Long non-coding RNAs have age-dependent diurnal expression that coincides with age-related changes in genome-wide facultative heterochromatin

Jinhee Park and William J. Belden *

Abstract

Background: Disrupted diurnal rhythms cause accelerated aging and an increased incidence in age-related disease and morbidity. The circadian clock governs cell physiology and metabolism by controlling transcription and chromatin. The goal of this study is to further understand the mechanism of age-related changes to circadian chromatin with a focus on facultative heterochromatin and diurnal non-coding RNAs.

Results: We performed a combined RNA-seq and ChIP-seq at two diurnal time-points for three different age groups to examine the connection between age-related changes to circadian transcription and heterochromatin in neuronal tissue. Our analysis focused on uncovering the relationships between long non-coding RNA (lncRNA) and age-related changes to histone H3 lysine 9 tri-methylation (H3K9me3), in part because the Period (Per) complex can direct facultative heterochromatin and models of aging suggest age-related changes to heterochromatin and DNA methylation. Our results reveal that lncRNAs and circadian output change dramatically with age, but the core clock genes remain rhythmic. Age-related changes in clock-controlled gene (ccg) expression indicate there are age-dependent circadian output that change from anabolic to catabolic processes during aging. In addition, there are diurnal and age-related changes in H3K9me3 that coincide with changes in transcription.

Conclusions: The data suggest a model where some age-related changes in diurnal expression are partially attributed to age-related alterations to rhythmic facultative heterochromatin. The changes in heterochromatin are potentially mediated by changes in diurnal lncRNA creating an interlocked circadian-chromatin regulatory network that undergoes age-dependent metamorphosis.

Keywords: Circadian rhythm, Aging, Long non-coding RNA, Diurnal, Heterochromatin, Histone H3 lysine 9 methylation

Background

Early circadian research showed non-circadian day lengths negatively impact longevity [1], even though the free-running rhythm only changes slightly with age [2]. More recent reports indicate circadian disruption contributes to increased morbidity [3] and diseases including: metabolic disorders [4–6], depressive disorders [7], cardiovascular disease [8], cancer [9], and advanced aging [10]. In addition, age-related changes to the circadian clock result in a dampened endocrine and neuroendocrine hormone rhythm [11] concurrent with a decrease in Clock and Bmal1 expression, while the rhythm in Per genes is largely unaffected [12]. There are also extensive age-related changes to clock-controlled genes (ccg) in the liver that correlate with age-related changes to metabolism which occurs with a change in acetylation [13, 14]. As a result of these reports and others, there is a growing appreciation of the connection among circadian disruption, advanced-aging and an increased incidence of age-related maladies, yet the mechanism underlying these remains cryptic.
The core circadian rhythm is maintained by a transcriptional negative feedback loop where the dimeric transcriptional activators CLOCK and BMAL1 [15–18] drive expression of the Period (Per1, Per2, Per3) [19–21] and Cryptochrome (Cry1, Cry2) genes [22, 23]. Once expressed, the PERs associate with the CRYs and form a multisubunit complex (approximately 2 MDa) that contains repressive chromatin modifiers including a histone H3 lysine 9 methyltransferase (KMT1/Suv39h), RNA binding proteins, and the nucleosome remodeling and deacetylase (NuRD) corepressor [24–27]. In a second feedback loop, RORα and REV-ERBa have a positive and negative role, respectively, in regulating rhythms in Bmal1 expression [28, 29]. The clock regulates, either directly or indirectly, a large percentage of protein-coding genes [30–36] and controls genome-wide chromatin states that oscillate between permissive and non-permissive states [31, 32, 37]. Facultative heterochromatin at the central clock gene(s) in Neurospora, Drosophila and mammals is observed during the repressive phase and is characterized by histone H3 lysine 9 di- and tri-methylation (H3K9me2 & H3K9me3) and HP1 binding [24, 38–42]. The extent of circadian-regulated facultative heterochromatin (CRFH) throughout the genome is unknown, but it does occur at D-element binding protein (Dbp) suggesting, at the very least, it is present at a ceg [41].

Long non-coding RNAs (lncRNAs) and natural antisense transcripts (NATs) are known to aid in establishing repressive chromatin and can function in both cis and trans [43, 44]. As such, they are a potential mechanism by which the circadian clock could establish permissive or non-permissive chromatin states to regulate gene expression. Support for this idea comes from numerous reports documenting a wide-array of rhythmic ncRNA [32] including a NAT that originates from Per2 [31, 32]. Based on studies on the Neurospora frequency NAT, it is possible that Per2AS may be necessary to help establish heterochromatin and assist in feedback repression [39].

Current theories on aging suggest that changes in heterochromatin may be responsible for age-related changes in gene expression and alterations in H3K9me3 may function as a driver of aging [45, 46]. For example, in models of Werner syndrome, a premature aging disorder in humans, there are genome-wide changes to H3K9me3. The changes in H3K9me3 in Werner syndrome are likely due to WRN-guided heterochromatin because WRN associates with KMT1/SUV39H and HP1 [46, 47]. The connections among the circadian clock, heterochromatin, lncRNA and aging led us to examine the role of lncRNA in age-related changes to the clock-chromatin circuit and ascertain whether this may contribute to age-related diseases. To accomplish this, we performed a comprehensive RNA-sequencing (RNA-seq) and H3K9me3 ChIP-seq on zebrafish brain tissue from animals that were 4, 12, and 20 months (M) old at zeitgeber time (ZT) 4 and ZT16. We observed age-related changes to diurnal H3K9me3 and RNA expression, with a subset of diurnal genes showing changes in H3K9me3. The results suggest that age-related changes to circadian regulated transcription occur in part due to changes in heterochromatin.

