Paraspeckles as rhythmic nuclear mRNA anchorages responsible for circadian gene expression

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ABSTRACT
Circadian clocks regulate rhythmic gene expression levels by means of mRNA oscillations that are mainly driven by post-transcriptional regulation. We identified a new post-transcriptional mechanism, which involves nuclear bodies called paraspeckles. Major components of paraspeckles including the long noncoding RNA Neat1, which is the structural component, and its major protein partners, as well as the number of paraspeckles, follow a circadian pattern in pituitary cells. Paraspeckles are known to retain within the nucleus RNAs containing inverted repeats of Alu sequences. We showed that a reporter gene in which these RNA duplex elements were inserted in the 3′-UTR region displayed a circadian expression. Moreover, circadian endogenous mRNA associated with paraspeckles lost their circadian pattern when paraspeckles were disrupted. This work not only highlights a new paraspeckle-based post-transcriptional mechanism involved in circadian gene expression but also provides the list of all mRNA associated with paraspeckles in the nucleus of pituitary cells.

KEYWORDS
circadian oscillators; circadian rhythm; nuclear RNA retention; paraspeckle; post-transcriptional mechanism

Most organisms have built-on time-measuring devices that are commonly known as circadian clocks. These circadian clocks allow them to anticipate the time of day and hence to temporally organize behavior as well as physiologic and biochemical processes. They therefore play a key role as adaptive mechanisms to permanent changes in the environment necessary for the individual survival and the sustainability of the species. In living systems ranging from bacteria to humans, circadian rhythms are generated endogenously through genetic control and regulate vital aspects of the organism physiology, from sleeping and waking to neurotransmitter secretion and cellular metabolism. At the center of these rhythms is the circadian clock machinery that consists in a transcription-translation feedback system regulated by a group of genes that oscillate in a circadian manner, the so-called clock genes. In mammals, the circadian system is hierarchically organized. Indeed, while molecular oscillations occur in most cells and tissues of the body, a central structure, the suprachiasmatic nucleus (SCN) of the hypothalamus, functions as the master regulator to synchronize the phase of the other slave oscillating tissues.

Briefly, the molecular mechanism that generates circadian rhythms involves the interacting positive and negative feedback loops of transcriptional or translational processes of clock genes (Fig. 1). In mammals, 2 basic helix-loop-helix transcription factors, Circadian Locomotor Output Cycles Kaput (CLOCK) and Brain and Muscle Aryl Hydrocarbon Receptor Nuclear Translocator-Like Protein 1 (ARNTL also named BMAL1), heterodimerize and subsequently bind to conserved E-box sequences in target gene promoters. In this manner, this complex activates the transcription of mammalian Period (Per1, Per2, Per3) and Cryptochrome (Cry1, Cry2) genes. The PERs and CRYs proteins are expressed, post-translationally modified, feedback to inhibit their own transcription and are then degraded to lead to a
new round of BMAL1:CLOCK mediated transcription
(For a review see ref. 7). Another regulatory loop is
mediated by the orphan nuclear receptors, the Reti-
noic Acid Receptor-Related Orphan Receptor
(RORα/b/g) and the Reverse Erb α/β (Rev-erb α/β),
that activate and inhibit, respectively, transcription of
Bmal1 through the retinoic acid Receptor Response
Element (RRE) in its promoter, leading it to oscillate
in a circadian manner (Fig. 1).

In addition to the core regulation at the level of
transcription or translation, circadian clock proteins
are also subjected to extensive post-translational
modifications that appear to control their cellular
localization, protein stability, and activity. For exam-
ple, Casein Kinase Iε and Iδ (CKIε/δ) are known to be
critical factors that regulate the turnover of PERs and
CRYs in mammals, 8–10 however, kinase CKIε also acti-
vates BMAL1-mediated transcription 9 (Fig. 1).

Importantly, circadian transcription factors not
only regulate their own transcription but also regulate
the expression of numerous other clock-controlled
genes 6 [CCGs; (Fig. 1)]. Over the past decade, clock
gene transcriptional regulation has been described in
many species and tissues, where it drives rhythmic

