1 rDNA nascent transcripts promote a unique spatial organization during

2 mouse early development

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31 Abstract

32 During the first cell cycles of the early development, the chromatin of the embryo is highly 33 reprogrammed alongside that embryonic genome starts its own transcription. The spatial 34 organization of the genome is a major process that contributes to regulating gene transcription 35 in time and space, however, it is poorly studied in the context of early embryos. To study the 36 cause and effect link between transcription and spatial organization in embryos, we focused 37 on the ribosomal genes, that are first silent and begin to transcribe during the 2-cell stage in 38 mouse. We demonstrated that ribosomal sequences are spatially organized in a very peculiar 39 manner from the 2-cell to the 16-cell stage with transcription and processing of ribosomal 40 RNAs excluding mutually. Using drugs inhibiting the RNA polymerase I, we show that this 41 organization, totally different from somatic cells, depends on an active transcription of 42 ribosomal genes and induces a unique chromatin environment that favors major satellite 43 sequences transcription after the 4-cell stage.

44

45 Abbreviations

- 46 5'ETS: 5' externally transcribed spacer
- 47 ActD: Actinomycin D
- 48 EGA: Embryonic Genome Activation
- 49 hphCG: hours post-injection of human Chorionic Gonadotrophin
- 50 H3K9ac: Histone 3 acetylated in lysine 9
- 51 H3K4me3: Histone 3 tri-methylated in lysine 4
- 52 ITS1: internal transcribed spacer 1
- 53 ITS2: internal transcribed spacer 2
- 54 Major sat: mouse major satellite sequences
- 55 NPB: Nucleolar Precursor Body
- 56 Nopp140: Nucleolar phosphoprotein of 140 kD
- 57 mPN: maternal pronucleus
- 58 pPN: paternal pronucleus
- 59 PVP: Poly Vinyl Pyrrolidone
- 60 RT-qPCR: Real-Time quantitative PCR
- 61 RNA polymerase I: RNA pol I
- 62 rDNA: ribosomal DNA genes/sequences
- 63 rRNA: ribosomal RNA
- 64 UBF: Upstream Binding Factor
- 65

66 Introduction

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In eukaryotes, the spatial organization of the genome within the interphasic nuclei is 68 not random. DNA fluorescent in situ hybridization (DNA-FISH) experiments have 69 70 demonstrated that chromosomes occupy a specific nuclear position called territories ((Cremer 71 and Cremer, 2001) and that transcriptional activity may dictated gene positioning reviewed in 72 (Meaburn, 2016). The three-dimension (3D) organization of chromatin acts as a key 73 component of the cell identity and can be correlated with highly differentiated cell types 74 (Solovei et al., 2009). However, the relationships between genome organization and gene 75 transcription is still a matter of debate, albeit after extensive studies reviewed in (van Steensel 76 and Furlong, 2019). For a decade, chromosome conformation capture techniques (3C, 4C and 77 Hi-C) offer the possibility to study whole-genome organization, long-distance interaction, and 78 loops between genomic regions or loci (reviewed in (Bonev and Cavalli, 2016). But 79 depending on the biological model, DNA-FISH and microscopy are still relevant and at least 80 complementary to these approaches (Szabo et al., 2020) allowing single-cell analysis in intact 81 nuclei (Gelali et al., 2019) and localization of repeated sequences such as pericentromeric and 82 centromeric regions of ribosomal DNA (rDNA) genes (Maiser et al., 2020; Mayer et al., 83 2005).

As for mammalian embryos, 3D-FISH of pericentromeric regions (comprising minor and major satellite sequences, (Guenatri et al., 2004)) has been extensively used to demonstrate that these regions undergo large-scale reorganization during the early embryonic development period (reviewed in (Jansz and Torres-Padilla, 2019). We and others have shown that dramatic changes occur throughout the two first cell cycles of mouse preimplantation development concomitantly with the onset of embryonic transcription (also called embryonic genome activation (EGA)) (Aguirre-Lavin et al., 2012; Bonnet-Garnier et al., 2018; Probst et

91 al., 2007). This spatial reorganization of heterochromatin was shown to be required for further 92 development (Casanova et al., 2013; Probst et al., 2010), highlighting its importance. 93 Remarkably, in 1-cell and early 2-cell mouse embryos, major (and minor) satellite sequences 94 surround dense, spherical structures called nucleolar precursor bodies (NPBs -(Fléchon and 95 Kopecny, 1998)). While, the NPBs were first described as a seed for embryonic nucleolus to 96 settle (Zatsepina et al., 2003), it is now believed that they serve rather as a structural platform 97 anchoring heterochromatin and allowing its remodeling (Fulka and Langerova, 2019). At the 98 time of EGA, satellite sequences indeed progressively disconnect from these NPBs while 99 forming round shape clusters as found in somatic cells and called chromocenters (Aguirre-Lavin et al., 2012). Recently, (Hamdane et al., 2016) have shown that embryos lacking the 100 101 nucleolar protein UBF also lack NPBs and display an abnormal distribution of 102 heterochromatin. However, the precise relationship between these sequences and the NPBs 103 remains unclear.

104 The inner organization of NPBs has been thoroughly investigated (Baran et al., 1995; 105 Baran et al., 2001; Koné et al., 2016) by immunofluorescent staining of nucleolar proteins 106 such as Upstream Binding Transcription Factor (UBTF), Fibrilarin, B23/Nucleophosmin 1 107 (NPM1), and Nopp140/NOLC1 (Nucleolar and Coiled-body phosphoprotein 1) showing 108 dynamic redistribution of the different nucleolar compartments (mainly the dense fibrillar 109 component, (DFC) and the fibrillar center, (FC)) between the 2-cell and blastocyst stages 110 (time of implantation). The NPBs also structurally support the ribosomal genes (rDNA) 111 (Romanova, 2006). Remarkably, reinitiating of ribosomal transcription (rRNA synthesis) has 112 been shown - by BrUTP incorporation (Zatsepina et al., 2003) - to occur in NPBs at the end of 113 the 2-cell stage, when pericentromeric repeats undergo massive reorganization. It could be 114 that the chromatin state of both rDNA and pericentromeric sequences influence each other as described in Embryonic stem cells (Guetg et al., 2010; Savić et al., 2014). Little is known 115

116 regarding the spatial organization and expression of rDNA genes in the nuclei of mouse 117 embryos.

118 In the somatic nuclei, the ribosomal genes are transcribed in the nucleolus by the RNA 119 polymerase I (RNA pol I) in a long precursor transcript called 47S pre-rRNA, that will be 120 cleaved by several exonucleases to separate the internal transcribed spacers 1 (ITS1) and 2 121 (ITS2) and the 5' and 3' external transcribed spacers (5'-ETS and 3'-ETS) from the mature 122 ribosomal RNAs (rRNAs): 18S, 5.8S, and 28S (Henras et al., 2015). These rRNAs are 123 associated with ribosomal proteins in pre-ribosomes particles, processed/matured by several 124 proteins, and exported in the cytoplasm to form the small and large ribosome subunits 125 (Mullineux and Lafontaine, 2012). Two proteins: the UBF and the Selective factor 1 (SL1) 126 are required to load the Pol I complex and initiate transcription of the rDNA (Moss et al., 127 2019). Experimental disruption of the rDNA transcription can be done using either 128 Actinomycin D that intercalates into DNA at the actively transcribed rDNA sites and inhibits 129 pre-rRNA chain elongation (Schöfer et al., 1996) or CX-5461 that specifically inhibits Pol I 130 transcription of rDNA genes by selectively targeting the SL1 transcription factor (Drygin et 131 al., 2011). Studies have shown that these inhibitors are able to affect nucleolar structure 132 organization, reviewed in (Grummt, 2013; Mangan et al., 2017; Potapova and Gerton, 2019).

With the use of these specific inhibitors, we will address the organization of rDNA in time and space with regards to major satellite sequences during early embryonic development by 3D-FISH. We will also analyze expression of the various rRNA transcripts to decipher the causal relationship between the spatial organization and the expression changes of these sequences.

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139 **Results**

140 Ribosomal genes (rDNA) 3D organization is linked to their transcription status during

141 preimplantation development

To determine the spatial distribution of rDNA repeats concomitantly with major satellite sequences in mouse early embryo nuclei, 3D-DNA-FISH was performed using probes specific to mouse rDNA repeats (Akhmanova et al., 2000; van de Nobelen et al., 2010) and major satellite sequences (Aguirre-Lavin et al., 2012), respectively. Preservation of the 3D structure of the whole embryos allowed us to do in-depth localization of these sequences from the 1-cell to the blastocyst stage (Fig. 1A and 1C, Fig. S1A and S1B).