Results

Identification of long non-coding RNAs

There are age-related changes to circadian transcription that coincide with changes in physiology, but the connection to changes in circadian noncoding RNA and chromatin are lacking. To further define age-related changes to circadian transcription, including changes in lncRNA expression, we performed a multi-dimensional RNA-sequencing (RNA-seq) experiment with a focus on lncRNA transcript discovery. We isolated total RNA from zebrafish brain tissue at two diurnal time points (ZT4 and ZT16) for 3 different ages (4 M, 12 M, 20 M). We processed the ribo-depleted, stranded RNA-seq data using TopHat2/Cufflinks (T/C) [48] and/or HISAT2/StringTie (H/S) [49] with the zebrafish genome GRCz10. In our analyses, we also included stranded RNA-seq data from developing zebrafish embryos [50] to enhance lncRNA identification. In order to detect the known per2AS transcript with H/S pipeline, we had to use an FPKM cut-off of 0.1 in StringTie. Stipulating per2AS identification as criteria in StringTie generated an excessive number of single-exon transcripts (> 150,000), so we proceeded with the analysis using T/C results (Additional files 1 and 2). The Cufflinks transcripts were further subdivided by class code and processed with bedtools intersect to identify potential overlapping lncRNA. This analysis revealed there were potentially 14,773 overlapping transcripts (OT). Next, we processed the OT with PLEK (predictor of long non-coding RNAs and messenger RNAs based on an improved k-mer scheme) to predict potential NATs [51] with the notion these may play a role in chromatin regulation. Overall, this analysis revealed 5299 lncRNAs overlapped with another gene, either protein-coding gene or overlapping lncRNAs. The inclusion of RNA-seq data from developing zebrafish embryos allowed us to perform a comprehensive analysis of lncRNA from early development through aged adult animals. To further parse the data, we took the 146,910 transcripts from Cufflinks and H3K9me3 ChIP-seq on zebrafish brain tissue from animals that were 4, 12, and 20 months (M) old at zeitgeber time (ZT) 4 and ZT16. We observed age-related changes to diurnal H3K9me3 and RNA expression, with a subset of diurnal genes showing changes in H3K9me3. The results suggest that age-related changes to circadian regulated transcription occur in part due to changes in heterochromatin.
separate regulated transcription versus nonspecific transcriptional noise. In total, 17,702 potential lncRNAs displayed differential expression from embryogenesis to aged adult. Further clustering of 17,702 transcripts into 8 groups based on expression revealed that expression of some lncRNA are restricted to specific times during embryo development, while others are restricted to adult fish (Additional file 3). In general, we found specific lncRNAs are largely confined to brain tissue and were either not present during development or were low abundance. One group (Group 6, Additional file 3 g) displayed a noticeable diurnal oscillation at 12 M where transcripts were highly expressed at ZT4, but largely absent at ZT16. These same lncRNAs appeared constant regardless of the time-of-day at other ages.

**Age-related changes to diurnal gene expression**

Next, we proceeded to examine diurnal expression across all three age groups. Transcripts were split into protein-coding genes and transcripts not currently annotated in the GRCh38 reference assembly (unannotated transcripts). Among the 87,327 unannotated transcripts, PLEK predicted roughly 58% were lncRNA (Additional file 3). Analysis of diurnal expression comparing ZT4 to ZT16 (q ≤ 0.05) revealed 273, 604, and 193 diurnal transcripts in 4, 12, and 20 M age groups, respectively (Fig. 1a). Unannotated transcripts represented 48.7% and 39.4% in 4 and 20 M groups, while they represented 83.4% of diurnally expressed transcripts in the 12 M group. The annotated diurnal genes for each age group can be found in Additional files 4-6. At each age, a portion of the diurnal transcripts overlapped with another transcript on the opposite DNA strand. In total, 9.5% (26 of 273), 6.6% (40 of 604), and 12.4% (24 of 193) of diurnal transcripts at 4, 12, and 20 M contained an overlapping transcript.

To examine age-related changes in diurnal gene expression, diurnally expressed genes from the three different ages were split into two groups. Those that peaked at ZT4 were classified as morning-specific and those that peaked at ZT16 were classified as evening-specific. Comparison across ages revealed the vast majority of diurnal genes change with age and only a small subset (32 morning-specific and 7 evening-specific) of genes maintained diurnal expression regardless of age (Fig. 1b,c). The morning-specific genes included *per2* and *cry1a*, while evening-specific genes included *bmal1*/*arntl1a* and *clock* (Fig. 2c). We further examined the age-related changes to diurnal gene expression by clustering transcripts elevated at ZT4 relative to ZT16 for each age (Additional file 7). In general, morning and evening specific genes changed expression as the animals aged.

To further define the physiological consequence of age-related changes in diurnal gene expression, we calculated gene ontology (GO) enrichment as a function of age using WEB-based GEnE SeT AnaLysis (WebGestalt) Toolkit [52, 53] (Fig. 1d). This analysis showed diurnal genes differed in their ontology as a function of age. For example, in 4 M-old fish diurnally expressed genes were mainly associated with nucleosome/chromatin assembly, protein-DNA complex assembly and response to stimuli such as light, abiotic factors, and stress. In 12 M fish, the top enriched GO terms included circadian rhythm along with biosynthetic and metabolic processes. In 20 M fish, the most significant GO terms included glucose metabolism and carbohydrate or monosaccharide catabolism. The results indicate that age-related changes in diurnal genes are chronologically clustered into different biological processes that include cellular growth and anabolic processes in young animals, changing to genes involved in cellular maintenance in adults, and switching to catabolism in older animals.

Next, we performed disease enrichment on diurnal-regulated genes at each age to compare age-related changes with known disease markers. Disease Ontology (DO) analysis was conducted using DOSE [54] and revealed a correlation between gene expression and age-related diseases (Additional file 8). In addition to mood and bipolar disorder that appeared to manifest in older animals, genes implicated in diseases originating in other tissues were also present, suggesting that separate alterations in clock output in peripheral tissue may be a causative event in disease manifestation. Furthermore, we used the Age-Related Disease (ARD) database [55] and found an association between gene expression changes and age-related cognitive decline. For example, we found that *c3a.1* and *anxa3b*, genes implicated in neurodegenerative diseases such as Alzheimer’s disease and Parkinson’s [56], lost their diurnal rhythm at 20 M (Additional file 8b).

**Circadian clock remains constant regardless of age**

We next focused on the 39 genes (32 morning-specific and 7 evening-specific) that maintained diurnal expression regardless of age (Fig. 2a). GO analysis revealed the vast majority of genes are involved in circadian clock regulation (Fig. 2b). This subset of genes includes *arntl2*, *clocka*, *cry1a*, *cry2*, *per1b* and *per2* as well as *hif2*, *bhlhe40*, *fech*, *nr1d1* and *aldh4a1*, which all show distinctive diurnal expression (Fig. 2c). Consistent with other reports [13, 14], these data indicate that core clock genes are largely unaffected by age, suggesting there is an undefined age-related network that alters circadian output beyond the normal plasticity found in circadian oscillators.

**Age-specific diurnal genes with overlapping transcripts**

A fraction of diurnally expressed genes had a corresponding overlapping transcript so we examined how
the expression of these pairs changed with age with the idea that they may engage in transcriptional interference of represent a chromatin-regulating lncRNA and it’s cognate cis target. This concept is rooted in the observation that lncRNAs are known regulators of chromatin and can influence gene expression in both cis and trans [44].

In support of this notion, we found that 9.5, 6.6 and 12.4% of diurnal genes for the 3 age groups had an OT. Expression of representative diurnal genes with corresponding OT is in shown in Fig. 3a–d. As previously noted per2, which maintained a diurnal rhythm regardless of age, has a corresponding antisense transcript (per2AS), although in the RNA-seq per2AS lacks a readily apparent rhythm for the 2 diurnal time points tested here (Fig. 3a). However, when we examine a 24-h time course using strand-specific RT-PCR an antiphase rhythm is apparent (data not shown). Diurnal expression of other genes with OT was restricted to specific ages. For example, myl10 has diurnal expression at 4 M but not 12 and 20 M. Like per2AS, myl10AS does not have a detectible rhythm at any age (Fig. 3b, Additional file 9a).

