An alternative *D. melanogaster* 7SK snRNP

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**Abstract**

**Background:** The 7SK small nuclear RNA (snRNA) found in most metazoans is a key regulator of P-TEFb which in turn regulates RNA polymerase II elongation. Although its primary sequence varies in protostomes, its secondary structure and function are conserved across evolutionary distant taxa.

**Results:** Here, we describe a novel ncRNA sharing many features characteristic of 7SK RNAs, in *D. melanogaster*. We examined the structure of the corresponding gene and determined the expression profiles of the encoded RNA, called snRNA:7SK:94F, during development. It is probably produced from the transcription of a lncRNA which is processed into a mature snRNA. We also addressed its biological function and we show that, like dm7SK, this alternative 7SK interacts in vivo with the different partners of the P-TEFb complex, i.e. HEXIM, LARP7 and Cyclin T. This novel RNA is widely expressed across tissues.

**Conclusion:** We propose that two distinct 7SK genes might contribute to the formation of the 7SK snRNP complex in *D. melanogaster*.

**Keywords:** 7SK snRNA, Drosophila, P-TEFb, Long non-coding RNA

**Background**

Transcriptional elongation by RNA polymerase II (RNA Pol II) is regulated by the positive transcription elongation factor b (P-TEFb) composed of Cyclin T1 and cyclin-dependent kinase 9 (CDK9) [1]. P-TEFb is required for the release of RNA Pol II, which is paused at proximal promoter, into productive elongation. This is accomplished by the phosphorylation of both the 5,6-dichloro-1-D-ribofuranosylbenzimidazole (DRB) sensitivity-inducing factor (DSIF) and the Negative Elongation Factor-E (NELF-E) [2], but also by the phosphorylation of serine 2 residues on the C-terminal domain of the large subunit of RNA Pol II [3–5]. The activity of P-TEFb is regulated by the reversible association of the kinase with the 7SK snRNP [6, 7]. P-TEFb is maintained in an inactive state in the 7SK snRNP by a direct interaction with the RNA-bound, hexamethylene bisacetamide-induced proteins (HEXIM1/2) [8, 9]. 7SK is a 332-nt long non-coding RNA (lncRNA) transcribed by RNA Pol III [10–12]. In the nucleus, the 7SK snRNA is constitutively associated with Methylphosphate Capping Enzyme (MePCE; BCDIN3 homolog in *Drosophila*) which methylates its 5′ end [13], and with La-Related protein 7 (LARP7) which binds to the 3′ end [14–17]. These two factors contribute to the stability of the RNA [18–20].

The genes encoding 7SK are found in all vertebrates and the sequence of the snRNA is highly conserved in mammals, whereas basal vertebrates only exhibit poorly conserved sequences (e.g. ~ 68% similarities in lamprey) [21]. Nevertheless, their 5′ and 3′ stem-loop structures remain evolutionarily conserved, both structurally and functionally, probably because they physically interact with HEXIM, P-TEFb, MePCE and LARP7. In 2008, Gruber and colleagues [22, 23] discovered 7SK snRNA-like sequences in the genome of several protostomes.
(arthropods, mollusks, and annelids). Further analyses of more basal protostomes have uncovered a 7SK snRNA candidate in Caenorhabditis species [24], although this is still debated as it could be a homologue of the U8 snoRNA [25]. A structural analysis of the already known 7SK snRNAs has shown several highly conserved motifs in bilaterians. Thus, M1, M3 and M8 motifs may form the core structure and are closed together to stabilize the entire structure of the RNA in a “closed” structural model [24, 26]. A recent and broader analysis of the structure of invertebrate 7SK RNAs defined an additional conserved inner stem-loop structure [27]. Overall, 7SK sequences are poorly conserved among protostomes, which explains why BLAST-based searches on mammalian sequences as bait have been mostly unsuccessful. We have previously identified and characterized the 7SK snRNP containing the 444-nts 7SK RNA, P-TEFb, HEXIM, LARP7, and MePCE in Drosophila [28]. This indicates that the 7SK snRNA pathway is conserved in metazoans.