148 As expected during the 1-cell stage, major satellite sequences progressively formed a 149 ring that surrounded NPBs (Fig. S1A, (Aguirre-Lavin et al., 2012). rDNA signals correspond 150 to large foci (Fig. 1A and Fig. S1A) and are always associated with major satellite sequences 151 irrespective of the origin of the pronucleus (mPN or pPN). At the 2-cell stage, rDNA FISH 152 signals have a small spots shape close to large signal of major satellite sequences and can be 153 divided into two categories: (i) some of them are not associated with NPBs (green arrows in 154 Fig. 1A) and (ii) some of them are embedded within the ring of major satellite sequences 155 surrounding the NPBs (black and white arrows in Fig. S1A). Interestingly, at the late 1-cell 156 and early 2-cell (2-cell E) stages, these rDNA signals are localized at the outer edge of the 157 NPBs (Fig. 1A) as demonstrated by plot profiles of fluorescence intensities for DAPI, major 158 sequences, and rDNA signals across NPBs (mPN, pPN and 2 -cell E in Fig. 1B). At the end of 159 the 2-cell stage (2-cell L) concomitantly with the embryonic genome activation (Zatsepina et 160 al., 2003), rDNA FISH signals change in shape and localization: forming pearl necklace 161 structures (arrowhead in Fig. 1A) juxtaposed to major satellites and extending inside the 162 NPBs (green pick inside NPB2, plot profile at 2-cell L in Fig. 1B).

163 From the early 4-cell to the late 8-cell stage, rDNA signals that were mostly located at 164 the surface of NPBs, lose the pearl necklace shape to form a dispersed cloud of smaller dots, 165 being less and less associated with major satellite sequences (Fig. 1C) and more and more 166 inside the NPBs (as shown by plot profiles drawn across NPBs, Fig. 1D). Finally, the 3D 167 rDNA organization changes one last time, between the 16-cell and the Morula stage: the FISH 168 signals fill the NPBs (16-cell E, 75hphCG in Fig. S1B) and acquire a nucleolus-like structure 169 (Morula) like in differentiated cells (Junera et al., 1995), with no differences between inner 170 cell mass (ICM) and trophectoderm cells (TE) at the Blastocyst stage (Fig. S1B).

171 Quantification of the number of NPBs per stage revealed that it decreased from the 172 late 2-cell to the 16-cell stage (black line in Fig. S1C) and that the remaining NPBs at the 16-173 cell stage are mostly those associated with the rDNA signal (dark blue bars in Fig. S1C). We 174 then analyzed how rDNA organization evolves and defined four types of NPBs (Fig. 2A, 175 upper panel): T1 corresponding to NPBs with a small number of round-shape rDNA spots; T2 176 corresponding to NPBs with larger spots distributed like a pearl necklace, T3 corresponding 177 to NPBs surrounded by a thin cloud of rDNA signals and T4 corresponding to NBPS with a 178 larger cloud of rDNA signals. Analysis of 350 nuclei from 4-cell to 16-cell stages embryos 179 (Fig. 2A lower panel) shows that T1 and T2 NPBs are predominant at the 4-cell stage and 180 decrease significantly up to the late 8-cell stage (p-value < 0.005). On the other hand, T3 181 NPBs are mostly observed in late 4-cell and early 8-cell stages and T4 NPBs number 182 increases significantly from 8-cell to 16-cell stage (p-value < 0.005, Mann-Whitney U test).

To complete our analysis of rDNA 3D-organization, Imaris software (version 9.6, Oxford Instruments) was used to measure the volume of rDNA FISH signals in whole embryos from the 2-cell to 16-cell stage (Fig. S1D) and the corresponding nuclear volume (stained with DAPI, Fig. S1E). To correct for the potential variation of the nuclear volumes (Fig. S1E), the total rDNA volume in a nucleus was divided by the volume of this nucleus

188 (Fig. 2B). While we did not observe variations at the 2-cell stage, earlier stages have always 189 significantly lower normalized mean volumes than later ones at 4 and 8-cell stages (Fig 2B, 190 S2D) potentially linked with the cell cycle since the chromatin is condensed into 191 chromosomes during mitosis. The late 4-cell, early, and late 8-cell stage displayed larger 192 rDNA volumes than earlier stages, suggesting an important decondensation of the rDNA 193 sequences as visualized on DNA-FISH images (Fig. 1B, lower panel). Remarkably, such 194 decondensation is confirmed by calculating the mean sphericity of rDNA spots (in a given 195 nucleus) at each stage (Fig. 2C). Indeed, 4-cell and 8-cell stage embryos have significantly 196 lower sphericity mean values (from 0.77to 0.82, p-value <0.005) when compared to those of 197 2-cell and 16-cell stage embryos (0.88 and 0.85), respectively. With regards to these 198 parameters (rDNA volume and sphericity), the 16-cell stage has to be considered apart as the 199 structure of the signal changes dramatically at this stage when compared to previous stages 200 and because nuclei become asynchronous, and early vs. late stage cannot be distinguished 201 anymore.

Altogether, our results show that rRNA sequences change in shape and distribution twice during early development: first between the early and the late 2-cell stage, when rDNA becomes transcriptionally active, and, secondly, between the late 8-cell stage and the 16-cell stage on the commencement of cell differentiation.

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207 *rRNA transcription during early development: a two-step dynamic*

To investigate the link between 3D organization of rRNA genes and their transcriptional status, localization of probes specific to the classical rRNA (18S and 28S) and to several pre-rRNA regions that are not retained after complete rRNA processing (5'ETS, ITS1 and ITS2, detailed in Materials and Methods section and Table S1) were studied using RNA-FISH on 3D-preserved embryos (Fig. 3A and 3B).

213 No RNA-FISH signal were detected in the nucleus of 1-cell (data not shown) and early 214 2-cell stages (36hphCG, Fig 3A, and Fig S2A and B) irrespective of the probe used. At the 215 late 2-cell stage, the 5'ETS signal (corresponding to the nascent transcripts) is located in 216 DAPI-free regions inside the nuclei (arrowhead in Fig 3A, 2-cell embryo at 45hphCG) and at 217 the periphery of bigger NPBs (arrow in Fig 3A). At the 4-cell and 8-cell stages, its 218 localization shifts from the periphery to the inner part of the NPBs. Finally, at the 16-cell 219 stage (75hphCG, Fig 3A), the 5'ETS signal fills the NPBs (Fig 3A - lower panel). As for the 220 5'ETS probe, the RNA-FISH signal of ITS1 and ITS2 tend to gain space inside the NPBs 221 during progression through early embryonic development (Fig S2 A and B). At the 16-cell 222 stage, ITS2 and 28S as well as ITS1 and 18S are completely intermingled (Fig S2A and B).

To complete RNA-FISH data with quantitative data, real-time quantitative PCR (RTqPCR) using specific pre-rRNA primers (Table S2) were performed. The level of nascent rRNA transcripts (assessed with 47S and 5'ETS specific primers), gradually increased from late 2-cell to early 8 cell stage and then the curve tends to a plateau (Fig 3C, upper panel). ITS1 and ITS2 specific primers allow us to assess the amount of 'in-process' pre-rRNAs: a significant increase was observed from the late 4-cell to the 8-cell stage (Fig 3C, lower panel).

229 Finally, to gain insight into a putative correlation between epigenetic marks and 230 transcription of rDNA, we have determined using an immuno-RNA-FISH approach, the 231 distribution of H3K9ac and H3K4me3 together with their co-localization with nascent 232 transcripts (5'ETS probes). These two histone post-translational modifications (PTMs) are 233 canonically associated with actively transcribed genes (Allis and Jenuwein, 2016). From the 234 2-cell to the 8-cell stages, we observed that some H3K9ac bright foci (green arrows in Fig 235 3D) are localized close to 5'ETS RNA-FISH signals (white arrows in Fig 3D). Notably, 236 H3K9ac foci are found at the periphery of 5'ETS signal at 2-cell and 4-cell stages and then dispersed in the 5'ETS cloud inside the NPBs at the 8-cell stage but do not co-localized with 237

the totality of the nascent transcripts (Fig 3D). The same results are found for H3K4me3(green arrows in Fig S2C).

240 In conclusion, in contrast to the nucleolus in the differentiated cells, rDNA 241 transcription and rRNA maturation processes are separated in time and space in the immature 242 nucleolus of the embryos at early stages. The first part of the early development (from the 243 late-2-cell to the 4-cell stage) is mainly dedicated to the initiation of rDNA transcription as 244 revealed by a large number of nascent transcripts and the presence of H3K9ac and H3K4me3 245 localized close to nascent rRNA. The second part of the early development (8-cell and 16-cell 246 stages) is characterized by the maintenance of the initiation of transcription but an increase in 247 the processing of immature pre-rRNAs.