In contrast, cishb and cishbAS both have diurnal expression with a similar phase (both transcripts are elevated at ZT4 and reduced at Z16) (Fig. 3c, Additional file 9b). Finally, myl1 exhibited diurnal expression at 20 M, but not 4 M and 12 M and its OT, myl1AS did not show a detectible rhythm at any age examined (Fig. 3d, Additional file 9c). RT-PCR results for these OT are consistent with the RNA-seq data, confirming OT are present at a subset of diurnal genes but do not...
necessarily display antiphasic rhythms; a finding consistent with results from liver [32]. However, our data are limited to 2 diurnal time-points, so we cannot rule out the possibility that these OTs have an oscillation with a different phase. In addition, the antisense transcripts were all expressed at much lower levels so detecting differences was inherently more difficult.

Diurnal and age-related changes to facultative heterochromatin

Given our observation of many diurnal lncRNAs and their potential role as regulators of chromatin, we proceeded to examine if there were age-related changes in facultative heterochromatin, specifically H3K9me3, which might account for the age-related changes in gene expression. We performed H3K9me3 chromatin immunoprecipitation (ChIP) from crosslinked zebrafish brain tissue at ZT4 and ZT16 at 4, 12 and 20 months of age.

Chromatin enriched with H3K9me3 was identified using MACS2 (q ≤ 0.01) [57] and differential diurnal distribution within and between ages defined by DiffBind [58]. Diurnal heterochromatin between ZT4 and ZT16 was observed at all ages (p ≤ 0.05) and H3K9me3 levels changed throughout the genome concurrent with change in age. For example there were 3424 rhythmic H3K9me3 peaks at 4 M, 3417 at 12 M and 9717 at 20 M (Fig. 4a a-c). To understand the distribution of H3K9me3 between coding and non-coding loci and the extent to which H3K9me3 oscillated at OT, the H3K9me3 peaks were broken down into those occurring in or near protein-coding genes (Fig. 4a d-f) or at loci containing predicted lncRNA (Fig. 4a g-i). The peaks were further classified into ones that occurred near OTs and these are highlighted in green. At 4 M, 14.2% of the diurnal H3K9me3 peaks occurred at annotated genes with an OT, while 8.3% occurred at a lncRNA with an OT; at...
12 M 22% occurred at annotated genes with an OT and 13.6% occurred at a lncRNA with an OT; and at 20 M, 11.2% of peaks occurred at annotated genes with an OT and 22.2% occurred at a lncRNA with an OT. We next examined the differences in genome location of H3K9me3 that occurred in protein-coding genes relative to regions containing lncRNA (Fig. 4b). At protein-coding genes, H3K9me3 tended to cluster in promoter regions 1–3 kb upstream of transcription start site, in introns and intergenic regions, whereas at lncRNAs, H3K9me3 mostly occurs in intergenic regions. This phenomena was likely observed because lncRNAs lack a 5' or 3' UTR and tend to be smaller with less splicing than their protein-coding counterparts. Collectively, the data indicate there are genome-wide diurnal oscillations in H3K9me3 that change in an age-dependent manner.

To further define age-related changes in facultative heterochromatin and examine the impact on age-related changes in gene expression, we began by selecting genes that displayed differential expression between 4 M and 20 M at the same time point (q ≤ 0.05). We found 266 transcripts had an age-related change at ZT4 and 462 at ZT16 (Fig. 5a, b). At ZT4, 76 annotated and 190 unannotated transcripts changed expression between 4 and 20 M. Error bars present SEM and the p-values are shown on the graphs.
20 M and at ZT16, 167 annotated and 295 unannotated transcripts displayed age-related changes (Additional files 10 and 11). Next, we sorted the genes with age-related changes into four groups: Group 1 was morning-specific genes that decrease with age, Group 2 was morning-specific genes that increase with age, Group 3 was evening-specific genes that decrease with age and Group 4 was evening-specific genes that increase with age. Consistent with previous findings [14], GO analysis revealed age-related changes to metabolic function were prevalent (Table 1). Other notable findings in our analysis indicate that cardiovascular fitness and regenerative potential decline with age (Group 1) whereas responses to hypoxia and low oxygen are endemic to aged animals (Group 4).

We next examined age-related changes to H3K9me3. Analysis of H3K9me3 between 4 and 20 M suggested H3K9me3 levels change for large portions of the genome as organisms age (Fig. 5c). This analysis shows an overall decrease in significant H3K9me3 peaks in 4 M versus 20 M at ZT4 and the opposite at ZT16. Changes occurred at protein-coding genes and lncRNA and were particularly pronounced for lncRNAs, some of which contained sense-antisense pairs. When compared with gene expression, H3K9me3 enrichment generally showed the expected and opposite pattern. For example, H3K9me3 was
elevated near slc4a1a at 4 M relative to 20 M and slc4a1a showed a corresponding decrease in expression at 20 M compared to 4 M (Fig. 5d, Additional file 12a-c). In this case, loss of H3K9me3 at ZT4 near slc4a1a with increasing age resulted in a gain of morning-specific diurnal expression that was exclusive to 20 M brain. Conversely, we also observed the opposite and found an age-related loss in diurnal expression and a corresponding...
increase in heterochromatin. This phenomenon is represented by *fbxoa32* (Fig. 5e), which displayed a diurnal oscillation at 4 M in the RNA-seq that was lost at 12 and 20 M. In this instance H3K9me3 levels increased with age at ZT16, which corresponded with the loss of expression. Despite the age-related changes in diurnal expression of the protein-coding genes and heterochromatin, the corresponding overlapping lncRNA did not always show an overt rhythm or expression differences among ages, indicating there is far more complexity to the system than just changes to lncRNA expression.

Of note, the size of H3K9me3 peaks around *slc4a1a* and *fbxoa32* were approximately the size of a single nucleosome (150–200 bp), as were many of the peaks in and around protein coding genes. There were also larger heterochromatin domains containing broader peaks and these typically occurred around unannotated transcripts (Additional file 13). In certain instances, the broad peaks had diurnal changes, but there were no apparent oscillations in the corresponding transcript.