In this work, we identified a novel non-coding snRNA in Drosophila genome with a bioinformatic analysis based on the structure of the Drosophila RNA Pol III-specific promoters [29]. We argue that this RNA is probably not transcribed by RNA Pol III but is more likely to be formed after the cleavage of an IncRNA precursor transcribed by RNA Pol II. We document its expression levels during the life cycle, embryogenesis and organogenesis of Drosophila. We also demonstrate that this snRNA physically interacts with dmHEXIM, and co-immunoprecipitates with dmP-TEFb. We propose that this novel snRNA named snRNA:7SK-94F is a Drosophila-specific alternative 7SK snRNA. This is the first description of two distinct 7SK gene products which could participate in the formation of the P-TEFb snRNP complex.

Results
A novel snRNA in D. melanogaster
The primary sequence of the 7SK snRNA is poorly conserved over large evolutionary distances, which limits the sensitivity of searches based on sequence similarities. Indeed, a BLAST search [30] failed to detect 7SK related sequences in Drosophila genome. Instead, we derived a profile Hidden Markov Model based on a structural alignment of chordates 7SK sequences (see Materials and Methods and Fig. 1a).

In total, six loci showed significant similarities to the 5′ or the 3′ end of the model, but only a single one of them showed >100 bp similarity with both the 5′ and the 3′ of the model, suggesting that it corresponds to a 7SK-like RNA gene. The other loci (chr4:1169608..1169632; chrX:4501439..4501471; chr2R:12354215..12354269; chr2L:17785962..17785988; chr3L:3230335..3230369) were spurious hits arising from bioinformatic noise and were not studied further. First, we identified the Transcriptional Start Site (TSS) of this gene by 5′ RACE PCR (see Materials and Methods), which is an adenine located at 3R:23432869 (Flybase release FB2018_06) [31] (Additional file 1 Fig. A1). If the 3′ end of the gene is defined by a double track of 4 and 6 thymidines at 3R:23433272, which serves as the termination signal for RNA Pol III transcription, the putative gene would be 404 bp long (Additional file 1 Fig. A1). It is located in the 94F3 cytological band, in the second intron of the CG4374-B transcript, which encodes a protein containing a zinc finger domain of unknown function (http://flybase.org). Then we analysed the upstream region of this gene by comparison to the well-known organization of the promoters of Drosophila snRNA genes [29, 32, 33]. In D. melanogaster, the promoters of RNA Pol II (U1 to U5) and RNA Pol III (U6) snRNA genes contain two elements: an upstream Proximal Sequence Element (PSE), and a PSEB (for RNA Pol II snRNA genes) [34] or a TATA box (for U6 snRNA gene) [35]. The upstream sequences display some similarities to snRNA promoter structure: a PSE is separated by 8 bp from a TATA box, which is located 18 bp upstream of the TSS (Fig. 1b). Of note, the PSE of 7SK-like RNA contains a TC dinucleotide at position 19 and 20, typically associated with snRNA transcription by RNA Pol III in Drosophila (Fig. 1c). The upstream region of this locus presents only slight similarities with a TATA box although it contains a G at its 3′ end, as in insects (Fig. 1d) [29]. The internal sequence of the 7SK-like RNA gene displays 5 tracks of 4 to 5 thymidines, mainly in the first hundred nts, which are efficient RNA Pol III transcription termination signals in 75 to 95% of genes [36]. This implies that this gene, despite its Pol III-like promoter structure, is probably not transcribed by RNA Pol III, in contrast to 7SK snRNA genes found in other organisms.

RNA Pol II transcription profiling by nascent transcripts analysis with PRO-seq (Precision Run-On Sequencing) on 2–4 h embryos [37] revealed that this locus is strongly transcribed over a 4.7 kbp region (Fig. 2a). Processing and analysis of RNA-seq data generated from total RNAs extracted from Drosophila pupae further show strong evidence for accumulation of RNAs at the expected location (Fig. 2a). RNAs accumulation is highly consistent with PRO-seq signal. In addition, the peak of PRO-seq is slightly offset, between 20 and 60 nts downstream of the transcription start site, as observed for most genes [38].

