Authors’ response: We have re-written in the revised manuscript.

Section “Results and discussion” line 164 – The expression “FPKM” should be indicated extensively.

Authors’ response: We have re-written the sentence in the revised manuscript.

Section “Methods” line 188 – The verb “were” might be “was”.

Authors’ response: Thanks, we corrected it in the revised version.

Section “Methods” line 196 – The sentence “Total RNA was extracted as described above” might be deleted.

Authors’ response: We have deleted the sentence in the revised manuscript.

Section “Methods” line 231 – “We” might be “we”.

Authors’ response: We are sorry about this mistake, and have corrected it in the revised version.

Reference number 8. The journal name should be abbreviated. The same goes for several other references.

Authors’ response: We fully checked the manuscript and revised similar question.

Reference number 34. Volume and pages are missing. The same happens also in some other reference.

Authors’ response: We fully checked the manuscript and revised similar question.

In the right part of Figure 1, the box named “Non-coding” might be modified by listing the four programs in the same order used to describe them in lines 93-100 of the section “Results and discussion”: iseeRNA, CPC, PhyloCSF, CPAT. In the present figure, CPAT is the second program and not the fourth.

Authors’ response: We have modified the description about Figure 9 in the “Methods” section.

The caption of Figure 2 is unclear. It should be re-written.

Authors’ response: We have revised it.

Figure 3 – Which units are used the measure the free energy? The data shown in the figure are taken from PH or Encode? It is also necessary to write what are the
thick horizontal black lines and what the error bars indicate. In fact, although most of the readers will understand this figure, it is mandatory to write an exhaustive legend.

Authors’ response: It has been revised according to the reviewer’s suggestion.

In Figure 4, “Encode” should be “ENCODE”.

Authors’ response: We are sorry about this mistake, and have corrected it in the revised version.

Figure 5 – See the observations about Figure 3.

Authors’ response: It has been revised according to the reviewer’s suggestion.

Figures 6 and 7 – These curves have very different shapes. Might the Authors try to describe them and write something about the differences?

Authors’ response: It has been revised according to the reviewer’s suggestion.

Reviewer #1 (Second Round): Prof Oliviero Carugo, University of Vienna, Austria

The revised version of the manuscript is considerably better than the original version. However, in my opinion, there is still a very modest direct and experimental evidence of the difference between overlapping and unique lincRNAs. As a consequence, the conclusions are extremely speculative.

Authors’ response: To explore the stability of two separated classes of lincRNAs by comparing both RNA-Seq datasets of K562 cells, we have analyzed minimum free energy distribution and measured the half-lives of the selected lincRNAs. The results supported the conclusion that the overlapping lincRNAs show more stable than the unique lincRNAs in K562 cell. Actually, the conclusion could be accepted in a general consideration that the overlapping lincRNAs expressed in both datasets should have more probability to be detected than the unique lincRNAs only expressed in one single dataset. Therefore, the lincRNAs existing in two datasets were likely more stable than those existing in one single dataset.

Reviewer #2 (First Round): Dr Alistair Forrest, Omics Science Center, Japan

Major comments

1. "The method is simpler and more effective than integrating several database annotations by their own scripts [29,30]." Unless you provide benchmarking this is an unsupported statement. Should be removed.

Authors’ response: We did not compare the effectiveness without their scripts, however, our pipeline is simpler and more convenient than integrating several database annotations by the scripts.

2. "our sequenced RNA-Seq dataset (PH)." Why is the dataset called PH?

Authors’ response: We are really grateful to the reviewer’s carefulness. A series of RNA-Seq datasets were sequenced in K562 cells by PMA or hemin treatment. PH was untreated RNA-Seq dataset.

3. The main point of the paper seems to be that reproducible lincRNAs have higher free energy and inferred secondary structure than lincRNAs only observed in one dataset. I think this is likely to just reflect abundance. More highly expressed lincRNAs are more likely to be observed in multiple datasets. Weakly expressed lincRNAs are less likely. The authors could perhaps strengthen their story by looking at the relationship between expression level and stability by breaking the lincRNAs into several bins (low, mid, high) and examine the free energy (and half-life) in box plots.

Authors’ response: Thanks for the reviewer’s helpful suggestions. We have taken this advice and analyzed the relation of the minimum free energy and expression level in the revised manuscript. However, they were not correlated, similar to no correlation between lincRNA (including lincRNA) expression and half-life.

4. "highly expressed in ENCODE, but hardly expressed in PH." For these kind of statements it is very important to explain what RNA-seq protocol was used for ENCODE and PH. If the methods do not match this may explain why you see differences.