### Discussion

Using multiple genomic approaches, the work reported here was designed as an initial test for the hypothesis that diurnal gene expression and age-related dysregulation is driven, in part, by lncRNA-guided heterochromatin changes. This hypothesis arose from a number of previous observations. First, H3K9me2 and H3K9me3-dependent facultative heterochromatin occurs at the clock genes during the repressive phase of the circadian cycle [24, 40, 42]. Second, the Per-complex contains KMT1/SUV39 along with a number of noncoding RNA binding proteins including NONO [24, 27]. Third, the *Per2* gene has an antisense transcript and in Neurospora, the *frq* NAT natural antisense transcript (NAT) *qrf* is needed for H3K9me3-dependent facultative heterochromatin [31, 32, 39]. Forth, there are multiple examples, from single-cell eukaryotes to mammals that reveal H3K9me3 can be directed by lncRNA ranging from centromeres in *S. pombe* [59] to H19-guided imprinting in mammals [60]. And finally, emerging models on aging suggest there is a redistribution of H3K9me3 and DNA methylation, which is either a direct cause or a consequence of aging. Taken together, these findings suggest age-related changes to the circadian transcriptome may occur in part due to age-related changes to

| GO annotation adj P value | Group 1: Decrease with Age | | Group 2: Increase with Age | | Group 3: Decrease with Age | | Group 4: Increase with Age |
|--------------------------|---------------------------|---|--------------------------|---|---------------------------|---|
| **Morning-Specific**      | heart contraction 3.70E-03| | short-chain fatty acid metabolic process 6.24E-05| | glycolysis 1.92E-06| | endothelial cell migration 2.00E-04|
| heart process 3.70E-03   | regulation of biological quality 2.00E-04| | monosaccharide metabolic process 3.32E-06| | monosaccharide metabolic process 3.32E-06| | immune system process 1.00E-03|
| fin regeneration 4.70E-03 | homeostatic process 6.00E-04| | hexose metabolic process 3.32E-06| | hexose metabolic process 3.32E-06| | response to decreased oxygen levels 1.10E-03|
| circulatory system process 5.50E-03| fatty acid metabolic process 1.40E-03| | glucose catabolic process 4.92E-06| | glucose catabolic process 4.92E-06| | response to hypoxia 1.10E-03|
| blood circulation 5.50E-03 | ion homeostasis 1.70E-03| | monosaccharide catabolic process 5.66E-06| | hexose catabolic process 5.66E-06| | response to oxygen levels 1.10E-03|
| tissue regeneration 7.80E-03 | chemical homeostasis 2.20E-03| | single-organism carbohydrate catabolic process 5.66E-06| | carbohydrate catabolic process 6.48E-06| | sprouting angiogenesis 1.50E-03|
| fin morphogenesis 1.48E-02 | monocarboxylic acid metabolic process 3.10E-03| | cellular lipid metabolic process 1.14E-02| | glucose metabolic process 1.52E-05| | erythrocyte homeostasis 2.70E-03|
| purine-containing compound biosynthetic process 1.88E-02 | cellular lipid metabolic process 1.14E-02| | carboxylic acid metabolic process 2.08E-02| | generation of precursor metabolites and energy 2.00E-04| | erythrocyte differentiation 2.70E-03|
| response to wounding 1.93E-02 | carboxylic acid metabolic process 2.08E-02| | o xoacid metabolic process 2.22E-02| | erythrocyte differentiation 2.70E-03| | embryonic hemopoiesis 2.90E-03|
| appendage development 1.93E-02 | monocarboxylic acid metabolic process 3.10E-03| | homeostasis of number of cells 3.10E-03| | homeostasis of number of cells 3.10E-03|

(Continued)
circadian IncRNA, which could then alter the underlying heterochromatin. The data presented here is by no means definitive proof this as an all-inclusive model, but does demonstrate a likelihood this occurs at some loci. The data also reveal other mechanisms must exist to account for changes in gene expression independent of H3K9me3.

In good agreement with previous observations, this report shows global reprogramming of the circadian transcriptome during aging even though the underlying circadian oscillator remains largely unaffected. For example, in the liver, age-induced changes in circadian expression are accompanied by changes in protein acetylation and NAD+ metabolism [14]. Similarly, extensive reprogramming of circadian transcriptome in aged stem cells causes a switch in expression of genes involved in homeostasis, to those involved in tissue-specific stress such as DNA damage or inefficient autophagy [13]. Our study contributes new insights into mechanisms underlying age-related changes to diurnal expression and further implicates epigenetic changes that include alterations to circadian IncRNA and genome-wide heterochromatin as potential regulators.

An additional objective of this study was to further identify IncRNAs in zebrafish brain. Considering the majority of expressed transcripts in mammals do not appear to code for proteins [61], these data should serve as a valuable resource to the zebrafish community. In particular, IncRNAs have diverse roles in regulation ranging from transcriptional interference in yeast, to X chromosome inactivation, imprinting, and epigenetic silencing in mammals, including directing H3K9me3 and H3K27me3 [62] so a more complete zebrafish noncode is warranted. Due to the growing evidence IncRNAs are targets of circadian regulation in a variety of tissues and organisms, it is important to get a base-line measure of IncRNAs to begin to understand their role in the clockworks. For example there are 112 IncRNAs showing differential expression in the light and dark in rat pineal gland [63] and mouse circadian transcriptome data suggests there are at least 1000 circadian regulated IncRNAs observed in multiple tissues [30]. Here, we expanded the zebrafish noncode by adding brain to a developmental dataset and explored whether any of these IncRNAs coincided with diurnal changes in H3K9me3. Further, we examined how IncRNA expression changed with age and time-of-day. Among the over 140,000 transcript isoforms identified via Cufflinks, more than 50% were previously unannotated in the GRCz10 and 50,524 are predicted to be IncRNAs by PLEK (both long intergenic non-coding RNA and OT). It remains to be determined how many of these newly identified transcripts are under direct control of the circadian clock and whether they function as heterochromatin regulators in either cis or trans.

Our findings of age-related changes to diurnal transcription are consistent with other reports showing the core circadian oscillator is relatively stable regardless of age, but ccg expression undergoes chronological changes over an animal's lifetime. In this study, not only are protein-coding ccgs changing with age, but also a significant number of rhythmic IncRNA also change as organism's age. From gene ontology analysis of protein-coding genes, it is clear the circadian transcriptome changes from anabolic to catabolic processes as one ages. The dynamic nature of age-related changes to clock output between 4 M and 12 M suggests the existence of a complex regulatory mechanism that partially resides at the level of chromatin and becomes unstable with age (20 M). This is quite consistent with the age-related changes in acetylation given deacetylation precedes heterochromatin.

We also examined age-related changes in diurnal genes by disease ontology analysis in hopes of identifying key factors that could provide insight into age-related disease. We specifically examined changes in the expression of Werner syndrome helicase (WRN) because mutations in WRN protein cause an advanced aging disorder known as adult progeria. WRN, like PER2, is associated with KMT1 (SUV39h) and HP1 and there is a reduction in heterochromatin in WRN null cells [47]. We did not observe any age-related changes to WRN (data not shown), nor did it have a diurnal expression (at least at the phase we examined). However, WRN interacting protein 1 (wrnip1) was lost with age (Additional file 8c), so it is interesting to speculate that an interaction between Wrinp1 and KMT1 might occur through Wrinp1 and this could potentially account for some of the age-related changes in heterochromatin. Alternatively, other alterations in heterochromatin may be mediated by changes in IncRNAs that potentially interact with the Per2 complex and direct Per-associated KMT1 to different loci; a speculative notion at this stage, but one that needs further study. Regardless, it is clear there is a complex regulatory system below the level of the master oscillator controlling circadian output that changes with age and occurs concurrent with changes in IncRNA, H3K9me3, and must include other chromatin changes.