This locus probably encodes a lncRNA that we named lncRNA:94F according to the current nomenclature. The 5′ end of this transcription unit corresponds precisely to the 7SK-like RNA sequence identified in the bioinformatic screen. It is likely that the lncRNA:94F RNA is a
precursor transcript later processed into the short mature 7SK-like snRNA.

The sequence of the lncRNA:94F gene is well conserved in all Drosophilidae species but not in other insects, particularly in the 5′ and 3′ ends (Fig. 2b). The predicted secondary structure of the 7SK-like RNA displays motifs similar to the M1, M3 and M8 motifs of the 7SK snRNA. These motifs are involved in the physical interactions within the 7SK snRNP complex [24] (Additional file 1 Fig. A2). There are also two evolutionary conserved regions, the stem A (previously called M4 and M5 motifs) [24] and the stem B [27], shared by Hexapoda. Moreover, the 5′ and 3′ ends are brought together to form the typical core structure of the 7SK RNA which is directly involved in the scaffolding of the P-TEFb inactive complex [18]. Interestingly, the different conserved structural motifs are located in the regions that are well conserved among the Drosophilidae (Fig. 2b). This structural analysis suggests that the 5′ end of the lncRNA:94F folds into a stable secondary structure promoting the downstream cleavage and the release of the short 7SK-like RNA, named snRNA:7SK:94F. In addition, to demarcate the processed RNA, we used RT-PCR with various forward and reverse primer sets spanning the genomic region (Additional file 1 Table A1). To increase the detection sensitivity, we carried out a southern blot on PCR-products using a radioactive probe mapping to the transcribed snRNA:7SK:94F region (Fig. 3a).

A band of the expected size is amplified only after reverse transcription (with primer PU15) followed by PCR with primers located in the predicted transcription unit (PU11 and PU12) (Fig. 3b). No stable transcript could be detected when using a reverse primer located just downstream of the putative cleavage site (PU14, Fig. 3a) or using a forward primer located upstream from the putative TSS (PU10, Fig. 3b).

The snRNA:7SK:94F is ubiquitously expressed
We first profiled snRNA:7SK:94F expression using RT-sqPCR, from the embryonic stages until adulthood (Fig. 4).
Notably, we found snRNA:7SK:94F transcripts before zygotic activation, suggesting that it is of maternal origin in the early developmental stages. Whereas snRNA:7SK:94F transcription levels were low at the first-instar larval stage, expression strongly increased from the second-instar larva and reached peak levels at the adult stage. Of note, this is significantly different from the expression levels of the dm7SK snRNA [23], which was highly expressed at all stages during the life cycle (Fig. 4). The expression level of snRNA:7SK:94F is lower than dm7SK snRNA, between 1 to 50 times less, depending on the developmental stage (Additional file 1 Fig. A3).

We next examined in more detail the expression patterns of snRNA:7SK:94F transcripts by in situ hybridization during embryogenesis and organogenesis. snRNA:7SK:94F transcripts are ubiquitously expressed during early embryonic development (Additional file 1 Fig. A4). During the formation and extension of the germ band, snRNA:7SK:94F transcripts were strongly expressed in this embryonic zone, which was the most transcriptionally active region of the embryos. It is also expressed at later stages of embryogenesis. Likewise, during organogenesis, snRNA:7SK:94F transcripts were found in all the imaginal discs tested (eye-antenna, leg or wing discs) and larval organ brain-optic lobes (Additional file 1 Fig. A4). We also performed control experiments to ensure that these signals are specific. Indeed, no signal was detected when a sense probe against snRNA:7SK:94F was added (data not shown).
snRNA:7SK:94F physically interacts with HEXIM

To test whether this novel snRNA shares functional properties with the 7SK RNA, we looked for physical interactions with the components of P-TEFb complexes [28]. We first performed immuno-precipitation assays with protein extracts from adult flies to explore interactions between the snRNA:7SK:94F RNA and dmHEXIM, dmLARP7 or dmCyclin T proteins. As expected, neither dmHEXIM nor dmLARP7 protein was detected in the bound fraction without antibody, whereas dmHEXIM antibodies efficiently pulled-down both dmHEXIM and dmLARP7 (Fig. 5a).