248

Inhibition of rRNA transcription and its consequences: elongation is impaired not initiation

251 To further investigate this temporal separation of initiation of the transcription and 252 processing of rRNA, we used two inhibitors of the RNA Polymerase I (RNA Pol I). Embryos 253 at the 1-cell stage were cultured with either CX-5461 (1µM) or Actinomycin D (ActD, 254 7.5ng/µL) and fixed 24h (2-cell stage) or 48h later (4-cell stage). We confirmed that both CX-255 5461-treated and ActD-treated embryos arrested their development at the 4-cell stage (Koné 256 et al., 2016). We then localized the nascent rRNA transcripts by immunoRNA-FISH with 257 5'ETS probe as well as two key nucleolar components, UBF and Nopp140 involved in the 258 initiation of rRNA transcription and processing of the pre-RNA, respectively (Koné et al., 259 2016). In both the treated groups, 5'ETS signal (green arrows in Fig 4A), Nopp140 (red 260 arrows in Fig 4A), and UBF (white arrows in Fig 4A) immunoRNA-FISH patterns are altered 261 compared to the control-untreated group (Fig 4A). As described in Koné et al (2016), in the control group of UBF and 5'ETS, signals are superposed and surrounded by Nopp140 signal 262

and in CX-5461-treated embryos, Nopp140 and UBF form nucleolar caps, reviewed in (van
Sluis and McStay, 2017) with 5'ETS signals still localized around NPBs. On the contrary,
after 48h of treatment, ActD-treated embryos show a higher disruption than CX-5461-treated
embryos with complete segregation between the UBF-5'ETS and Nopp140 signals, suggesting
that nascent transcripts are not processed anymore.

268 To test this hypothesis, the number of nascent pre-rRNAs (using primers for 47S-pre 269 rRNA and the 5'ETS regions) and the number of pre-rRNAs in the process (using primers 270 specific to the ITS1 and the ITS2 regions) were assessed by RT-qPCR (Fig 4B and Fig 4C). 271 There was no significant difference between control and treated-embryos at the 2-cell stage. 272 In the CX-5461 condition, the quantity of 47S, 5'ETS, and ITS1 pre-rRNAs were significantly 273 increased (1.8 fold change) between control and the treated 4-cell embryos (Fig 4B, p value< 274 0.01) although we observed a slight decrease in the RNA-FISH signal compared to control 275 (5'ETS FISH signals in Fig 4A, ITS1 and ITS2 FISH signals in Fig 4D). A putative 276 explanation is that CX-5461-treated embryos compensate a lower number of rDNA loci 277 engaged in transcription by a higher rate of transcription on rDNA that was able to initiate 278 transcription by recruitment of SL1 and UBF. On the contrary, we observed a significant (p-279 value <0.005) decrease of 5'ETS, ITS1, and ITS2 pre-rRNAs in ActD-treated embryos both 280 by RT-qPCR quantification (Fig 4C) and by RNA-FISH (Fig 4A and 4E). In agreement with 281 the hypothesis that Actinomycin D inhibits RNA Pol I elongation, ITS1 and ITS2 display a 282 stronger reduction (Fig4 E) than 5'ETS (Fig 4A, left panel). CX-5461 has no specific effect on 283 rRNA processing compared to Actinomycin D (in congruance with Mars et al 2020).

Altogether, these results demonstrate that RNA Pol I inhibition with Actinomycin D treatment during early embryonic development induces transcriptional arrest at the 4-cell stage and a massive reorganization of nucleolar proteins due to an incomplete transcription and processing of the long 47S pre-rRNA.

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289 RNA Pol I inhibition induces changes on the 3D organization of rDNA

290 To assess the consequences of RNA Pol I inhibition on rDNA organization and the 291 nearby major satellite sequences, DNA-FISH approach was used (Fig. 1). To compare the 292 organization of rDNA and major satellite sequences between the control and treated embryos, 293 the shape of both DNA signals was visualized and analyzed using 3D reconstruction at the 2-294 cell and 4-cell stages (Fig.5A and 5B) respectively. At the 2-cell stage, the rDNA signals are 295 compacted and juxtaposed to major satellite signals both in the control and treated embryos 296 (green and black arrows in Fig 5A) but their position in regards to NPBs is different between 297 the control (inside NPBs, Ctrl in Fig 5C) and the treated (outside NPBs, CX-5461 and ActD 298 in Fig 5C) embryos, as shown in plot profiles. At the 4-cell stage, the rDNA signal in the 299 treated embryos is less intense than in the control (green arrows in the upper panel, Fig 5B), 300 its spatial distribution is more compact (clusters seem bigger and in lower number) and in 301 ActD-treated embryos, it forms clusters as in 2-cell stage embryos. Plot profiles drawn across 302 the nuclei show that in the CX-5461, rDNA are still inside the NPBs as in the control 303 condition, but not in the ActD condition where rDNA are outside the NPBs, juxtaposed to 304 major satellite sequences (Fig 5D).

305 To assess and quantify the differences in terms of compaction and shape between the 306 control and the treated embryos, rDNA FISH signals were segmented using Imaris software 307 (as previously in Fig 2B and 2C) to measure the volume and the sphericity of the rDNA signal 308 at 2-cell and 4-cell stages (Fig 6A and 6B). In the controlled condition, the mean volume of 309 the rDNA signal increases from the 2-cell to 4-cell stage, and its shape (measured by the 310 sphericity) becomes less round meaning that rDNA sequences decondensed concomitantly 311 with being actively transcribed. At the 2-cell stage, the shape of the rDNA signal is 312 significantly different between the control and the treated embryos (left panel in Fig 6B, p-

value <0.005) suggesting that ribosomal sequences are less round in the treated embryos and
putatively less compact. At the 4-cell stage, the mean volume of rDNA signal is significantly
(p-value <0.005) different between the control and the treated embryos (Fig 6B, 5.31 +/- 1.63
in Ctrl vs 0.85+/-0.48 and 0.93+/- 0.27 in CX-5461 and ActD treated embryos, respectively)
meaning that rDNA sequences stay tightly compact and clustered upon RNA Pol I inhibition.

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319 Disruption of 3D organization of rDNA with RNA Pol I inhibition has consequences on the 320 transcription of pericentromeric sequences

321 By paying attention to the major satellite signals (shape and volume) assisted us in 322 pinpointing that their distribution is modified by the drug treatment. Indeed, after 48h of RNA 323 Pol I inhibition, major satellite DNA-FISH signals show a croissant shape- as in 2-cell 324 embryos (red and black arrows in Fig 5B)- whereas they are normally clustered into 325 chromocenters at the 4-cell stage in controls (black arrowhead in Fig 5B). To confirm this 326 further, the mean volume and the sphericity of the DNA-FISH signals of major satellite 327 sequences were measured using Imaris in the control and the treated embryos (Fig S3A and 328 S3B and Fig 6C and 6D). The mean volume of major satellite sequences FISH signals is 329 significantly lower (p-value <0.005) at the 2-cell stage in the treated embryos compared to the 330 control (right panel, Fig 6C). At the 4cell stage, the volume in the control decreases, while 331 there is no change in the treated embryos (left panel, Fig 6C). Furthermore, the decreased 332 sphericity in the treated embryos compared to the control at the 4-cell stage (right panel, Fig 333 6D) means that the major satellite sequences are less round and compact in the treated 334 embryos. Similarly, the mean sphericity of the 4-cell stage in the treated embryos is not 335 significantly different from that of the 2-cell control embryos.

336 Previous studies highlighted that major satellite sequences are specifically transcribed
337 at the 2-cell stage when their shape is less round (Casanova et al., 2013; Probst et al., 2010),

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338 therefore we decided to assess major satellite transcription in the treated vs. control embryos 339 by RNA-FISH and RT-qPCR. No difference in the major satellite RNAs was detected 340 between the control and treated embryos at the 2-cell stage, neither by RNA-FISH nor by RT-341 qPCR (Fig 7A, B and C) however we detected a significant decrease of transcription in the 4-342 cell stage ActD-treated embryos (Fig 7B, p < 0.005). On the other hand, a housekeeping gene 343 such as Hprt (Fig S3C) does not vary in ActD-treated embryos suggesting a reciprocal 344 influence between rDNA and major satellite transcription. As a matter of fact, after the 2-cell 345 stage burst of transcription, the remaining major satellite RNA-FISH spots were always 346 visualized inside the cloud of 5'ETS FISH signal (green arrows in Fig 7D), at the 4-cell, 8cell, and 16-cell stages (red arrow in Fig 7D) suggesting that rDNA transcription might 347 348 induce a chromatin environment that favors transcription of major satellite sequences.

349

350 **Discussion**

351 Spatial organization of rDNA during early development

352 In this study, we demonstrated that the spatial positioning of ribosomal genes (rDNA) 353 correlates with their transcription activity. In the context of early embryonic development, 354 ribosomal repeat sequences are first clustered at the surface of NPBs, flanked by ring-shaped 355 major satellite sequences (1-cell and early 2-cell). From the late 2-cell to the 16-cell/morula 356 stage, the sequences then expanded in the inner space of the NPBs until complete filling. 357 During this period, rDNA gradually separated spatially from major satellite sequences 358 although they are physically attached on the same chromosome. The cloudy organization of 359 the active ribosomal sequences we observed fits with the localization of UBF as previously 360 described (Fulka and Langerova, 2014; Hamdane et al., 2016; Koné et al., 2016) and is 361 specific to the early embryonic period. This unique nucleolus organization is very different from the somatic ones (Maiser et al., 2020) that can only be found at the morula and 362

blastocyst stages (Lavrentyeva et al., 2015). Interestingly, this second switch in the nucleolus
conformation is associated with a second burst of embryonic genes transcription at the
morula-to-blastocyst transition described in transcriptomic data (Hamatani et al., 2004;
Hamatani et al., 2006; Wang et al., 2004).