As a final note, it is important to emphasize that the goal of this study was not to identify circadian protein-coding genes per se, this has been done exhaustively by others. As such, we intentionally ignored established guidelines for circadian rhythm research [64] and instead set out to examine a potential mechanism for age-related changes to gene expression, with the main focus being IncRNA discovery and heterochromatin. To that end, we limited our analysis to two diurnal time points, sequenced to extremely high depth (90 million paired-end reads), and preserved strandedness to augment transcript discovery. This focus on diurnal regulation...
proved fruitful as it allowed us to identify a much broader range of lncRNAs and analyze H3K9me3 in greater detail. It is now clear that a more comprehensive analysis of age-related changes to lncRNAs and corresponding chromatin structural changes on the circadian time scale is warranted to uncover more detailed mechanism(s) underlying aging.

**Conclusions**

There are age-related changes to lncRNA and H3K9me3 that coincide with changes in *ccg* expression. Based on the known function of lncRNA guiding epigenetic modifications in cellular events such as imprinting, X-chromosome inactivation, and NAT-mediated heterochromatin, it is possible that a subset of diurnal lncRNA may guide age-related epigenetic changes that affect *ccg* expression without altering the master circadian oscillator. This theory is supported by our observations showing diurnal and age-related changes in gene expression, including lncRNAs, coincide with diurnal and age-related changes in H3K9me3. This notion provides one potential solution to the perplexing observation that clock output is dynamic as organism’s age, but the core clock does not change appreciably [13, 14]. Our findings reveal that chromatin structure, in the form of circadian-regulated facultative heterochromatin, is one facet that determines whether a gene is rhythmic in young or older adults or when it is expressed in given block of time over one’s lifespan. Based on these and other findings, this may be one of many underlying causes of age-related disease manifestation observed with circadian disruption.

**Methods**

**Animal care**

Wild-type zebrafish (*Danio rerio*) (SAT) were obtained from ZIRC (Zebrafish International Resource Center, Oregon), housed in system water (conductivity 600 ppm, pH 7.4) at 27 °C, and fed twice daily. For diurnal entrainment, the fish were placed under a 12:12 light:dark cycle, but otherwise maintained at 14:10 light:dark cycle for breeding. Embryos and young larvae were raised in egg water (30 mg/L instant ocean in deionized water). For tissue extraction, fish were sacrificed by emersion in cold MS-222 (300 mg/l, Sigma), decapitated and dissected under Phosphate Buffered Saline (PBS) using microdissection tools.

**RNA-Seq**

Four Month (M), 12 M and 20 M zebrafish were maintained under a 12:12 light:dark cycle for one week prior to tissue removal. Zebrafish were sacrificed at ZT4 and ZT16 and brain tissue was removed as described [65]. Tissue was immediately snap frozen in liquid nitrogen and RNA was extracted from all samples simultaneously using TRIzol (Invitrogen). RNA-Seq libraries (*n* = 2) were processed at the Columbia Genome Center (New York, NY). Briefly, ribosomal RNA was depleted with RiboZero Gold (Illumina) and converted to cDNA with TruSeq Stranded Library Kit (Illumina). The libraries were sequenced on an Illumina 2500 to a depth of 90 million 100 bp paired-end reads per library. The data are deposited in Gene Expression Omnibus (GEO) under the accession number GSE109856.

**ChIP-Seq**

Zebrafish were maintained under identical conditions described for the RNA-seq. Brain tissue was removed and immediately cross-linked with 1% formaldehyde for 10 min then quenched with 0.1 M glycine for 10 min prior to being snap frozen in liquid nitrogen. Frozen tissue was lysed by mechanical disruption in the presence of ChIP lysis buffer [50.0 mM Hepes (pH 7.4), 150 mM NaCl, 1.0 mM EDTA, 1% Triton X-100, 0.1% deoxycholate] containing protease inhibitors (2.0 μg/mL leupeptin, 2.0 μg/mL pepstatin A, 1.0 mM PMSF). Additional cell disruption and chromatin shearing were done in a Misonix cup sonicator 5 times for 30 s at 20% power. The resulting lysates were cleared of cellular debris by centrifugation at 16,000 x g for 1 min. The average size of sheared chromatin was 500 bp. For the ChIP, 200 μg of lystate was mixed with 3 μg H3K9me3 antibody (Abcam ab8898) prebound to Protein A magnetic beads (Dynal) then incubated overnight at 4 °C. The ChIP samples were washed 5x with RIPA buffer (10 mM Tris pH 8.0, 1 mM EDTA, 1% Triton X-100, 0.1% SDS, 0.1% sodium deoxycholate, 140 mM NaCl) and eluted with 0.1 M sodium bicarbonate, 1% SDS. Isolated samples were heated at 65 °C to reverse crosslinks and protein was removed by the addition of protease K. DNA was further purified by phenol-chloroform extraction then precipitated with 1/10 volume 3 M NaOAc pH 5.4 and 2.5 volumes 95% ethanol. Purified DNA from each H3K9me3 ChIP done in triplicate was used for library preparation. DNA was sequenced 50 million 50-bp single-end reads on the Illumina HiSeq 2500 at Duke University Genome Center. The data are deposited in Gene Expression Omnibus (GEO) under the accession number GSE109856. H3K9me3 from the ChIP-seq was validated by qPCR using oligonucleotides contained in Additional file 14.

**RNA-seq analysis**

The RNA-seq data were processed using two different analysis pipelines roughly outlined in Additional file 1: Figure S1 and S2 for transcript discovery. The first method used Tophat2-Cufflinks-Cuffdiff and second used Hisat2-Stringtie-Cuffdiff [48, 49, 66]. To enhance transcript discovery, we also included previously published RNA-seq dataset from developing zebrafish embryos (GSE32989) [50]. The combined analysis yielded a single
ChIP-seq analysis
The reads of H3K9me3 ChIP-seq were mapped to the zebrafish GRCz10 genome using Bowtie2 [68]. Bam files of each triplicate were further processed using MACS2 for peak calling (FDR ≤ 0.01) [57]. Differential H3K9me3 enrichment between different conditions was analyzed by Diffbind and visualized with MAplot [58]. ChIPseeker was used to analyze genomic features of the H3K9me3 enrichment [69]. The H3K9me3 peaks were visualized with IGV [67].

Quantitative reverse transcription PCR
Zebrafish (n = 3) were maintained and entrained under identical conditions described for RNA-seq. Brain tissue was removed and immediately snap frozen in liquid nitrogen. TRIzol reagent was used for the extraction of total RNA from the brain tissue. 500 µg of total RNA was then used as a template with the SuperScript III First-Strand Synthesis kit (Invitrogen) to produce cDNA. The specific targets were amplified by RT-PCR using oligonucleotides in Additional file 14 using hmbs as the internal control. T-tests were performed to calculated p-value to determine significant difference between samples.