Fig. 4 snRNA:7SK:94F is expressed during the life cycle of D. melanogaster. The transcription levels of the housekeeping rp49 gene (lower panel) were used to normalize, by semi-quantitative RT-PCR, the expression levels of snRNA:7SK94F (upper panel) and dm7SK (middle panel). Larva 1, 2 and 3 correspond to larval stages. The size of the RT-PCR products are indicated on the right side of the picture.
Also, dmCyclin T antibodies pulled-down dmHEXIM and dmLARP7, showing that they form a complex in vivo. Importantly, the snRNA:7SK:94F RNA co-purifies with dmHEXIM, dmLARP7 and dmCyclin T, and is completely depleted from the unbound fractions (Fig. 5b). In agreement with previous reports [28], the dm7SK RNA was also pulled-down with this complex (Fig. 5b).

We then used the yeast three-hybrid system [39, 40] to map the interaction domains of HEXIM and the snRNA:7SK:94F RNA (Fig. 6a).

Since RNAs longer than 150 to 200-nt may affect the efficiency of the assay [42], we performed the experiments with the full-length snRNA:7SK:94F RNA (nt 15–404), but also with three truncated fragments (nt 15–163; nt 164–304, and nt 305–404). The positive control is composed of human 7SK (h7SK) snRNA (nt 1–175) and human HEXIM1 (hHEXIM), which are known to interact with each other [41]. Anti-sense sequences were used as negative controls. As expected, the co-expression of MS2-h7SK (nt 1–175) and hHEXIM produced a dark blue color with X-Gal assay (Fig. 6b), indicating physical interaction between these two partners. In contrast, a light blue color was observed with MS2-h7SK (nt 1–330) and hHEXIM1, as already observed (not shown) [41]. A light blue color was detected with both dm7SK or snRNA:7SK:94F and dmHEXIM, suggesting a weak or transient (but significant) interaction (Fig. 6b). Interestingly, the h7SK RNA displayed a strong interaction with dmHEXIM indicating that hHEXIM and dmHEXIM are functionally interchangeable. By contrast, Drosophila snRNA:7SK:94F did not interact with hHEXIM (Fig. 6b). The activation of the reporter genes using these constructs was also confirmed by monitoring the β-galactosidase activity in liquid assays (Fig. 6c). We observed a 3-fold increase of the β-galactosidase activity when MS2-dm7SK:94F fragments (nt 15–404 or nt 305–404) were co-expressed with dmHEXIM (Fig. 6c). These observations suggest that snRNA:7SK:94F interacts with dmHEXIM mainly through its 3′ end. Thus, in Drosophila, this experiment confirms that dmHEXIM interacts with two distinct 7SK snRNAs.
Discussion

In this study, we discover a novel 7SK-like snRNA in *Drosophila* genomes. We propose that it is generated after cleavage of a 4.7 kb IncRNA precursor, probably transcribed by RNA Pol II. The mature RNA displays strong structural similarities with known 7SK snRNA, and engages physical interactions with the various components of the P-TEFb complex. For these reasons, we propose that this novel processed IncRNA is an analogue of the 7SK snRNA and shares overlapping biological activities. According to the international nomenclature of ncRNA, we named the precursor, IncRNA:94F and the mature RNA, snRNA:7SK:94F.

The 7SK snRNA is highly abundant in eukaryotic cells and is well known for its crucial role in the regulation of RNA Pol II-mediated transcription. Although they have been found in the genome of many vertebrates (from humans to lampreys) [21], homologues of 7SK snRNA have been only recently described in several protostomes [22–24, 27]. Notably, the 7SK *Drosophila* homologue is able to sequester P-TEFb into a large and catalytically inactive complex contributing to RNA Pol II elongation control [28, 43], as seen in vertebrates.