Authors’ response: We appreciate the reviewer’s suggestion. ENCODE and PH datasets were sequenced following the manufacturer’s instructions of Illumina. And we analyzed them using the same method (see Methods).

5. "Unstable lincRNAs are very sensitive, respond rapidly when transcription changes and act almost immediately after transcription without producing a functional gene product in the nucleus." Unsupported statement. You do not demonstrate that unstable lincRNAs are inherently ‘rapid responders’, neither do you demonstrate ‘without producing a functional gene product’. The manuscript should be re-read critically examining whether your statements are supported by your analysis or from a primary reference.

Authors’ response: We added the cited paper and revised the sentence.

6. "Pervasive transcription of lincRNAs in K562 cells". This section doesn’t really add anything. "514 lincRNAs (FPKM > =1) in ENCODE (80 for FPKM > =10), whilst there were 312 lincRNAs (FPKM > =1) in PH (30 for FPKM > =10). 89 overlapping lincRNAs of both datasets expressed with FPKM > =1." This does in no way suggest pervasive.

Authors’ response: We really thank for the reviewer’s question. We have re-written the section. The distribution and expression of lincRNA were showed in the revised Figure 1.

Minor comments

1. "LncRNAs, located and transcribed from intergenic genomic regions that are proved the importance in cancer
by genome-wide studies, are named intergenic lncRNAs (lincRNAs)." Do you mean the lncRNAs or the genomic regions where they are found are associated with cancer? The references 5, 6 do not directly correspond to a cancer link.

Authors' response: We have re-wrote the introduction and deleted the sentence.

2. "lincRNA" and 'lncRNA' are used interchangeably in the paper. lincRNA corresponds to 'intergenic', whereas lncRNA are just long. You should make a definition and stick to one, probably lncRNA.

Authors' response: LncRNA is a subgroup of lncRNA. The features of IncRNA also are showed for lincRNA. Some cited papers were associated with lncRNA, however, IncRNA included lincRNA.

3. "It is found that a large proportion of overlapping lincRNAs are more stable than unique lincRNAs". I suggest the authors not use the term 'overlapping lincRNAs' as this suggests genomic overlap. I think what the authors mean is 'lncRNAs observed in multiple K562 datasets are more stable than lncRNAs unique to one K562 dataset'.

Authors' response: We have annotated overlapping lincRNAs and unique lincRNAs in the "Results" section. We compared PH and ENCODE datasets to attain overlapping and unique lincRNAs in venn diagram (Figure 2). Unique linRNAs presented in only PH or ENCODE dataset, and overlapping lincRNAs presented in both ENCODE and PH datasets.

4. "In light of this, we acquired a great deal of intergenic transcripts involved possible novel lincRNAs and annotated lincRNAs with Ensembl or Gencode". Unclear what "transcripts involved possible novel" means.

Authors' response: It has been revised according to the reviewer's suggestion.

5. "Hexamer usage bias" Bias

Authors' response: Thanks, we have corrected it in the revised version.

6. "1,804 lincRNAs were indentified" Identified

Authors' response: We are sorry about this mistake, and have corrected it in the revised version.

7. "We randomly testified the half-lives of 7" tested

Authors' response: We have revised it.

Reviewer #2 (Second Round): Dr Alistair Forrest, Omics Science Center, Japan

Major points

1. OK. The authors have removed the sentence, still the section is entitled 132

   "Improved pipeline for lincRNA analysis". I would change to just "Pipeline for lincRNA analysis" as still you have provided no evidence that the pipeline is 'improved'.

Authors' response: It has been revised according to the reviewer's suggestion.

2. FIX. The authors must add a sentence at the very first use of PH explaining that PH is (PMA or Hemin treated K562s).

Authors' response: We have added the annotation at the very first use of PH in the revised manuscript. In this study, PH was only named for our RNA-Seq dataset of untreated K562 cells.

3. FIX. Figure 4 should not be split into A and B on 1 FPKM. Only one set of boxplots should be shown.

Authors' response: Figure 4 splits into A and B on 1 FPKM in order to correspond to the minimum free energy (A and B in Figure 3). It can clearly illuminate the cases of FPKM $\geq 1$ and FPKM < 1.

4. OK. Actually GSM765405 was sequenced using the CSCL RNA-seq protocol not Truseq. However should be comparable.

Authors' response: In this study, we have compared the protein-coding RNAs from both datasets by different sequencing library construction methods. We found that 2546 (87.4%) protein-coding RNAs in PH also presented in ENCODE, which showed that the results from CSCL RNA-seq protocol and Truseq are comparable.