Gene and Disease ontology
WebGestalt (WEB-based GEne SeT AnaLysis Toolkit) was used for Gene Ontology (GO) enrichment using an adjusted p-value ≤0.01 as the cutoff, with multiple-hypothesis correction to calculate the enrichment p-value. The GO clusters with significant p-value were taken for further analysis. The top GO terms were converted to log₁₀ adjusted p-value and visualized with a radial graph or in tabular format [52, 53]. In order to use human Disease Ontology database, the zebrafish genes were first matched to their human orthologues by using Ensembl biomart (www.ensembl.org/biomart) and we used Bioconductor DOSE packages with p-value ≤0.05 cut to identify corresponding genes.

Additional files

Additional file 1: TopHat2/Cufflinks revealed 146,910 transcripts/isosforms with the min-isoform-fraction (-F) set to 0.05 (default setting). 16,532 out of the 146,910 isoforms were known protein-coding genes (class code =). Overlapping transcripts (OT) were identified using Cuffcompare by extracting transcripts with class code x and s from the merged GTF file. Transcripts with class code i, o and u were also extracted and further processed using Bedtools intersect (option -S) to sort antisense OT. In total, there were 14,774 OT fitting these criteria. The 14,774 OT were subjected to PLEK analysis to predict IncRNAs. PLEK revealed 5299 OT IncRNAs. 75% of the OT IncRNAs are unannotated intergenic transcripts with class code u. The remaining 25% are IncRNAs that overlap with an exon of a protein-coding gene. Of note, after PLEK, there were no single exon transcripts. (EPS 1498 kb)

Additional file 2: The HISAT2/StringTie analysis revealed 318,442 transcripts/isosforms with FPKM cutoff set to 0.1 in StringTie. This cutoff, which was an order a magnitude lower than the default, was necessary to identify the known per2AS transcript. HISAT2/StringTie identified 15,245 protein-coding transcripts (class code =). OTs were identified using a similar pipeline as described in Additional file 1: Figure S1 except we used Cuffcompare instead of Cuffmerge. Transcripts with class codes x, i, o or u were extracted and those with class code i, o, or u were processed with Bedtools intersect. In all, this revealed 183,635 potential OT. The 183,635 OTs were processed using PLEK yielding 131,635 predicted IncRNAs. However, after removing single exon transcripts, only 31,629 OTs remained. Of these, 76% were intergenic and the remaining 24% were contained within another gene. (EPS 1699 kb)

Additional file 3: Expression patterns of IncRNAs from early development through adult aged zebrafish. To enhance transcript discovery, we included stranded RNA-seq data from zebrafish embryogenesis [50]. In addition to allowing enhanced detection of IncRNAs, this dataset provided the ability to examine dynamic changes in IncRNA expression from bud stage to 20 M old fish. To identify all the IncRNAs, we filtered the RNA-seq data using the flowchart in panel a. We started with the 146,910 transcripts from Cufflinks and found 86,327 undefined transcripts (class code u). Next, PLEK predicted 50,524 potential IncRNA. The 50,524 transcripts were further filtered by their expression changes based on the criteria that a transcript must have a statistical change in expression in at least one-time point. This analysis yielded 17,702 unique IncRNAs. The 17,702 transcripts were then clustered based on similarities in the timing of expression. (b-i) Expression patterns of the 17,702 IncRNA from the bud stage of embryogenesis to 20 M fish segregated into eight groups based on similarity in expression. The thick black lines are the average pattern for each group. Also shown is a representative transcript from each group visualized with IGV (left panels). Certain IncRNAs were largely restricted to adult fish ≥4 M (Groups 1, 3, 5 and 6) while others were largely restricted to developmental stages (Groups 2 and 8). Group 7 appeared to span the later stages of development into adult fish ≥12 M. In contrast, Group 4 IncRNAs were expressed at every time point. Of note, the IncRNAs in Group 6 were largely restricted to adult fish and acquired a diurnal change when the fish were 12 M old. (EPS 1670 kb)
Additional file 8: Disease Ontology enrichment analysis of diurnal transcripts at different ages. We performed DO on protein-coding genes that had diurnal expression. (a) The top-ranked DO terms of diurnal genes in the specific age are illustrated in a dot graph ($p \leq 0.05$). (b) Bar plots of complement component 3a (C3a.1) and annexin 3b (annxa3b). Expression changes in both C3a.1 and annexa3b correlate with age-related cognitive decline [56]. (c) We also examined the expression of wimp1 and found that it is lost with age. (EPS 1310 kb)

Additional file 9: Expression of mylf10, cishb, and myl1 sense and antisense transcripts. IGV illustration showing relative expression of sense and antisense transcripts originated from (a) mylf10, (b) cishb and (c) myl1 for both time points for all three age-groups. The panel on the right shows relative expression validated by RT-PCR (n = 3). Error bars represent the SEM with asterisks indicating $p$-value < 0.05. (EPS 1721 kb)

Additional file 10: Genes that changed expression between 4 M and 20 M at ZT4. (XLSX 27 kb)

Additional file 11: Genes that changed expression between 4 M and 20 M at ZT16. (XLSX 38 kb)

Additional file 12: Expression and H3K9me3 for slc4a1a. (a-b) The expression of slc4a1a sense and antisense transcripts was confirmed by RT-PCR (n = 3). (c) H3K9me3 ChIP-qPCR at slc4a1a. Error bars represent the SEM and asterisks indicates p-value < 0.05. (EPS 962 kb)

Additional file 13: Representative view of H3K9me3 broad peaks. Broad peaks over 1 kb around unannotated transcripts were observed and illustrated in the IGV viewer. The H3K9me3 broad peaks were found in an unannotated transcript XLOC_030565 (a) and downstream of an unannotated transcript XLOC_028020 (b). The broader peaks spanning multiple nucleosomes showed a diurnal rhythm between ZT4 and ZT16. (EPS 932 kb)

Additional file 14: Oligonucleotides used in this report. (EPS 1201 kb)

Abbreviations

ChiP-seq: Chromatin Immunoprecipitation with DNA sequencing; CRF-H: Circadian regulated facultative heterochromatin; DO: Disease Ontology; GO: Gene ontology; H5S: HISAT2/StringTie, PLEK; IncRNA: Long noncoding RNA; OT: Overlapping transcripts; RNA-seq: RNA sequencing; T/C: TopHat2/Cufflinks

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Availability of data and materials

The RNA-seq and ChiP-seq datasets are available in the Gene Expression Omnibus (GEO) under accession number GSE109856.

Authors’ contributions

WJB conceived the project. JP performed the experiments. JP and WJB analyzed the data and wrote the manuscript.

Ethics approval and consent to participate

All animal procedures were performed in accordance with guidelines from the National Institutes of Health and approved by Rutgers University Institutional Animal Care and Use Committee (IACUC).