7SK snRNA homologues have been found in bilaterians and follow precisely the phylogenetic distribution of HEXIM [24]. This is not surprising since the components of molecular complexes often exhibit very similar phylogenetic distribution. RNAs generally co-evolve with their respective partners and they are subject to strong structural constraints, albeit with a relaxed selection at the sequence level [44, 45]. As such, the sequence of IncRNA is often poorly conserved and distant homologues/analogues are notoriously difficult to find and many of these may have been missed in genomes. Of note, the internal part of 7SK snRNAs is less constrained by physical interactions with a partner (e.g. HEXIM, P-TEFb, LARP7 and MePCE), which certainly relaxes locally the selection pressure and may favours sequence variations. This easily explains the difference of the gene length observed in Drosophilidae (~ 400 nts) compared to Vertebrates (~ 300 nts). In addition, the evolution of IncRNA is known to occur at a fast pace and genome
There are many examples where the biological function of lncRNA does not depend on its primary sequence but rather on its 3D structure [47]. All the known 7SK snRNAs display a core structure containing M1, M3, and M8 motifs [24]. The stem A is well conserved as well, although it has not yet been linked to a function. In invertebrates, the 7SK snRNAs, together with snRNA:7SK:94F, present an additional conserved stem B structure [27]. We were able to detect this novel 7SK analogue because of the exquisite sensitivity of HMMs, but also because its 3’ end (which physically interacts with LARP7) is fairly similar to that of humans. Importantly, this novel 7SK displays all the most conserved regions found among the invertebrates 7SK RNA.

An intriguing fact is that the sequence of snRNA:7SK:94F presents several internal tracks of thymidines which act as RNA Pol III transcriptional termination signals. It is therefore highly unlikely that it is an RNA Pol III gene. An interesting hypothesis would be that the RNA Pol III promoter structure evolved over time into an RNA Pol II promoter which overcomes premature termination. Accordingly, the promoter specificity is not dictated by the TATA box but rather by the sequence of the PSE [32]. Such a functional shift would only require a few point mutations and would be very rapid over evolutionary time. The transcription unit almost certainly depends on its primary sequence but rather on its 3D structure [46]. There are several internal tracks of thymidines which act as RNA Pol III transcriptional termination signals. It is therefore highly unlikely that it is an RNA Pol III gene. An interesting hypothesis would be that the RNA Pol III promoter structure evolved over time into an RNA Pol II promoter which overcomes premature termination. Accordingly, the promoter specificity is not dictated by the TATA box but rather by the sequence of the PSE [32]. Such a functional shift would only require a few point mutations and would be very rapid over evolutionary time. The transcription unit almost certainly depends on its primary sequence but rather on its 3D structure [46].

What are the functional connections (if any) between these two 7SK genes found in Drosophila? Do they functionally cooperate or is their functions antagonistic or independent of each other (despite interacting with the same regulatory complex)? The fact that more of one 7SK sequence is present in a genome is not new. For example, mammalian genomes contain a single functional 7SK gene but they are populated with numerous 7SK-like retro-pseudogenes [49]. This raises the question of the possible role of those pseudogenes as molecular sponge, similarly to PTENP1 which is the pseudogene of PTEN protein [50]. Moreover, many lncRNAs function as molecular sponges for miRNA or RNA Binding Protein (RBP) to limit the abundance of their functional binding sites [51, 52]. In previous study, two 7SK gene copies were found in C. elegans genome, but the expression of only one could be detected by Northern blot, probably due to the very low expression level of the other. The authors suggest that they are paralogues subjected to concerted evolution because their sequence is nearly identical [23]. Likewise, expression of snRNA: 7SK:94F is challenging to measure as we were unable to detect it by Northern blot and we had to use highly optimized RT-PCR protocol to detect it. Nonetheless, the case for Drosophila is quite different from C. elegans: the sequences of the dm7SK and the proposed snRNA:7SK:94F are poorly conserved and display distinct expression patterns during development. This suggests that the two genes are not fully redundant and may have specific biological functions. Also, one can note that the phylogenetic distribution of snRNA:7SK:94F is restricted to the Drosophila genus (Fig. 2), suggesting that this snRNA is an innovation specific to this group and may have a Drosophila-specific function.