367 In parallel to these drastic changes in the spatial organization of the sequences, we 368 show that rRNA synthesis begins at the mid-2-cell stage, first in the small spherical structures 369 outside the NPBs. Then, nascent rRNAs later appear at the surface of some NPBs as 370 previously described (Koné et al., 2016; Lavrentyeva et al., 2015). Therefore, at very early 371 stages (1-cell and 2-cell stages), NPBs can not be considered as a reservoir for pre-rRNAs 372 waiting to be cleaved confirming previous studies (Lavrentyeva et al., 2015; Shishova et al., 373 2015a; Shishova et al., 2015b). Interestingly, immuno-RNA-FISH revealed that bright spots 374 of H3K9ac and H3K4me3 are located at the vicinity of 5'ETS signal at the 2-cell stage and 375 that these spots are found inside the 5'ETS cloud at later stages. These results are in 376 agreement with ChIP data obtained on mouse embryonic fibroblast or 3T3 cells (Herdman et 377 al., 2017; Zentner et al., 2014) describing an enrichment of these two epigenetic marks on the 378 spacer promoter (UCE core) at the actively transcribed rDNA repeat (Grummt and Längst, 379 2013; Moss et al., 2019).

380

381 Inhibition of RNA Pol I and its impact on rRNA processing

To decipher the link between the spatial organization of rDNA and their transcriptional activity, we used two drugs that inhibit RNA Pol I activity in two different mechanisms. CX-5461 is a molecule that inhibits the initiation of transcription by impairing the formation of the pre-initiation complex (PIC) (Drygin et al., 2011) whereas Actinomycin D is a DNA intercalant that inhibits elongation by the RNA polymerases (all of them) in a dose-dependent manner (Perry and Kelley, 1970). Several recent studies (Bruno et al., 2020;

Mars et al., 2020; Sanij et al., 2020) discussed the specific targets of CX-5461 (SL1 or Topoisomerase II or RRN3) and its mode of action (G-quadruplex stabilization, DNA damage, impairing of DNA replication fork). Notably, Mars et al., (2020) emphasis that RNA Pol I inhibition by CX-5461 is not reversible and can lead to cell death because of DNA damage. To get read of these putative side effects of CX-5461, ActD was also used in this study.

394 In the early embryonic context, we observed that CX-5461 disrupts the classical 395 tripartite (i.e. FC, DFC and Granular component (GC) (Boisvert et al., 2007; Pederson, 2011)) 396 nucleolus organization (this study & Koné et al 2016). In agreement with (Mars et al., 2020), 397 our study demonstrates that CX-5461 has less impact on the elongation process than 398 Actinomycin D. In the treated embryos, Actinomycin D treatment also inhibits nascent rRNA 399 synthesis but induces a complete stop of the long pre-rRNA production (labeled with ITS1 400 and/or ITS2) which in turn induces both dissociation of transcription initiation (labeled by 401 5'ETS and UBF) and processing of rRNA (labeled with Nopp140). It is worth-mentioning that 402 the treated embryos arrested their development at the 4-cell stage most likely because 403 maternally inherited ribosomes are not renewed as demonstrated in the embryos invalidate for 404 UBF (Hamdane et al., 2014; Hamdane et al., 2016).

405

406 The link between major satellite and rDNA sequences transcription

Lastly, we compared rDNA shape, distribution, and transcriptional activity in time and space with regards to major satellite sequences during early development. Indeed, major satellite sequences are drastically reorganized at the time of EGA (Aguirre-Lavin et al., 2012; Probst et al., 2007) and this reorganization is dependent on major satellite transcripts (Casanova et al., 2013; Probst et al., 2010). On the other hand, localization of major satellite sequences around the NPBs is mandatory at the 1-cell stage as demonstrated by enucleolation 413 experiments (Fulka and Langerova, 2014; Kyogoku et al., 2014; Ogushi and Saitou, 2010; 414 Ogushi et al., 2008) and at the early 2-cell stages (Jachowicz et al., 2013) even if in-depth 415 knowledge of the mechanisms involved are still unknown. Growing pieces of evidence 416 propose that NPBs, rather than acting as a precursor of nucleoli, act as a platform or a Velcro 417 for major satellite sequences (Fulka and Aoki, 2016; Padeken and Heun, 2014) thanks to 418 nucleoplasmin 2 (NPM2, (Burns et al., 2003; Inoue and Aoki, 2010; Kyogoku et al., 2014)). 419 In fact injection of sole NPM2 mRNA rescue developmental failure of enucleolated oocytes 420 (Ogushi et al., 2017). Surprisingly, the absence of NPBs induces the earlier formation of 421 chromocenter-like structures at the zygotic stage (Ogushi et al., 2017) although transcription 422 of major satellite sequences is repressed after the end of the 2-cell stage (Fulka and 423 Langerova, 2014). In this study, we show that inhibition of rDNA transcription induces major 424 satellite sequences to change from a chromocenter-like shape to a ring-like shape without 425 increasing their transcription activity. Surprisingly, Actinomycin D treatment induces a 426 decrease of major satellite transcription while other RNA polymarase II-dependent genes are 427 not affected.

428 rDNA spatial organization is disrupted on the inhibition of RNA Pol I and rRNA 429 sequences aggregate, exhibiting a round shape comparable to earlier inactive stages. 430 Similarly, major satellite sequences exhibit a spatial organization that looks like an early 2-431 cell stage. This loss of chromatin organization in RNA Pol I inhibited embryos can partially 432 explain the decrease of major satellite sequences transcription. After the 2-cell stage, major 433 satellite transcription is severely repressed and only one or two spots of major satellite RNA 434 can be detected per nucleus by RNA-FISH (Probst et al., 2010). Remarkably, these spots were 435 often found inside the 5'ETS cloud (this study), suggesting transcription of major satellite 436 sequences after the 2-cell stage may require a suitable environment that favor transcription. A 437 putative explanation -among others- is that transcription of major satellite sequences, in 4-cell

- 438 and 8-cell embryos, need the rDNA peculiar chromatin state (Moss et al., 2019; Potapova and
- 439 Gerton, 2019): nucleosome-free chromatin with UBF that promotes loops formation
- 440 (Stefanovsky et al., 2001).

441

442

443 Material and Methods

444 Ethics

Animal care and handling were carried out according to the national rules on ethics and animal welfare in the Animal facility (IERP, INRAE, Infectiology of fishes and rodent facility, doi: 10.15454/1.5572427140471238E12, Jouy-en-Josas, C78-720). This work was approved by the French Ministry of Higher Education, Research and Innovation (n°15-55) and the local ethical committee (INRAE Jouy-en-Josas Center). Departmental veterinary regulatory services have delivered habilitation to work with laboratory animals to ABG (n°78–184) who supervised the work.

452

453 Collection of mouse embryos and culture

454 Embryos were collected upon superovulation and mating of C57/CBAF1 mice as previously 455 described (Koné et al., 2016). Zygotes were obtained by dissecting the ampulla of the oviduct 456 24 hours post-injection of human chorionic gonadotropin (hCG) and treated briefly with 457 1mg/ml of hyaluronidase (Sigma) in M2 medium (Sigma) to remove follicular cells. Then, 1-458 cell (24h phCG) embryos were cultured in vitro in M16 medium (Sigma) at 37°C in a 459 humidified atmosphere of 5% CO₂ until they reached the adequate stages to be processed: 460 early 2-cell (36h phCG - 2CE), late 2-cell (48h phCG - 2CL), early 4-cell (54h phCG - 4CE), 461 late 4-cell (58h phCG - 4CL), early 8-cell (60h phCG - 8CE), late 8-cell (72h phCG - 8CL), 462 16-cell (75h phCG), Morula (96h phCG) and Blastocyst (110h phCG).

463

464 CX-5461 and Actinomycin D treatment

Two different RNA Polymerase I (RNA Pol I) inhibitors were used in this study: CX-5461 that impairs the formation of the RNA Polymerase I pre-initiation complex (PIC) by disrupting the recruitment of SL1 (Selective factor 1) with UBF (Upstream biding factor) to the rDNA promoter (Drygin et al., 2011) and Actinomycin D that intercalates into the DNA and thereby inhibits elongation of all the RNA polymerases activity, although with a higher affinity for RNA Pol I (Bensaude, 2011; Perry and Kelley, 1970). These two drugs induce segregation of the nucleolus components dedicated either to transcription (fibrillar and dense fibrillar components -FC and DFC) or to rRNA processing (granular component, GC). These components can be visualized by fluorescent immuno-detection of the proteins involved in these processes (UBF and Nopp140 respectively, as described in (Koné et al., 2016)).

Embryos collected at the 1-cell stage (24hphCG) were cultured in M16 medium containing either 1 μ M of CX-5461 (Adooq) or 7.5ng/ μ L of Actinomycin D (Sigma) to inhibit RNA Pol I as described respectively in (Bellier, 1997; Koné et al., 2016). Every 24h, the embryos were transferred into new droplets of medium containing the RNA Pol I inhibitors. The treated embryos were fixed at two-time points: 24h and 48h after exposure to the drug and then processed for immunoRNA-FISH / DNA-FISH or frozen dry in low RNA binding tubes and kept at -80°C for RT-qPCR assays.