Consent for publication

Not applicable

Competing interests

The authors declare they have no competing interests.

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References

1. Pittendrigh CS, Minis DH. Circadian systems: longevity as a function of circadian resonance in Drosophila melanogaster. Proc Natl Acad Sci U S A. 1972;69(6):1537–9.
2. Pittendrigh CS, Daan S. Circadian oscillations in rodents: a systematic increase of their frequency with age. Science. 1974;186(4163):548–50.
3. Davidson AJ, Sellis MT, Daniel I, Yamazaki S, Menaker M, Block GD. Chronic jet-lag increases mortality in aged mice. Curr Biol. 2006;16(21):R914–6.
4. Reinke H, Asher G. Circadian clock control of liver metabolic functions. Gastroenterology. 2016;150(3):574–80.
5. Bass J, Takahashi JS. Circadian integration of metabolism and energetics. Science. 2010;330(6009):1349–54.
6. Green CB, Takahashi JS, Bass J. The meter of metabolism. Cell. 2008;134(5):728–42.
7. Landgraf D, Long JE, Poulov CD, Barandas R, Malinow R, Welsh DK. Genetic disruption of circadian rhythms in the suprachiasmatic nucleus causes helplessness, behavioral despair, and anxiety-like behavior in mice. Biol Psychiatry. 2016;80(11):827–33.
8. Young ME, Bray MS. Potential role for peripheral circadian clock dysynchrony in the pathogenesis of cardiovascular dysfunction. Sleep Med. 2007;8(8):656–67.
9. Masri S, Kinouchi K, Sassone-Corsi P. Circadian clocks, epigenetics, and cancer. Curr Opin Oncol. 2015;27(5):50–6.
10. Kondratov RV, Kondratova AA, Gorbacheva YY, Wkhovanets OV, Antoch MP. Early aging and age-related pathologies in mice deficient in BMAL1, the core component of the circadian clock. Genes Dev. 2006;20(14):1868–73.
11. Hofman MA, Swaab DF. Living by the clock: the circadian pacemaker in older people. Ageing Res Rev. 2006;5(1):33–51.
12. Kolker DE, Fukuyama H, Huang DS, Takahashi JS, Horton TH, Turek FW. Aging alters circadian and light-induced expression of clock genes in golden hamsters. J Biol Rhythm. 2003;18(2):159–69.
13. Sato S, Solanas G, Peixoto FO, Ebrahimi V, Madrigal I, Sone C, Moreira D, Datta D, Symeonidi A, Castellanos A, Weis PS, Caballero JM, et al. Aged stem cells reprogram their daily rhythmic functions to adapt to stress. Cell. 2017;170(4):678–92 e620.
14. Sato S, Solanas G, Peixoto FO, Bee L, Symeonidi A, Schmidt MS, Brenner C, Masri S, Benitha SA, Sassone-Corsi P. Circadian reprogramming in the liver identifies metabolic pathways of aging. Cell. 2017;170(4):664–77 e611.
15. Antoch MP, Song EJ, Chang AM, Vitaterna MH, Zhao Y, Willsbacher LD, Sangoram AM, King DP, Pinto LH, Takahashi JS. Functional identification of the mouse circadian clock gene by transgenic BAC rescue. Cell. 1999;89(4):655–67.
16. King DP, Zhao Y, Sangoram AM, Willsbacher LD, Tanaka M, Antoch MP, Steeves TD, Vitaterna MH, Kornhauser JM, Lowrey PL, et al. Positional cloning of the mouse circadian clock gene. Cell. 1997;89(4):641–53.
17. Vitaterna MH, King DP, Chang AM, Kornhauser JM, Lowrey PL, McDonald JD, Dove WF, Pinto LH, Turek FW, Takahashi JS. Mutagenesis and mapping of a mouse gene, clock, essential for circadian behavior. Science. 1994;264(5159):719–25.
18. Hogenessch JB, Gu YZ, Jain S, Bradfield CA. The basic-helix-loop-helix-PAS orphan MOP3 forms transcriptionally active complexes with circadian and hypoxia factors. Proc Natl Acad Sci U S A. 1998;95(10):5474–9.
19. Zheng B, Larkin DW, Albrecht U, Sun ZS, Sage M, Eichele G, Lee CC, Bradley A. The mPer2 gene encodes a functional component of the mammalian circadian clock. Nature. 1999;398(6728):627–30.
20. Cermakian N, Monaco L, Pando MP, Dierich A, Sassone-Corsi P. Altered behavioral rhythms and clock gene expression in mice with a targeted mutation in the Period1 gene. EMBO J. 2001;20(15):3967–75.
21. Shearman LP, Jin X, Lee C, Reppert SM, Weaver DR. Targeted disruption of circadian resonance in Drosophila melanogaster. Proc Natl Acad Sci U S A. 1997;94(7):2968–72.
22. van der Horst GT, Muijtjens M, Kobayashi K, Takano R, Kanno S, Takao M, de Wit J, Verkerk A, Bier WP, van Leenen D, et al. Mammalian Cry1 and Cry2 are essential for maintenance of circadian rhythms. Nature. 1999;398(6728):627–30.
24. Duong HA, Weitz CJ. Temporal orchestration of repressive chromatin modifiers by circadian clock period complexes. Nat Struct Mol Biol. 2014; 21(2):126–32.

25. Duong HA, Robbins MS, Knutti D, Weitz CJ. A molecular mechanism for circadian clock negative feedback. Science. 2011;332(6036):1436–9.

26. Kim YJ, Kwak PB, Weitz CJ. Specificity in circadian clock feedback from targeted reconstitution of the NuRD corepressor. Mol Cell. 2014;56(6):73–48.

27. Brown SA, Ripperger J, Kadener S, Fleury-Olela F, Vilbós F, Rosbash M, Schibler U. PER1/DO1-associated proteins modulate the negative limb of the mammalian circadian oscillator. Science. 2005;308(5722):693–6.

28. Sato TK, Panda S, Miraglia LJ, Reyes TM, Rudic RD, McNamara P, Naik KA, FitzGerald GA, Kay SA, Hogness JB. A functional genomics strategy reveals Rora as a component of the mammalian circadian clock. Neuron. 2004;49(4):527–37.

29. Preitner N, Dimioli F, Lopez-Molina L, Zakany J, Duboule D, Albrecht U, Schibler U. The orphan nuclear receptor REV-ERBalpha controls circadian transcription within the positive limb of the mammalian circadian oscillator. Cell. 2002;110(2):251–60.

30. Zhang R, Lahens NF, Balasse EI, Hughes ME, Hogeness JB. A circadian gene expression atlas in mammals: implications for biology and medicine. Proc Natl Acad Sci U S A. 2014;111(45):16219–24.

31. Kolke N, Yoo SH, Huang HC, Kumar V, Lee C, Kim TK, Takahashi JS. Transcriptional architecture and chromatin landscape of the core circadian clock in mammals. Science. 2012;338(6105):349–54.