Proteins can often accommodate various molecular functions. For example, Actin proteins are components of the cytoplasmic acto-myosin network, but they are also nuclear components of the chromatin remodeling complex [53]. Beta-catenin, a membrane bound protein and transcription factor, is another example [54]. This is much less common for RNA, and although several cases have been reported [55, 56], their corresponding function is often restricted to the biology of nucleic acids. For example, U1 snRNA presents different roles in regulation of genes expression [57]. In humans, the 7SK snRNA itself has different roles, and some are dependent on the P-TEFb complex [58] and some are independent [59, 60]. Therefore, given the known function of HEXIM, it is tempting to speculate that the function of the HEXIM/snRNA:7SK:94F complex may be related to transcriptional regulation. It is clear however, as stated above, that snRNA:7SK:94F and dm7SK snRNAs do not have redundant functions because snRNA:7SK:94F is differentially expressed with lower levels in the larval and pupal stages whereas dm7SK displays high and constant expression during development. Although snRNA:7SK:94F physically interacts with the components of the P-TEFb complex, it does so in a slightly different manner to dm7SK, since the GAUC stem, which is the contact interface between HEXIM and 7SK, is mutated into GAUG [61]. Mutations in this 5’-hairpin have been extensively studied and some can affect the binding efficiency to HEXIM [62]. Thus, the point mutation found in snRNA:7SK:94F may have a functional impact in vivo. An interesting hypothesis (albeit untested) would be that snRNA:7SK:94F could be a molecular sponge aimed at
regulating the abundance of the proteins forming the 7SK snRNP complex.

Materials and methods
Identification of a 7SK-like analogue in D. melanogaster
Given that ncRNA 7SK sequences are poorly conserved, we used the HMMER package (www.hmmer.org) to look for signatures of 7SK sequences. To this end, a structural alignment built from the 7SK sequence of humans, mice, chicken, xenopus, fugu, tetraodon, lamprey, myxin and amphioxus was used to set up a nucleotidic multiple alignment, which was used to derive a profile HMM model with hmmbuild. The model was used to scan the *Drosophila* genome (FB2018_06 release) [31] with hmmssearch. Alignments were local for both the genome and the model.

Processing of NGS datasets
Sequence reads were mapped on Dmel v6 genome available from UCSC website by using the bowtie short reads aligner [63] with stringent mapping parameters (−m 1 -n 1 -l 49 -p 20). Reads were clipped down to 50 bp, as this does not limit mapping efficiency and specificity but prevents issues with spliced alignments. Genome wide density profiles were computed with genomeCoverageBed from the BEDTOOLS package [64] and converted to bigwig with the wigToBigWig software, from the Jim Kent suite available at http://hgdownload.soe.ucsc.edu/admin/jksrc.zip. Visualization of reads density and the genomic environment is carried out with the JBROWSE genome browser [65]. RNA-Seq datasets (SRA reference PRJNA644503; https://www.ncbi.nlm.nih.gov/bioproject/PRJNA644503) from wild type pupae are from [66] and PRJNA644503; https://www.ncbi.nlm.nih.gov/bioproject/PRJNA644503). The secondary structure of hs7SK snRNA, snRNA:7SK:94F, and dm7SK snRNA was computed with the mfold software, a predictive RNA-folding program [68]. The structure of hs7SK snRNA served as a control, since it has been extensively studied by different methods [12, 24]. Using mfold, the hs7SK snRNA is folded into a structure similar to the one determined by Wassarman and Steitz [12].

Secondary structure
In situ hybridization
In situ hybridizations of embryos (a mix of 0-4 h and 0-18 h aged embryos) and larval tissues were carried out with standard procedures [69] using appropriate primers (Additional file 1 Table A1). Each experiment was replicated three times on at least 30 embryos and 15 larvae, and lead to identical results. Embryos or imaginal discs were mounted on slides and captured by microscope (Zeiss, X100).