482

483 **Probes for DNA and RNA-FISH**

484 rDNA and major satellite FISH probes were prepared as described in (Aguirre-Lavin et al., 485 2012) and (Bonnet-Garnier et al., 2013). Briefly, cosmid containing the entire transcribed 486 sequences (n°13, detailed in (Akhmanova et al., 2000; van de Nobelen et al., 2010) was used 487 for the ribosomal sequences (kind gift from N. Galjart). The amplified rDNA gene fragments 488 were purified using nucleobond AX 100 columns for Miniprep System (Macherey-Nagel, ref 489 740521.100). The rDNA sequences were then labeled with Digoxigenin-11-dUTP by nick 490 translation according to the manufacturer's protocol (Roche, Ref 11277065910). For the 491 detection of major satellites, a probe prepared by PCR on genomic mouse DNA with the primers 5'-CATATTCCAGGTCCTTCAGTGTGC-3' and 5'-CACTTTAGGACGTGAAAT 492

493 ATGGCG-3' was labeled with Cy3 by random priming according to the kit instruction
494 (Invitrogen Kit, Ref 18095–011).

495 Specific oligo-probes (designed from the complete 45S mouse ribosomal sequences -496 X82564.1 or adapted from (Le Bouteiller et al., 2013; Shishova and others, 2011; Shishova et 497 al., 2015b) directly labeled with a fluorophore in 3' end (sequences are detailed in Table S1) 498 were purchased from Eurogentec. These probes located (Fig. 2B) on different regions of the 499 47S pre-RNA (Kent et al., 2008) are named: 5'ETS for the probe specific to the external 500 transcribed spacer (rapidly removed, allowing the detection of early transcription sites, i.e. 501 nascent rRNA transcripts), ITS1: specific for the internal transcribed spacer 1 located between 502 the 18S and the 5.8S regions and ITS2: specific for the internal transcribed spacer 2 located 503 between the 5.8S and the 28S region. ITS1 and ITS2 enable the mapping of the region where 504 rRNA processing occurs. Probes specific for 18S and 28S regions were also used to assess if 505 mature or pre-rRNAs were already accumulated in NPBs before the onset of rDNA 506 transcription. Of note, the 18S and 28S probes signal can be detected in the cytoplasm 507 because they are components of ribosomes.

508

509 DNA-FISH

510 For DNA-FISH, DNA probes described above specific to either ribosomal or major satellite 511 sequences were used together on the whole embryos. During all the manipulations, drying 512 was avoided to preserve the 3D of nuclei. This protocol is based on (Miyanari and Torres-513 Padilla, 2012) with slight modifications. The embryos collected at a specific stage were 514 transferred in a 10-well glass dish warm at 37°C. The embryos were rinsed in M2 medium, 515 submitted briefly (less than 15s) to Tyrode's acid solution (Sigma-Aldrich), washed quickly in 516 an M2 with 10mM phenylmethanesulfonyl fluoride (PMSF, Fluka) solution then in a 0.5% 517 polyvinyl-pyrrolidone (PVP) in PBS with 10mM PMSF solution and finally incubated in a 518 fixation/permeabilization solution (4% paraformaldehyde (PFA), 0.5% Triton X-100, 10mM 519 PMSF, 0.5% PVP in PBS) for 15min at 37°C. The embryos were then washed twice in a 520 PBS/PVP 0.5% solution at RT and let at +4°C (at least overnight) in a humid chamber until 521 fixation of other stages. To remove the zona pellucida (ZP), all the embryos were incubated 522 under a stereomicroscope to monitor the ZP vanishing in a 0.1N HCl (Prolabo) solution for 20 523 to 30s (depending on the stage) at RT and transferred to PBS/PVP 0.5%. The ZP removal was 524 achieved by successive passages through a very fine glass pipette when needed. The embryos 525 were permeabilized in a 0.5% Triton X-100 / 200µg/ml RNase solution for 30 min at 37°C 526 and washed twice in 2X saline-sodium citrate (2XSSC) pH 6.3 for 5 min at 37°C. The embryos were then transferred for 3h at 37°C in a humidified chamber in a 20µl drop of 527 528 hybridization buffer (50% formamide, SCC 2X, Denhardt 1X, 40 mM NaH2PO4, 10% 529 dextran sulfate) containing the DNA-FISH probes mix (14µl of rDNA probe solution at 530 50ng/µl and 1µl of major satellite probe solution at 100 ng/µl completed to 20µl with 531 hybridization buffer). The embryos were denatured at 85°C for 10 min on a heating plate and 532 then let three days for hybridization at 37°C in a humid chamber. After hybridization, the 533 embryos were rinsed twice with 2X SSC / 0.1% Triton X-100 / 0.5% PVP for 10 min at 42°C. 534 Embryos were then incubated in a blocking solution (4X SSC / 2% BSA) for 1h at RT and 535 incubated with a primary antibody diluted in blocking solution (sheep anti-digoxigenin, 1/200, 536 Roche) overnight at $+4^{\circ}$ C in a dark humidified chamber. The embryos were washed twice in 537 4X SSC / 2% BSA / 0.05% Tween20 before and after incubation with the secondary antibody 538 (mouse anti-sheep Ig coupled to FITC, 1/200, 713-095-147, Jackson ImmunoResearch 539 Laboratories Inc., USA) for 1h at RT. The embryos were washed in 2X SSC / 0.5% PVP and 540 gently mounted to preserve the 3D structure of the nuclei between slides (Menzel Superfrost 541 Plus, Thermo Scientific) and cover-slip with a large amount of Vectashield antifading agent 542 (Abcys) containing 10µg/mL DAPI (Invitrogen).

543

544 **RNA-FISH**

545 For RNA-FISH, the embryos were processed almost as for DNA-FISH with two main 546 differences: 1) the ZP was not removed by HCl treatment; 2) an RNAse inhibitor (1µL/mL of 547 RNasin - Promega) was added to all the solutions after the fixation/permeabilization step. 548 Briefly, after fixation/permeabilization, embryos were further permeabilized in 0.5% Triton X100 /0.5% PVP-PBS for 45min at 37°C, washed and transferred for 30min at 50°C in the 549 550 pre-hybridization mix containing 50% Formamide (Sigma-Aldrich); 0.5µg/µL tRNA (Sigma-551 Aldrich); 1X hybridization buffer (2X hybridization buffer was prepared beforehand with 20% 552 dextran sulfate, 4X SSC; 1mM EDTA, 40mg/mL BSA, 2mg/mL PVP, 0.1% Triton X100, 553 diluted in pure grade water). In parallel, a solution containing the fluorescent oligonucleotide 554 probes (100ng/µl diluted in 1X hybridization buffer) was denatured 10 min at 85°C and 555 immediately put on ice to avoid renaturation of the DNA. The embryos were then transferred 556 in this hybridization mix and incubated overnight at 42°C. The embryos were washed twice 557 in 2XSSC; 0.5%PVP; 0.1% Triton X100 (diluted with pure grade water) for 10min at 37°C 558 and mounted on slides with coverslips as described in the DNA-FISH section.

559

560 ImmunoRNA-FISH

Immunofluorescence labeling followed by RNA-FISH was performed as described in Kone et al. (2016) with slight modifications. Briefly, the embryos were fixed as described in the RNA-FISH section before immunostaining. The embryos were then permeabilized in 0.5% Triton X100 / 0.5% PVP-PBS for 30min at 37°C and transferred in 2% BSA-PBS 1h at RT. The embryos were incubated with primary antibody overnight at 4°C, washed thrice in 0.5% PVP-PBS, and then incubated with secondary antibody for 1h and post-fixed in 2% PFA/0.5% PVP-PBS for 10 min at RT. The embryos were then further processed for RNA-FISH as

described in the RNA-FISH section. All antibodies were diluted in 2% BSA-PBS. Again, RNAse inhibitor (1 μ L/mL of RNasin - Promega) was added to all the solutions after the fixation/permeabilization step.

571

572 Antibodies

Primary antibodies used: anti-UBF (1/100; mouse polyclonal antibody H00007343-M01;
Novus Biologicals), anti-Nopp140 (1/150; rabbit polyclonal antibody, RF12 serum; a gift
from U. Thomas Meier, Department of Anatomy and Structural Biology, New York, USA),
anti-H3K9ac (1/100, rabbit polyclonal antibody, 39917) and anti-H3K4me3 (1/400, rabbit
polyclonal antibody, ab8550, Abcam). Secondary antibodies used: IgG donkey anti-mouse
Cy3 (1/200, 715-025-151, Jackson ImmunoResarch) and IgG donkey anti-rabbit Cy5 (1/200,
111-175-152, Jackson ImmunoResarch).