Figure 1. Molecular mechanisms of the circadian clockwork in mammals. In mammals, rhythmically transcribed BMAL1 heterodimerizes
with CLOCK, and together they bind to target E-boxes in the promoters of Per, Cry, Rev-erb, and Ror. PER and CRY proteins are synthe-
sized in the cytoplasm and may be phosphorylated by CK1 kinases including CK1ε/δ. PER/CRY heterodimers translocate to the nucleus
where they repress their own transcription, generating a near 24-h feedback loop. A second feedback loop represses or activates the
transcription of Bmal1 through the actions of REV-ERB or ROR, respectively. BMAL1/CLOCK heterodimer drives oscillating expression of
clock-controlled genes with E-box or RORE containing promoters. Post-transcriptional events in the life cycle of a (pre-) mRNA that have
been reported to directly influence the circadian clock and/or to be controlled in a circadian manner include splicing, nuclear retention
and cytoplasmic export, regulation by miRNA and polyadenylation at the 3′ end. Translation and degradation of the mature mRNA are
rhythmic processes as well.
mRNA expression. By use of techniques such as microarrays, a large fraction of the mRNA population (up to 10-15% of all mRNAs in a single mammalian tissue) has been shown to display a rhythmic expression that has been initially assumed to result from temporal changes in transcription. However, data from mouse liver demonstrate poor correlation between the activation of a promoter and the amount of the corresponding transcript for genes that are rhythmic at the steady-state level. Actually, with the development of high-throughput sequencing, results obtained in the last years indicate that approximately 43% of the mammalian genome is rhythmic and analysis of circadian nascent RNA has allowed to show that less than 30% of circadian mRNA are regulated by de novo transcription, suggesting that post-transcriptional regulation contributes mostly to rhythmic mRNA expression (Fig. 1). Much of what we initially knew about post-transcriptional regulation came from studies of fungi, plants and flies (For a review see), but circadian post-transcriptional mechanisms involved in rhythmic control of mRNA expression have now also been reported in mammals at many different levels (For a review see), such as RNA splicing, poly-adenylation, mRNA stability, mRNA cytoplasmic export and RNA nuclear retention (Fig. 1).

RNAs can be retained in the nucleus by particular bodies called paraspeckles. These nuclear bodies are found in almost all of the cultured cell lines and primary cultures from tissues, except for embryonic stem cells. Paraspeckles are detected as discrete dots found in inter-chromatin space, close to nuclear speckles. A long noncoding RNA, nuclear-enriched abundant transcript one (Neat1) is the structural component (Fig. 2). While a short and a long transcript previously identified as MENe (Neat1–1) and MENb (Neat1–2), respectively, are generated from the same promoter, Neat1–1 alone cannot induce paraspeckle formation since specific depletion of Neat1–2 leads to disruption of paraspeckles. While paraspeckles detected by RNA FISH of Neat1 appeared as round foci when visualized under a confocal microscope, we showed that they appeared more likely as oblong structures with smaller dimensions after use of a combination of Neat1 RNA FISH and Super Resolution STORM analysis (as designed in Fig. 2). Paraspeckles have been shown to retain in the nucleus RNAs containing duplex structures. This has been shown for the mouse cationic amino acid transporter 2 (Cat2) transcribed nuclear RNA, CtnRNA, an alternatively spliced form of the Cat2 mRNA, which contains a dsRNA structure resulting from inverted short inter-spersed nuclear elements (SINEs) in its 3'-UTR. In primate cells, the most common inverted repeated SINEs are Alu elements. Alu elements are unique to primates and account for almost all of the human SINEs and for more than 10% of the genome and inverted repeat structures (inverted repeated Alu elements [IRAlus]) occur frequently in gene regions. Paraspeckles have been shown to retain in the nucleus mRNAs containing IRAlus in their 3'-UTRs like Nicolin 1 (NICN1) or Lin28.