Co-immunoprecipitation
Co-immunoprecipitations were performed using rabbit antibodies against isoform A of dmHEXIM [28], and sheep antibodies against dm Cyclin T [70]. Ten micrograms of purified antibodies were bound and crosslinked.
to 30 μl of G/A sepharose magnetic beads (Crosslink magnetic CoIP kit; Pierce). Then, 750 μg of protein extract from 60 adult flies were incubated overnight at 4 °C with antibodies. The unbound fraction was harvested and the bound fraction was eluted into 50 μl. Protein partners were analysed by western blot using primary antibodies dmHEXIM (1/2000) and dmlARP7 (1/2000) [28]. A Clean Blot IP buffer (Thermo-Scientific) was used as a secondary antibody at 1/1000. Immunodetection was performed with ECL (GE Healthcare) and imaged with the Fusion Fx7 system (BioRad). RNA partners from the bound or unbound fractions were extracted and analysed with specific primers (Additional file 1 Table A1).

Yeast three-hybrid system
The yeast three hybrid system was used as previously described [39, 42]. pIIIA/MS2-derived plasmids carry the URA3 gene and express snRNA:7SK:94F fragments fused to two MS2 sites. The pACTII-derived plasmids carry the LEU2 gene and drive the production of the dmHEXIM isoform A (dmHEXIM A) [28] fused to the Gal4 activation domain. A dmHEXIM A PCR fragment was cloned into the BamHI and Xhol sites of pACTII, after amplification from the pET21-HEXIMA plasmid [28] using specific primers (Additional file 1 Table A1). Full-length or truncated snRNA:7SK:94F fragments were amplified from genomic DNA of D. melanogaster CantonS strain (Bloomington Drosophila Stock Center) and cloned into the Smal sites of pIIIA/MS2.1 using specific primers (Additional file 1 Table A1). Qualitative and quantitative analyses of β-galactosidase activities were performed as previously described [42].

Supplementary Information
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Authors’ contributions
Conceptualization, DN, NB and PU. performed experiments, DN, NB, OF and PU. Analysed data, DN, NB, DHP and PU. Resources and methodology, AAM and OB. Original and critical writing, NB, DHP, OB and PU. All authors have contributed to the work, read and approved the manuscript.

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Availability of data and materials
The datasets analysed during the current study are available from the SRA website under the reference PRJNA644503, and the GEO repository under the reference GSE131160.

Declarations
Ethics approval and consent to participate
Not applicable.

Consent for publication
Not applicable.

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

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References
1. Price DH. P-TEFb, a cyclin-dependent kinase controlling elongation by RNA polymerase II. Mol Cell Biol. 2000;20(8):2629–34. https://doi.org/10.1128/MCB.20.8.2629-2634.2000.
2. Fujinaga K, Irwin D, Huang Y, Taube R, Kurosu T, Peterlin BM. Dynamics of human immunodeficiency virus transcription: P-TEFb phosphorylates RD and dissociates negative effectors from the transactivation response
element. Mol Cell Biol. 2004;24(2):787–95. https://doi.org/10.1128/mcb.24.2.787-795.2004.

3. Bowman EA, Kelly WG. RNA Polymerase II transcription elongation and Pol II CTD Ser2 phosphorylation: A tail of two kinases. Nuc (United States). 2014;5:224–36.

4. Kwik H, Lis JT. Control of transcriptional elongation. Annu Rev Genet. 2013;47(1):483–508. https://doi.org/10.1146/annurev-genet-110711-155440.

5. Zhou Q, Li T, Price DH. RNA helicase and 7SK snRNP elongation control. Annu Rev Biochem. 2012;81(1):119–43. https://doi.org/10.1146/annurev-biochem-051010-102419.

6. McNamara RP, Bacon CW, D’Oro I. Transcription elongation control by the 7SK snRNP complex: releasing the pause. Cell Cycle. 2015;14(16):2115–23. https://doi.org/10.1124/mol.115.398410.2016.1181241.

7. Peterlin BM, Brogie JE, Price DH. 7SK snRNA: a noncoding RNA that plays a major role in regulating eukaryotic transcription. Wiley Interdiscip Rev RNA. 2011;2(2):103–119. https://doi.org/10.1002/wrna.106.

8. Nguyen VT, Kiss T, Michels AA, Bensaud O. 7SK small nuclear RNA binds to and inhibits the activity of CDK9/cyclin T complexes. Nature. 2001;414(6861):222–5. https://doi.org/10.1038/35104581.