580

581 Confocal microscopy and image analyses

Imaging was performed with a ZEISS LSM 700 confocal laser scanning microscope (MIMA2, INRAE, Microscopy and Imaging Facility for Microbes, Animals and Foods, https://doi.org/10.15454/1.5572348210007727E12) equipped with a 63X (1.4 NA) oil immersion objective. Z-stacks were acquired with a frame size of 512×512 or 1024×1024 , a pixel depth of 8 bits, and a z-distance of 0.37 µm between optical sections. Fluorescence wavelengths of 405, 488, 555, and 639 nm were used to excite DAPI, Alexa-488, or FITC, Cy3, and Cy5, respectively.

589 Fluorescent profiles measurement were generated in Fiji (Schindelin et al., 2012; Schindelin 590 et al., 2015; Schneider et al., 2012) and 3D reconstructions using AMIRA 3.1 software 591 (Mercury Computer Systems, Berlin, Germany, (Stalling et al.) 592 Quantitative image analysis was performed manually using Imaris 9.6 software (Oxford 593 Instruments) available at the MIMA2 ISC INRAE. Briefly, in 10 embryos of each stage, 594 nuclei were first segmented using an automatically set threshold (wizard function in Imaris 595 surfaces package). Major satellites and rDNA signals were then segmented in a specified 596 region of interest (ROI defined by the nucleus label) following the same process. For all 597 detected objects, several values (area, volume, sphericity, numbers) were calculated by Imaris 598 and exported for statistical analysis in Microsoft Excel file format. For each analyzed nuclei, 599 the sum of the volume of DNA-FISH signals was calculated to obtain the total volume of 600 rDNA or major satellite DNA-FISH signals in a given nucleus. Then this total volume was 601 normalized by the volume of the given nucleus using this formula: (Vol_tot_{FISH} / 602 Vol_tot_{nucleus})x 100, to circumvent any nucleus size effect when comparing between stages.

603

604 **RT-qPCR**

605 The embryos at each stage were pooled by a batch of 10 and snap-frozen. Total RNA was 606 prepared without purification using the SingleShot Cell Lysis kit (Biorad) that includes 607 DNase treatment. cDNA was prepared directly from the lysates with random hexamers (300 608 ng) (Lifetechnologies, France) following the supplier's protocol (25°C for 5 min, 50°C for 60 609 min and 70°C for 15 min) in the presence of 10 fg of Luciferase RNA (Promega) using the 610 SuperSript III reverse transcription kit (Invitrogen). Negative controls without reverse 611 transcriptase were prepared the same way. RT-qPCR was performed on 0.2 or 0.032 embryo 612 equivalent (as described in Salvaing et al 2016), depending on the genes, using KAPA SYBR 613 FAST (Roche) or SYBR Green (Applied, for Major satellites) master mix on a StepOne Plus 614 cycling machine (Applied). The difference in amplification of positive and negative samples 615 was at least 5 Ct (5 to 9). To avoid biais due to a highly variable amount of transcript from 616 endogenous genes (Hprt1) before the EGA, the results were finally normalized to Luciferase 617 (as described in Bui et al). Primers are listed in Table S2. Three to six biological replicates618 were used.

619

620 Statistical analyses

All the statistical analyses and tests were performed using R packages. Several R packages such as ggplot2 and lattice were used in complement to Rcmdr packages to generate graphs. The normality and homogeneity of variances were tested using the 'shapiro.test' and 'bartlett.test' R-packages, respectively. As the sample size was small (<30), non-parametrical tests were used. Two by two comparison were done using 'wilcox' packge (Mann-Whitney-Wilcoxon) and comparison of the two consecutive stages were done with 'nparcomp' package (Nonparametric Multiple Comparisons).

628

629 Supplementary material

630 Figs. S1–S5 show 3D organization of ribosomal sequences (rDNA) at 1-cell stage, 16-cell and 631 morula to blastocyst stages using DNA-FISH, the number of nucleolar precursor bodies 632 (NPBs), Quantification of the volume of nucleus and rDNA FISH signals using Imaris, 633 Localization by RNA-FISH of several ribosomal RNA (rRNA) from 2-cell to 16-cell stages, 634 Localization of H3K4me3 and nacent transcripts of rRNA by immunoRNA-FISH, 635 Quantification of major satellite sequences volume and sphericity from 2-cell up to 16-cell 636 stage and relative expression of an housekeeping gene (Hprt) by RT-qPCR in control and 637 treated embryos.

Tables S1–S2 show the list of oligoprobes used for the RNA-FISH and the list of primersused in th RT-qPCR experiments respectively.

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657

658 Authors contribution

659 M. Chebrout set up the RNA-FISH and immunoRNA-FISH experiments, performed majority 660 of the experiments, and analyzed the data. M.C. Koné, M. Cournut and H.U Jan performed 661 and analyzed DNA-FISH experiments. R. Fleurot and T. Aguirre-Lavin set up and performed 662 the DNA-FISH experiments. A. Jouneau and N. Peynot performed and analyzed the RTqPCR experiments. N. Beaujean secured funding, gave guidance during initial data 663 664 production and analysis, and made early corrections to the manuscript. A. Bonnet-Garnier 665 supervised, directed, and designed the study, analyzed, and interpreted the data, made the 666 figures, and wrote the article. All the authors reviewed and commented on the manuscript.

667

668 Ethical approval

All applicable international, national, and/or institutional guidelines for the care and use of

animals were followed. This article does not contain any studies with human participants

671 performed by any of the authors.

672

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- 889
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891 **Figures Legend:**

892

893 Figure 1: 3D organization of ribosomal sequences (rDNA) is correlated with their

894 transcriptional state

895 (A) 3D organization of rDNA and major satellite sequences at 1-cell and 2-cell stages 896 detected by 3D DNA-FISH using specific probes. Upper panel : single z-section of a 897 representative nucleus of 1-cell (24hphCG), early (E, 36hphCG) and late (L, 48hphCG) 2-cell 898 stages. Lower panel: Amira 3D representation of each nucleus. White (or black, lower panel) 899 arrows indicate clustered rDNA FISH signal juxtaposed to major satellite signal at NPBs 900 surface; green arrows indicated rDNA juxtaposed to round shape major satellite signal and 901 white (or black, lower panel) arrowhead indicate pearl necklace rDNA signal at NPBs 902 surface. (B) Localization of rDNA and major satellite signals in regards to NPBs boundaries 903 at 1-cell and 2-cell stages. Left panel: single z-section of the nucleus with the line 904 corresponding to the Intensity profile measurement in Fiji. Right panel: Fluorescence intensity 905 plot profile along the line across multiple channels (blue dot line for DAPI, the green line for 906 rDNA signal, and red line for major satellite signal). (C) 3D organization of rRNA genes 907 from 4-cell to 8-cell stages detected by 3D DNA-FISH. rDNA FISH signal expands in the 908 inner of the NPBs up to fulfil it completely. Upper panel: z-section of a representative nucleus 909 of early (E) or late (L) 4-cell and 8-cell stage. Lower panel: Amira 3D representation of each 910 nucleus. (D) Localization of rDNA and major satellite signals in regards to NPBs boundaries 911 at 4-cell and 8-cell stages. Left panel: single z-section of the nucleus with the line 912 corresponding to the Intensity profile measurement in Fiji. Right panel: Fluorescence intensity 913 plot profile along the line across multiple channels (blue dot line for DAPI, the green line for 914 rDNA signal, and red line for major satellite signal).DNA is in blue or grey, rDNA in green, 915 and major satellite sequences in magenta. Scale bar = $5\mu m$. mPN: maternal Pronucleus; pPN : 916 paternal Pronucleus and NPB: Nucleolar Precursor Body; h phCG : hours post-injection of 917 human Chorionic Gonadotrophin, major: mouse major satellite sequences, rDNA: ribosomal 918 DNA.

919

920 Figure 2: Features of rDNA spatial organization during pre-implantation development

921 (A) Spatial organization rDNA surrounding the NPBs. <u>Upper panel:</u> Four kinds of NPBs can
922 be defined depending on their rDNA pattern: Type 1 (T1) displays small dots of rDNA signal,
923 Type 2 (T2) shows spots distributed like a pearl necklace, Type 3 (T3) harbors a thin cloud of
924 rDNA signal and Type 4 (T4) a large cloud of rDNA signal. Stars indicated the NPBs

925 representative of each type. Scale bar = $5\mu m$. Lower panel: Quantification of the number of 926 each type of rDNA/NPBs pattern per stage at the 4-cell, 8-cell (early (E) and late (L)) and the 927 early 16-cell stage. Mann-Whitney test was used to compare (two by two) the distribution of 928 the type between stages. **, p-value< 0.01; ***, p-value < 0.001. (**B** and **C**) Quantification of 929 rDNA total volume (B) and average sphericity (C) per nucleus and stage based on the DNA-930 FISH signal of the ribosomal sequences. All measurements were done using Imaris 9.6 931 (Oxford Instruments) from the 2-cell to 16-cell stage. Statistical significance was evaluated 932 using a non-parametric multiple comparison test (nparcomp, R package). ***, p-value < 933 0.001. The number of nuclei examined per stage is indicated below the stage name.