The nuclear retention of mRNAs containing IRAlus by paraspeckles can be regulated by a methyl-transferase, CARM1 (coactivator-associated arginine methyltransferase), which control both the binding capacity of some paraspeckle proteins to mRNAs containing IRAlus as well as NEAT1 transcription and then paraspeckle formation. Among the 40 paraspeckle proteins identified thus far, researchers classified 4 RNA-binding proteins, including 3 members of the Drosophila melanogaster behavior human splicing (DBHS) family proteins (NONO, PSPC1 and SFPQ) and RNA-binding motif protein 14 (RBM14) as major paraspeckle protein components (Fig. 2). We had previously reported that 2 of these major protein components of paraspeckles, namely NONO and SFPQ, display a circadian expression pattern in primary cultures of pituitary cells as well as in a rat pituitary cell line, the GH4C1 cells. In this latter cell line, we found thereafter that 2 other major paraspeckle proteins, PSPC1 and RBM14, display also a circadian pattern. All 4 proteins further bind rhythmically to Neat1 and Neat1 itself displays a circadian expression pattern. The expression of the long form of Neat1 RNA, Neat1–2, that is known to be sufficient for the formation of paraspeckle displays also a circadian pattern. In addition we showed that circadian expression of these different components leads to rhythmic variations in paraspeckle number within the cells. Thanks to their circadian expression pattern and given their presumed functions in gene expression through corresponding mRNA nuclear retention, we asked whether paraspeckle bodies can rhythmically retain RNAs in the nucleus leading to a rhythmic expression of the corresponding gene. This hypothesis was first tested with a reporter gene. Indeed, by using...
a construct of the EGFP reporter gene fused to an IRAlu and by transfecting the construct into GH4C1 cells, we obtained evidence that IRAlu elements inserted in 3′-UTR of egfp reporter mRNA allow for its circadian retention within the nucleus. This rhythmic nuclear retention is abolished after disruption of paraspeckles by siRNA or Neat1 antisens oligonucleotides. Using real-time video microscopy, IRAlu elements inserted in 3′-UTR of egfp reporter mRNA was shown to cause rhythmic cytoplasmic expression of the EGFP protein. Paraspeckles through their circadian expression, could then control circadian expression pattern of a reporter gene containing IRAlu elements in 3′-UTR. We then asked whether this circadian post-transcriptional regulation exerted by paraspeckle bodies applies to endogenous genes. To address this issue, it was necessary to determine which mRNA are associated with paraspeckle bodies in the nucleus of GH4C1 cells. To this end, we developed a hybridization-based strategy that uses complementary oligonucleotides to purify Neat1 RNA together with its RNA targets from reversibly cross-linked extracts. Two antisense biotinylated oligonucleotide probes that target accessible regions of Neat1 RNA, as predicted by modeling its secondary structure by bioinformatics, were designed and used for Neat1 RNA specific pull-down whereas one biotinylated irrelevant probe was used for Neat1 RNA non-specific pull-down. Genes which were considered specifically associated with paraspeckles exhibited values of fragment per kilobase per million of mapped reads (FPKM) higher than 1 and were common to the lists obtained with the 2 specific probes. Overlapping list represented 65% of the list obtained with one probe and 83% of the list obtained with the other one. By comparing this gene list to available data in the literature such as the list of rhythmic mRNA in the mouse pituitary or the mRNA whose rhythm is post-transcriptionally controlled in the liver of mice, we achieved significant recovery rates of 18 and 27%, respectively. These rates are probably underestimated since they come from the comparison of gene lists obtained in different structures and species.

By selecting a few genes from our list, we showed that the mRNA of these genes exhibit a circadian rhythm of their nuclear retention and that this rhythm is abolished when paraspeckles are disrupted. These results obtained on endogenous mRNA, together with results obtained with a reporter gene, allow to conclude that paraspeckles participate in the post-transcriptional control of rhythmic gene expression (Fig. 2). Besides deciphering a post-transcriptional mechanism involved in the circadian regulation of gene expression, a problematic anchored in the field of chronobiology, our study from an endocrine point of view allows to assign a role in the physiology of the pituitary gland to the long non-coding RNA Neat1 whose functions remain enigmatic.

Although it has been shown that paraspeckles can retain in the nucleus genes which have sequences of IRAlu type in their 3′-UTR, endogenous mRNA we found associated with Neat1, actually do not contain such sequences or equivalent IR-SINE sequences in our model of rat pituitary cells. This suggests

Figure 2. Structure and functional implication of paraspeckles in the rhythmic expression of mRNA. Paraspeckles are schematically drawn as oblong structures organized around the short (Neat1–1) and long (Neat1–2) transcripts of the long non-coding Neat1 RNA. Major protein entities of paraspeckles are also shown in the scheme. The roles of paraspeckles in circadian gene expression is schematically represented. The rhythmic number of paraspeckles inside the cells drives a circadian nuclear retention of paraspeckles-associated mRNA and thus leads to a circadian gene expression.
either that paraspeckles can recognize IR-sequences localized anywhere along the mRNA or that double-stranded RNA structures with such IR-sequences are not the only one recognized by paraspeckles. The challenge will be now to elicit the nature of the sequences together with their position along the mRNAs that are involved in the binding of paraspeckles and therefore form the basis of circadian expression.

In conclusion, circadian regulation has been investigated mainly at the transcriptional or the post-translational level, and RNA-based mechanisms contributing to this regulation are only beginning to emerge. Therefore, the impact of such mechanisms on circadian biology remains to be evaluated. However, given the finding that the majority of circadian mRNAs is not regulated by de novo transcription and given from the study presented here, the high number of mRNA that are potentially regulated by the new post-transcriptional mechanism we described, we predict all these post-transcriptional mechanisms will be shown to play extensive and widespread roles in circadian biology. It is also tempting to imagine that after the identification of mechanisms involved in post-transcriptional control of the circadian clock, connections to diseases will follow. Actually presumably because circadian rhythms play a key role as adaptive mechanisms to permanent changes in the environment, their deregulation is associated with many diseases the best known of which are neuropsychiatric, metabolic, cardiovascular disorders and cancers. A better understanding of the molecular mechanisms that govern the mammalian circadian clock is thus required to first identify connections to diseases, which may then give opportunities for new therapeutic concepts. Therapeutic targeting of paraspeckles could be for instance the basis for new strategies to control circadian rhythms, with the aim of improving human disease associated with rhythmicity dysfunction, in particular rhythmic hormonal dysfunctioning.

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