5. OK

6. OK

New minor comments

1. line 139 PhloCSF [34], PhyloCSF

Authors' response: We have corrected the spelling mistake in the revised version.

2. expressive abundance (FPKM). expression level or transcript abundance

Authors' response: It has been revised according to the reviewer's suggestion.

Reviewer #3 (First Round): Prof Manju Bansal, Indian Institute of Science, India

The authors evaluate the stability of long intergenic non-coding RNA (lincRNA) in human K562 cell using the RNA-seq data. Two datasets of lincRNA are used in the study. The authors developed a pipeline to enrich lincRNA compared to that seen in ENCODE dataset. The stability of the lincRNA in these two dataset is compared, using mfe predicted by secondary structure program RNAfold. The reason to carry out this comparison is not clearly explained. It is no surprise that non-coding regions have lower stability than protein-coding regions.

Authors' response: The secondary structure of lincRNA is important for its stability. We estimated the stabilities of overlapping and unique lincRNAs by bioinformatic method (minimum free energy). Furthermore, through the analysis of minimum free energy, our result agreed with previous report that lincRNA have lower stability than protein-coding RNA. That is, it was verification that the stability of lincRNA could be assessed using minimum free energy.
Further, the dataset is divided into overlapping and unique. Stability study shows that overlapping lincRNAs are more stable than unique lincRNAs. The explanation for the observed difference is not clearly stated. I cannot understand why lincRNA identified in two datasets (overlapping) should be more stable.

Authors' response: We showed that a large proportion of overlapping lincRNAs were more stable than the unique lincRNAs through the analysis of differences in minimum free energy distribution and lincRNA half-lives.

RNA half-life studies are carried out on few lincRNAs under each category. The findings of the RNA half-life studies, using very few samples (5-7), cannot be generalized for > 700 odd lincRNAs in each group. The conclusion that lincRNA stability may be related to function is already shown in Ref [26,28]. Overall, the conclusions made by the authors are rather weak or already established.

Authors' response: Since lots of experiments may need too much time, we have added some experiments of lincRNA half-lives. We have revised the conclusions and suggested that overlapping lincRNAs (relatively stable lincRNAs) and unique lincRNAs (relatively unstable lincRNAs) have different functions.

The manuscript has several grammatical mistakes, starting from the very first sentence in the Introduction section. Many paragraphs have loose statements and irrelevant information. E.g. See line 116 to 120. LincRNA and lncRNA are used interchangeably in many places.

Authors' response: We really appreciate the reviewer’s advice. We have revised the whole paper (including introduction) and deleted some irrelevant information.

The methods section dealing with RNAfold need to be elaborated. Figures need to have proper units and labelling. E.g. MFE units.

Authors' response: It has been revised according to the reviewer’s suggestion.

Reviewer #3 (Second Round): Prof Manju Bansal, Indian Institute of Science, India

I had requested the author to explain the purpose of dividing the dataset into Overlapping and Unique. See earlier comment “Further, the dataset is divided into overlapping and unique. Stability study shows that overlapping lincRNAs are more stable than unique lincRNAs. The explanation for the observed difference is not clearly stated. I cannot understand why lincRNAs identified in two datasets (overlapping) should be more stable”. The author’s explanations are not satisfactory and have merely reiterated the aim of the work.

Authors’ response: The lincRNAs are considered to play very important roles in gene regulation which shows dynamic properties in cellular processes. Therefore, identification of lincRNA stability is important to annotate its functions. In this paper, we classified the lincRNAs into overlapping lincRNAs and unique lincRNAs by comparing two RNA-Seq datasets of the same K562 cell line with venn figure (Figure 2). In general, it is easy to understand that the overlapping lincRNAs are more stable than the unique lincRNAs, because the expressing probability of lincRNA should have high value if it could be observed in two separated experiments from ENCODE and our own detection in K562 cells. That it, lincRNAs expressed in both experiments statistically have higher expressing probability than lincRNAs only expressed in one single experiment. We have carried out RNA-Seq of K562 cell line, compared our RNA-Seq dataset with the corresponding ENCODE dataset, and found that a large proportion of protein-coding RNAs (86.7%) of our dataset appeared in ENCODE, but relative small proportion of lincRNAs (44.1%) of our dataset appeared in ENCODE (Figure 7). We speculated this phenomenon arises due to the instability of lincRNAs during the cellular processes. We classified the overlapping part (overlapping lincRNAs) and the unique part (unique lincRNAs) from two datasets, and compared their stabilities. We have proved that a large proportion of overlapping lincRNAs were more stable than the unique lincRNAs through the analysis of differences in minimum free energy distribution and lincRNA half-lives.