934

935 Figure 3: rRNA transcripts organization upon early development

936 (A) rDNA transcripts (green) are visualized by RNA-FISH using a specific probe for the 937 5'ETS region. Upper panel: representative nuclei of 2-cell (36h and 45hphCG), 4-cell 938 (60hphCG), 8-cell (72hphCG) and 16-cell (75hphCG) stages embryos. Arrow indicates 939 nascent rRNA labeled with a 5'ETS probe at the surface of an NPB and arrowhead indicates 940 nascent rRNA outside NPB. Lower panel: 3D representation of a whole embryo at each stage 941 using Imaris 9.6 for channel segmentation. The hours indicated are "post-injection of hCG". 942 DNA is labeled in blue and rDNA transcripts in green. Scale bar = $5\mu m$. (B) Schematic 943 representation of the rDNA sequence and position of the rRNA FISH probes used in this 944 study. Each repeat of the mouse ribosomal sequence (X 82564.1) is transcribed in a long 945 mRNA (47S) encoding for 18S, 5.8S, and 28S rRNAs, separated by two external transcribed 946 spacers (5'ETS and 3'ETS) and two internal transcribed spacer (ITS1 and ITS2). (C) 947 Ouantification of the rRNA transcription at 2-cell (early (E) and late (L)), 4-cell (e and L), 8-948 cell (E and L), and 16 cell stages by Real-Time qPCR (RT-qPCR) using primers (described in 949 Table S2) specific of 47S long pre-rRNA, of 5'ETS, ITS1 and ITS2 regions. Four to six 950 biological replicates were done by stage, the expression of these pre-RNA was normalized to 951 a fixed amount of luciferase mRNA added before the reverse transcription. Statistical 952 significance was evaluated using a non-parametric multiple comparison test (nparcomp, R 953 package). ***, p-value < 0.001. (D) Localization of H3K9ac (green) and nascent rRNA 954 (magenta) by immuno-RNA-FISH from 2-cell to 8-cell stage. Green arrows indicate H3K9ac 955 bright and big foci close or inside the 5'ETS RNA-FISH signal. Pixels (point by white arrows) that correspond to co-localization between H3K9ac and 5'ETS fluorescent channels were 956 957 identified using Imaris 9.6 software.

958

959 Figure 4: Inhibition of rRNA transcription by CX-5461 and ActD.

960 (A) Localization using immuno RNA-FISH of nascent rRNA (5'ETS probe in green), 961 Nopp140 (red) and UBF (grey) at 4-cell stage in non treated (Ctrl) and treated embryos (CX-962 5461 and ActD), nuclei were counterstained with DAPI (blue). Every single z-section of a 963 confocal stack image show representative nuclei. Green arrows point 5'ETS FISH signals, red 964 arrows Nopp140 immunostaining, and grey arrows UBF immunostaining. Scale bar = $5\mu m$. 965 (B and C) Quantification by RT-qPCR of the relative expression of rRNAs amplified with 966 specific primers for 47S, 5'ETS, ITS1, and ITS2 in control embryos (2-cell, 4-cell and 8 cell 967 stages, green spots) and treated embryos (2-cell and 4-cell stages): with CX-5461 (B, orange 968 spots) and actinomycin D (C, purple spots). Statistical significance between control and 969 treated embryos was evaluated using the Mann-Whitney test (wilcox, R-package). *, p-value 970 < 0.05, **, p-value < 0.01. (D and E) Representative confocal images of 4-cell stages nucleus 971 in non-treated (Ctrl) and treated embryos with CX-5461 (D) or Actinomycin D (E). In process 972 pre-rRNAs species were localized by RNA-FISH with specific fluorescent oligoprobes for 973 ITS1 (red) - 18S (green), left panel or ITS2 (red) - 28S (green), right panel. Nuclei were 974 counterstained with DAPI (blue). Scale bar = $5\mu m$.

975

976 Figure 5: Consequences of RNA Pol I Inhibition on the spatial organization of ribosomal 977 sequences

978 (A) 3D organization of rDNA and major satellite sequences at 2-cell stages detected by 3D 979 DNA-FISH using specific probes. Upper panel: single z-section of representative nuclei of 980 non-treated (Ctrl) and treated embryos (CX-5461 and ActD). Lower panel: Amira 3D 981 representation of 2-cell embryos. Green arrows indicate clustered rDNA FISH signal 982 juxtaposed to major satellite signal at NPBs surface (B) 3D organization of rDNA and major 983 satellite sequences at 4-cell stages detected by 3D DNA-FISH. Upper panel: single z-section 984 of representative nuclei of non-treated (Ctrl) and treated embryos (CX-5461 and ActD). 985 Lower panel: Amira 3D representation of each embryo. Green arrows indicate clustered 986 rDNA FISH signal and red arrows major satellite sequences FISH signal with a ring/half-ring 987 shape. White arrowhead point major satellite sequences FISH signal with a round shape called 988 chromocenter. (C and D) Localization of rDNA and major satellite signals in regards to 989 NPBs boundaries at 2-cell stages (C) and 4-cell stages (D) in non-treated (Ctlr) and treated 990 (CX-5461 or ActD) embryos. Left panel: single z-section of a representative nucleus with the 991 line used to draw with Fiji the Intensity plot profile. Right panel: Fluorescence intensity plot 992 profile along the line across multiple channels (blue dot line for DAPI, the green line for

rDNA signal, and red line for major satellite signal). DNA is in blue or grey, rDNA in green, and major satellite sequences in magenta. *Scale bar* = $5\mu m$. mPN: major : mouse major satellite sequences, rDNA: ribosomal DNA.

996

Figure 6: Consequences of RNA Pol I Inhibition on rDNA and major satellite sequences 3D conformation (volume and shape).

999 (A and B) Quantification of rDNA total volume (A) and average sphericity (B) per nucleus 1000 and stage based on the DNA-FISH signal of the ribosomal sequences at 2-cell and 4-cell 1001 stages in non-treated (Ctrl) and treated (CX-5461 or ActD) embryos. The number of nuclei 1002 examined per condition is indicated below the name of the condition (Ctrl in green, CX-5461 1003 in orange, and ActD in purple). (C and D) Quantification of major satellite sequence total 1004 volume (C) and average sphericity (D) per nucleus and per stage based on the DNA-FISH 1005 signal, at 2-cell and 4-cell stages in non-treated (Ctrl) and treated (CX-5461 or ActD) 1006 embryos. The number of nuclei examined per condition is indicated below the name of the 1007 condition (Ctrl in pink, CX-5461 in orange, and ActD in purple). All measurements were 1008 done using Imaris 9.6 (Oxford Instruments). Statistical significance was evaluated using a 1009 non-parametric multiple comparison test (nparcomp, R package). ***, p-value < 0.001.

1010

1011 Figure 7: Consequences of RNA Pol I Inhibition on the expression of major satellite

1012 sequences

1013 (A and B) Quantification by RT-qPCR of the relative expression of ncRNA amplified with

1014 specific primers for major satellite sequence in control embryos (2-cell, 4-cell and 8 cell

1015 stages, pink/red spots) and treated embryos (2-cell and 4-cell stages): with CX-5461 (A,

1016 orange spots) and actinomycin D (B, purple spots). Three to four biological replicates were

1017 done by condition, the expression of these ncRNA was normalized to a fixed amount of

1018 luciferase mRNA added before the reverse transcription. Statistical significance between

1019 control and treated embryos was evaluated using the Mann-Whitney test (Wilcox, R-

1020 package). **, p-value < 0.01. (C) Major satellite transcripts (magenta) visualized using RNA-

1021 FISH in non treated (Ctlr) in non-treated (Ctrl) and treated (CX-5461 or ActD) embryos from

- 1022 2-cell (early (E) and late (L)) to 4-cell and 8-cell stages. ncRNAs spots are indicated by red
- 1023 arrows in treated embryos. (D) Localization of nascent rRNAs (green) and major satellite
- 1024 ncRNAs (red) in mouse embryos from the 4-cell to 16-cell stages by RNA-FISH using
- 1025 specific probes for 5'ETS region and major satellite sequences. Green arrows indicate a 5'ETS

- 1026 RNA-FISH signal closed to a major satellite signal (red arrows). Nuclei were counterstained
- 1027 with DAPI (blue). Scale bar = $5\mu m$.
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- 1029