9. Yang Z, Zhu Q, Luo K, Zhou Q. The 7SK small nuclear RNA inhibits the CDK9/cyclin T1 kinase to control transcription. Nature. 2001;414(6861):317–22. https://doi.org/10.1038/35104575.

10. Murphy S, Tripodi M, Melli M. Nucleic acids research-a sequence upstream from the coding region is required for the transcription of the 7SK RNA genes. Nucleic Acids Res. 1986;14(23):9243–50. https://doi.org/10.1093/nar/14.23.9243.

11. Kührer W, Benecke BJ. Structural and functional analysis of a human 7 S K RNA gene. J Mol Biol. 1987;195(1):31–41. https://doi.org/10.1002/2000-0022-2836(1987)90325-1.

12. Wassarman DA, Steitz JA. Structural analyses of the 7SK ribonucleoprotein (RNP), the Most abundant human small RNP of unknown function; 1991.

13. Krüger W, Benecke BJ. RNA Polymerase II transcription elongation and Pol II CTD Ser2 phosphorylation: A tail of two kinases. Nuc (United States). 2014;5:224–36.

14. Kwik H, Lis JT. Control of transcriptional elongation. Annu Rev Genet. 2013;47(1):483–508. https://doi.org/10.1146/annurev-genet-110711-155440.

15. Zhou Q, Li T, Price DH. RNA helicase and 7SK snRNP elongation control. Annu Rev Biochem. 2012;81(1):119–43. https://doi.org/10.1146/annurev-biochem-051010-102419.

16. Muniz L, Egloff S, Kiss T, Michels AA, Bensaud O. 7SK small nuclear RNA binds to and inhibits the activity of CDK9/cyclin T complexes. Nature. 2001;414(6861):222–5. https://doi.org/10.1038/35104581.

17. Uchikawa E, Natchiar KS, Han X, Proux F, Roblin P, Zhang E, et al. Structural analyses of the 7SK ribonucleoprotein (RNP), the Most abundant human small RNP of unknown function; 1991.

18. Kwik H, Lis JT. Control of transcriptional elongation. Annu Rev Genet. 2013;47(1):483–508. https://doi.org/10.1146/annurev-genet-110711-155440.

19. Zhou Q, Li T, Price DH. RNA helicase and 7SK snRNP elongation control. Annu Rev Biochem. 2012;81(1):119–43. https://doi.org/10.1146/annurev-biochem-051010-102419.

20. McNamara RP, Bacon CW, D’Oro I. Transcription elongation control by the 7SK snRNP complex: releasing the pause. Cell Cycle. 2015;14(16):2115–23. https://doi.org/10.1124/mol.115.398410.2016.1181241.

21. Gürsoy HC, Koper D, Benecke BJ. The vertebrate 7S K RNA separates hagfish from higher vertebrates: a phylogeny based on RNA-protein interactions. Nucleic Acids Res. 2017;45(11):6864–68. https://doi.org/10.1093/nar/gkw159.

22. Grüber AR, Koper-Emde D, Marz M, Donath A, Verstraete N, Nguyen VT, et al. The vertebrate 7S K RNA separates hagfish from higher vertebrates: a phylogeny based on RNA-protein interactions. Nucleic Acids Res. 2017;45(11):6864–68. https://doi.org/10.1093/nar/gkw159.

23. Grüber AR, Koper-Emde D, Marz M, Donath A, Verstraete N, Nguyen VT, et al. The vertebrate 7S K RNA separates hagfish from higher vertebrates: a phylogeny based on RNA-protein interactions. Nucleic Acids Res. 2017;45(11):6864–68. https://doi.org/10.1093/nar/gkw159.

24. Marz M, Donath A, Verstraete N, Nguyen VT, Stadler PF, Bensaude O. Evolution of 7SK RNA and its protein Partners in Metazoa. Mol Biol Evol. 2009;26(12):2821–30. http://www.ncbi.nlm.nih.gov/sutils/static/. https://doi.org/10.1093/molbev/msp198.
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