32. Vollmers C, Schmitz RJ, Nathanson J, Yeo G, Ecker JR, Panda S. Circadian pathways in the mouse by the circadian clock. Cell. 2002;109(3):307–18.

33. Kay SA, Takahashi JS, Hogenesch JB. Coordinated transcription of key circadian clock genes by the circadian clock. Cell. 2002;110(2):251–62.

34. De Seta MP, Miller BH, Si AI, Schook AB, Straume M, Schultz PG, Kay SA, Takahashi JS, Hogeness JB. Coordinated transcription of key oscillators of protein-coding and regulatory RNAs in a highly dynamic mammalian liver epigenome. Cell Metab. 2012;16(6):833–45.

35. Kramer C, Lorus JJ, Dunlap JC, Crosthwaite SK. Role for antisense RNA in regulating circadian clock function in Neurospora crassa. Nature. 2003;421(6926):948–52.

36. Panda S, Antoch MP, Miller BH, Si AI, Schook AB, Straume M, Schultz PG, Kay SA, Takahashi JS, Hogeness JB. Circadian gene expression and regulation in the mouse. Proc Natl Acad Sci U S A. 2010;107(3):207–12.

37. Hurley JM, Dasgupta A, Emerson JM, Zhou X, Ringelberg CS, Knabe N, Lipzen AM, Lindquist EA, Daum CG, Barry KW, et al. Analysis of clock-regulated genes in Neurospora reveals widespread posttranscriptional control of metabolic potential. Proc Natl Acad Sci U S A. 2014;111(48):16995–7002.

38. Storch KF, Lipian O, Leykin I, Wixanathan N, Davis FC, Wong WH, Weitz CJ. Extensive and divergent circadian gene expression in liver and heart. Nature. 2002;417(6884):78–83.

39. Takahashi JS. Transcriptional architecture of the mammalian circadian clock. Nat Rev Genet. 2017;18(3):164–79.

40. Belden WJ, Lewis ZA, Selker EU, Loros JJ, Dunlap JC. CHD1 remodels chromatin at frequency and represses light-activated gene expression. G3 – Genes, Genomes, Genet. 2011;7(7):680–7.

41. Robinson JT, Thorvaldsdottir H, Winckler W, Guttman M, Lander ES, Getz G, Mesirov JP. Integrative genomics viewer. Nat Biotechnol. 2011;29(1):24–7.

42. Cribbs DH, Berchtold NC, Pernaa V, Coleman PD, Rogers J, Tenner AJ, Cotman CW. Extensive innate immune gene activation accompanies brain aging, increasing vulnerability to cognitive decline and neurodegeneration: a microarray view. J Neuroinflammation. 2012;9:179.

43. Zhang Y, Liu T, Meyer CA, Eckhoute J, Johnson DS, Bernstein BE, Nusbaum C, Myers RM, Brown M, Li W, et al. Model-based analysis of ChIP-Seq (MACS). Genome Biol. 2008;9(9):R137.

44. Marchini JS, Myers RM, Abecasis GR, Cardon LR, Torkamani A, Fry B, et al. A new method for the detection of complex interactions in genome-wide association studies. Nat Genet. 2010;42(11):937–44.

45. Robinson JT, Thovardottir H, Winckler W, Guttman M, Lander ES, Getz G, Mesirov JP. Integrative genomics viewer. Nat Biotechnol. 2011;29(1):124–6.

46. Langmead B, Trapnell C, Pop M. Ultrafast and memory-efficient alignment of short DNA sequences to the genome. Genome Res. 2009;19(10):1547–54.

47. Zhang W, Li J, Suzuki K, Qu J, Wang P, Zhou J, Liu X, Ren R, Xu X, Ocampo A, et al. Aging stem cells. A Werner syndrome stem cell model unveils heterochromatin alterations as a driver of human aging. Science. 2015;348(6239):1160–3.

48. Trapnell C, Roberts A, Goff L, Pertea G, Kim D, Kelley DR, Pimentel H, Saberberg SL, Rinn JU, Pachter L. Differential gene and transcript expression analysis of RNA-seq experiments with TopHat and cufflinks. Nat Protoc. 2012;7(3):562–78.

49. Pertea M, Kim D, Pertea GM, Leek JT, Salzberg SL. Transcript-level expression analysis of RNA-seq experiments with HISAT, StringTie and Ballgown. Nat Protoc. 2016;11(9):1650–67.

50. Pauli A, Valen E, Lin MF, Garber M, Vastenhouw NL, Levin JZ, Fan L, Sandellin A, Rinn JU, Regev A, et al. Systematic identification of long noncoding RNAs expressed during zebrafish embryogenesis. Genome Res. 2012;22(5):577–91.

51. Li A, Zhang J, Zhou Z. PLEK: a tool for predicting long non-coding RNAs and messenger RNAs based on an improved k-mer scheme. BMC Bioinformatics. 2014;15:311.

52. Zhang B, Kirov S, Snouddy J. WebGestalt: an integrated system for exploring gene sets in various biological contexts. Nucleic Acids Res. 2005;33(Web Server):W741–8.

53. Wang J, Duncan D, Shi Z, Zhang B. WEB-based Gene Set Analysis toolkit (WebGestalt): update 2013. Nucleic Acids Res. 2013;41(Web Server issue):W7–83.

54. Wang J, Duncan D, Shi Z, Zhang B. WEB-based Gene Set Analysis toolkit (WebGestalt): update 2013. Nucleic Acids Res. 2013;41(Web Server issue):W7–83.

55. Yoon J, Wang LG, Yan GR, He QY. DOSE: an R/Bioconductor package for disease ontology semantic and enrichment analysis. Bioinformatics. 2015;31(6):688–9.

56. Taub C, Strauch LJ, Brennan BM, Golumbic JM, Tison L, Yip M, et al. Diaphanosoma brachyurum as a model organism to study circadian rhythms in nematodes. J Mol Biol. 2012;421(6):948–59.

57. Robinson JT, Thovardottir H, Winckler W, Guttman M, Lander ES, Getz G, Mesirov JP. Integrative genomics viewer. Nat Biotechnol. 2011;29(1):124–6.

58. Langmead B, Trapnell C, Pop M, Saberbarg SL. Ultrafast and memory-efficient alignment of short DNA sequences to the genome. Genome Res. 2009;19(10):1547–54.

59. Yu G, Wang LG, Yan GR, He QY. DOSE: an R/Bioconductor package for disease ontology semantic and enrichment analysis. Bioinformatics. 2015;31(6):688–9.

60. Pfitzner N, Damiola F, Lopez-Molina L, Zakany J, Duboule D, Albrecht U, Schibler U. The orphan nuclear receptor REV-ERBalpha controls circadian transcription within the positive limb of the mammalian circadian oscillator. Cell. 2002;110(2):251–60.