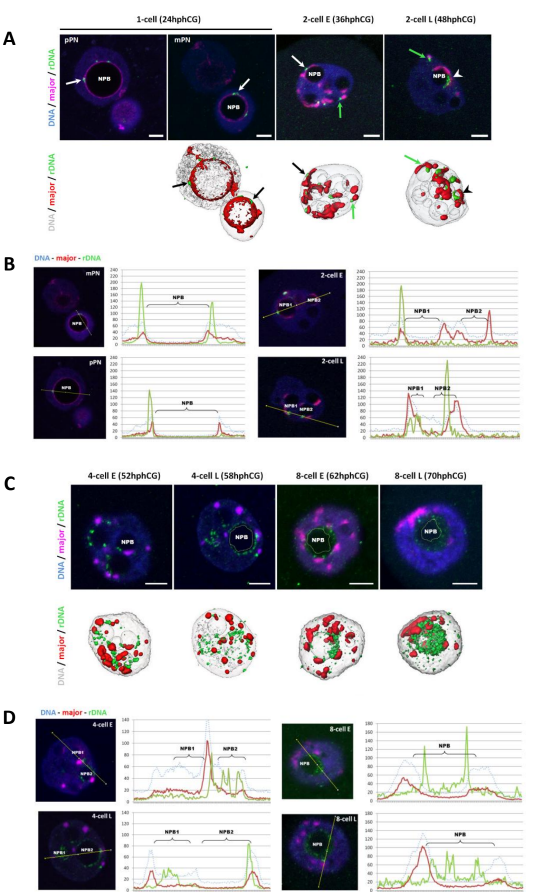
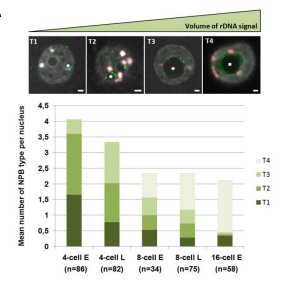
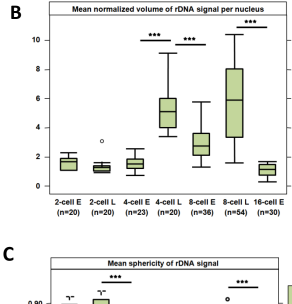


Figure 1 : 3D organization of ribosomal sequences (rDNA) is correlated with their transcriptional state





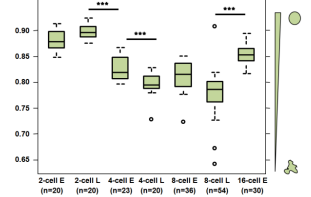
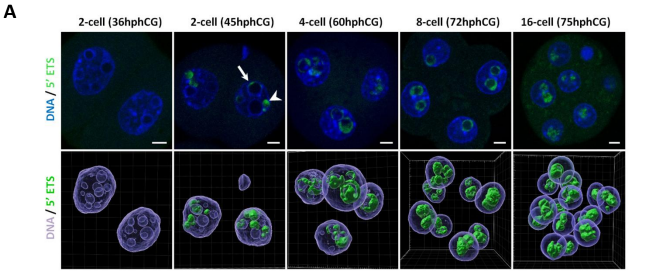


Figure 2 : Features of rDNA spatial organization during pre-implantation development



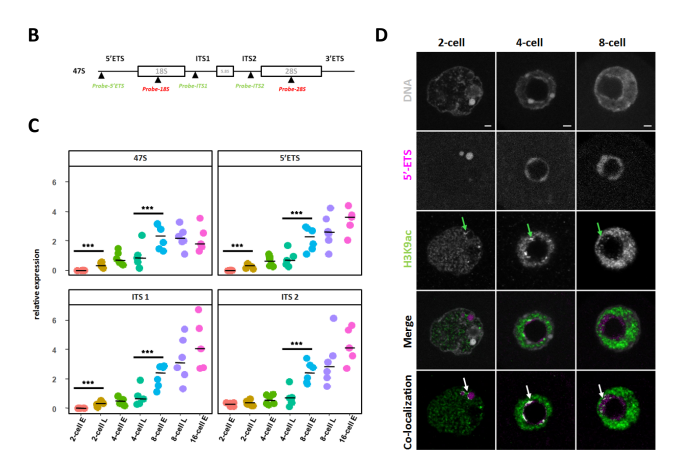
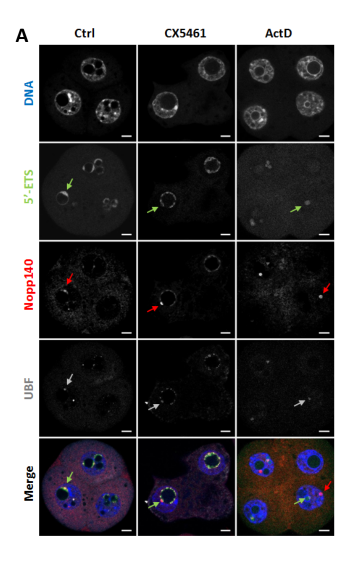
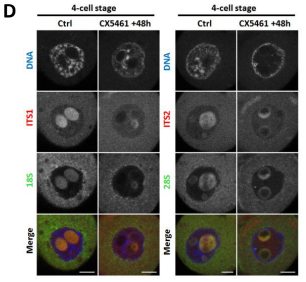
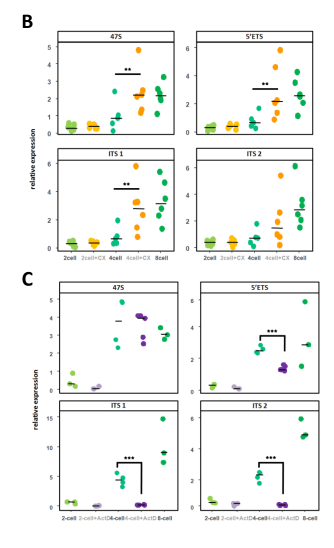


Figure 3: rRNA transcripts organization upon early development







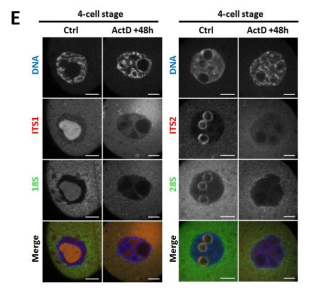
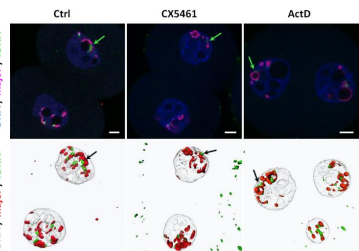
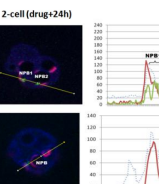
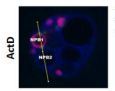


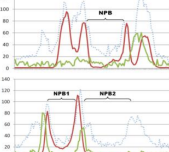
Figure 4: Inhibition of rRNA transcription by CX5461 and ActD

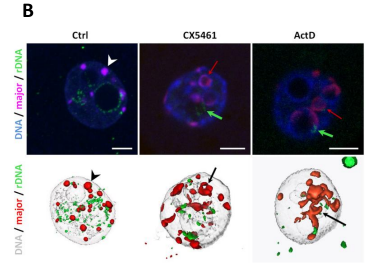












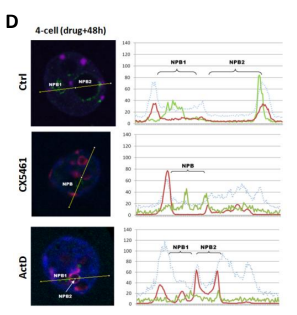


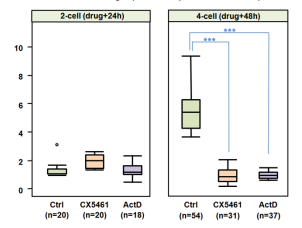
Figure 5: Consequences of RNA Pol I Inhibition on the spatial organization of ribosomal sequences

С

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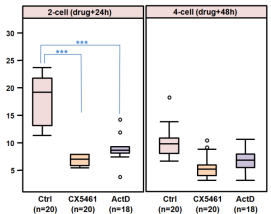
CX5461

volume of rDNA signal (normalized by the volume of nucleus)



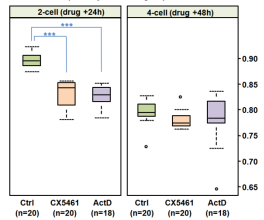
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В

Mean sphericity of rDNA signal per nucleus



D

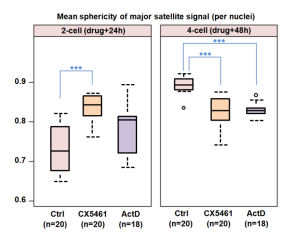
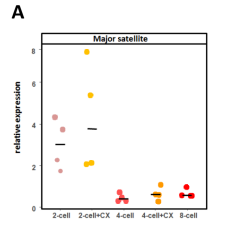
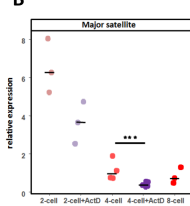


Figure 6: Consequences of RNA Pol I Inhibition on rDNA and major satellite sequences 3D conformation (volume and shape)





16-cell

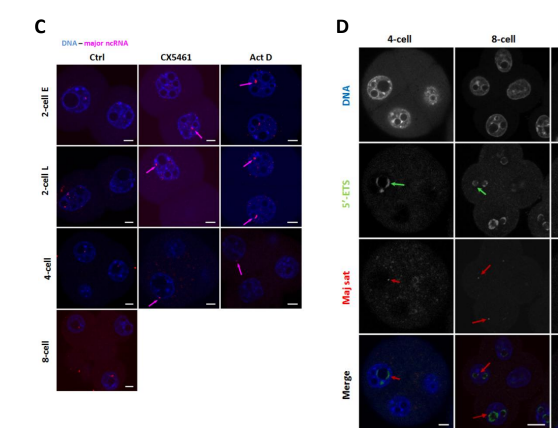


Figure 7 : Consequences of RNA Pol I Inhibition on the expression of major satellite sequences

В