The authors postulate that the unique lincRNAs present in ENCODE or PH dataset should be functionally different. This implies that the RNAs that are identified and annotated as lincRNAs by both ENCODE and PH pipeline have similar function and this not true.

Authors' response: We suggested that overlapping lincRNAs (relatively stable lincRNAs) and unique lincRNAs (relatively unstable lincRNAs) can be related to different functions, because they might be related to different cellular processes.

Further, minimum free energy of RNA is dependent on the GC content. A comment on the GC content of the overlapping and unique lincRNAs can explain the observed difference between the groups (Figure 3). Similar observation is warranted for half-life based stability analysis. If the GC content of the randomly selected lincRNAs in the two groups (overlapping (10 sequence) and unique (7 sequence)) is different then the selection of lincRNAs is biased by the sequence property. Moreover, mRNA expression is known to be affected by GC content. (G Kudla - & 8206;2006, PLoS Biology.). Authors can check this phenomenon by binning the lincRNAs based on GC content and correlated with the level of expression.

Authors’ response: In the previous studies, there is no correlation between lncRNA expression and its stability, although the signification correlation has been found for all RNAs (e.g. Clark et al. Genome Research, 2012, 22:885-898).

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Authors’ response: The lincRNAs are considered to play very important roles in gene regulation which shows dynamic properties in cellular processes. Therefore, identification of lincRNA stability is important to annotate its functions. In this paper, we classified the lincRNAs into overlapping lincRNAs and unique lincRNAs by comparing two RNA-Seq datasets of the same K562 cell line with venn figure (Figure 2). In general, it is easy to understand that the overlapping lincRNAs are more stable than the unique lincRNAs, because the expressing probability of lincRNA should have high value if it could be observed in two separated experiments from ENCODE and our own detection in K562 cells. That it, lincRNAs expressed in both experiments statistically have higher expressing probability than lincRNAs only expressed in one single experiment. We have carried out RNA-Seq of K562 cell line, compared our RNA-Seq dataset with the corresponding ENCODE dataset, and found that a large proportion of protein-coding RNAs (86.7%) of our dataset appeared in ENCODE, but relative small proportion of lincRNAs (44.1%) of our dataset appeared in ENCODE (Figure 7). We speculated this phenomenon arises due to the instability of lincRNAs during the cellular processes. We classified the overlapping part (overlapping lincRNAs) and the unique part (unique lincRNAs) from two datasets, and compared their stabilities. We have proved that a large proportion of overlapping lincRNAs were more stable than the unique lincRNAs through the analysis of differences in minimum free energy distribution and lincRNA half-lives.

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Authors’ response: In the previous studies, there is no correlation between lncRNA expression and its stability, although the signification correlation has been found for all RNAs (e.g. Clark et al. Genome Research, 2012, 22:885-898).
Additional files

Additional file 1: Table S1. The bed file of the unique lincRNAs in PH.

Additional file 2: Table S2. The bed file of the unique lincRNAs in ENCODE.

Additional file 3: Table S3. The bed file of the overlapping lincRNAs of both datasets.

Additional file 4: Table S4. The result for running Cuffcompare script to compare lincRNAs of both datasets.

Abbreviations
lincRNAs: Long intergenic non-coding RNAs; RNA-Seq: RNA Sequencing; lincRNAs: Long non-coding RNAs; ENCODE: The Encyclopedia of DNA Elements; TUCP: The transcripts of uncertain coding potential; FPKM: Fragments Per Kilobase of exon model per Million mapped fragments; ORF: Open reading frame; NMD: Nonsense mediated decay; qPCR: Quantitative polymerase chain reaction; ActD: Actinomycin D; miRNA: microRNAs.

Competing interests
The authors declare that they have no competing interests.

Authors’ contributions
2L conceived this study. LW and DZ designed the experimental plan, drafted and revised the manuscript. DZ carried out the field experiment. JT and YW participated in preparing, treating and collecting samples. LW analyzed and revised the manuscript. DZ carried out the field experiment. JT and YW participated in preparing, treating and collecting samples. LW analyzed and revised the manuscript. DZ carried out the field experiment. JT and YW participated in preparing, treating and collecting samples. LW analyzed and revised the manuscript. DZ carried out the field experiment. JT and YW participated in preparing, treating and collecting samples. LW analyzed and revised the manuscript. DZ carried out the field experiment.

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