#### ARTICLE





### Histone H2Bub1 deubiquitylation is essential for mouse development, but does not regulate global RNA polymerase II transcription

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#### Abstract

Co-activator complexes dynamically deposit post-translational modifications (PTMs) on histones, or remove them, to regulate chromatin accessibility and/or to create/erase docking surfaces for proteins that recognize histone PTMs. SAGA (Spt-Ada-Gcn5 Acetyltransferase) is an evolutionary conserved multisubunit co-activator complex with modular organization. The deubiquitylation module (DUB) of mammalian SAGA complex is composed of the ubiquitin-specific protease 22 (USP22) and three adaptor proteins, ATXN7, ATXN7L3 and ENY2, which are all needed for the full activity of the USP22 enzyme to remove monoubiquitin (ub1) from histone H2B. Two additional USP22-related ubiquitin hydrolases (called USP27X or USP51) have been described to form alternative DUBs with ATXN7L3 and ENY2, which can also deubiquitylate H2Bub1. Here we report that USP22 and ATXN7L3 are essential for normal embryonic development of mice, however their requirements are not identical during this process, as  $Atxn7l3^{-/-}$  embryos show developmental delay already at embryonic day (E) 7.5, while Usp22<sup>-/-</sup> embryos are normal at this stage, but die at E14.5. Global histone H2Bub1 levels were only slightly affected in Usp22 null embryos, in contrast H2Bub1 levels were strongly increased in Atxn7l3 null embryos and derived cell lines. Our transcriptomic analyses carried out from wild type and  $Atxn7l3^{-/-}$  mouse embryonic stem cells (mESCs), or primary mouse embryonic fibroblasts (MEFs) suggest that the ATXN7L3-related DUB activity regulates only a subset of genes in both cell types. However, the gene sets and the extent of their deregulation were different in mESCs and MEFs. Interestingly, the strong increase of H2Bub1 levels observed in the Atxn7l3<sup>-/-</sup> mESCs, or Atxn7l3<sup>-/-</sup> MEFs, does not correlate with the modest changes in RNA Polymerase II (Pol II) occupancy and lack of changes in Pol II elongation observed in the two  $Atxn7l3^{-/-}$  cellular systems. These observations together indicate that deubiquitylation of histone H2Bub1 does not directly regulate global Pol II transcription elongation.

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#### Introduction

During mouse embryonic development, dynamic modifications of chromatin are essential, as loss of chromatinmodifying enzymes, both writers and erasers, can lead to embryonic lethality, although with different severity [1]. Histone H2B can be modified by the dynamic addition of a

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single ubiquitin (ub1) molecule on lysine 120 in mammals (H2Bub1). The deposition of mono-ubiquitin onto H2B is catalyzed by the RNF20/RNF40 complex in mammals [2–4]. The exact cellular function(s) of the H2Bub1 chromatin mark is not yet fully understood, however, it was suggested that H2Bub1 weakens DNA-histone interactions and therefore disrupts chromatin compaction [5]. H2Bub1 mark was suggested to play a role in several DNA-related and epigenetically regulated processes, such as transcription, repair, replication, homologous recombination, as well as in mRNA processing and export [6-15]. Indeed, chromatin immunoprecipitation coupled to sequencing studies revealed that H2Bub1 is found at gene bodies of expressed genes and is absent from non-expressed chromosomal regions, suggesting that H2Bub1 may be involved in transcription elongation [16–20]. Intriguingly, however, when H2Bub1 deposition was disrupted in mammalian cells by knockdown of RNF20 or knock-out of RNF40, the expression of only a small subset of genes was affected [7, 12, 21]. H2Bub1 has also been implicated in histone crosstalk and shown to be a prerequisite for trimethylation of histone 3 lysine 4 (H3K4me3) around promoter regions both in yeast and mammalian cells [22–27].

H2Bub1 is erased by the deubiquitylation module (DUB) of the co-activator Spt-Ada-Gcn5 acetyltransferase (SAGA) complex [28–31], composed of the ubiquitin-specific protease 22 (USP22) and the ATXN7, ATXN7L3, and ENY2 adaptor proteins, needed for the full activity of USP22 enzyme [32]. ATXN7L3 is critical for directing the DUB module substrate specificity toward H2Bub1 [33]. In human cells, depletion of either ENY2 or ATXN7L3 adaptor protein resulted in a nonfunctional USP22 enzyme [15, 19, 32, 34]. Two additional USP22-related ubiquitin hydrolases (called USP27X or USP51) can also interact with ATXN7L3 and ENY2, and deubiquitylate H2Bub1 independently of the SAGA complex [34]. Thus, in mammalian cells the cellular abundance of H2Bub1 is regulated by the opposing activities of the ubiquitin E3 ligase complex, RNF20/RNF40, and three related DUB modules, each containing one of the homologous deubiquitylases: USP22, USP27X, or USP51 [34, 35]. These different DUB modules have also non-histone substrates [36–43]. As  $Usp22^{-/-}$ mouse embryos die around embryonic day (E) E14.5 [5, 40, 44], the alternative USP27X- and/or USP51-containing DUB modules cannot completely fulfill the role of the USP22-containing DUB module. Usp22<sup>-/-</sup> mouse studies suggest that USP22 is required to regulate apoptosis by deubiquitylating/stabilizing the SIRT1 histone deacetylase and by suppressing p53 functions during embryogenesis, and/or by regulating key signaling pathways crucial for mouse placenta vascularization [40, 44].

To better understand the role of USP22- and/or ATXN7L3-containing DUB modules in vivo, we have

generated mice lacking USP22 or ATXN7L3.  $Atxn7l3^{-/-}$  embryos show developmental delay as early as E7.5, while  $Usp22^{-/-}$  embryos are normal at this stage, and die at E14.5 similarly to previous publications [40, 44]. These results indicate that USP22 and ATXN7L3 are essential for normal development, however, their requirements are not identical. H2Bub1 levels were only slightly affected in  $Usp22^{-/-}$  embryos, while in contrast they strongly increased in  $Atxn7l3^{-/-}$  embryos and derived cellular systems. The genome-wide increase of H2Bub1 retention in mESCs and MEFs lacking ATXN7L3 was investigated and consequences of Atxn7l3 loss of function on cellular homeostasis, differentiation, and RNA polymerase II (Pol II) transcription were analyzed.

#### Materials and methods

### Generation and maintenance of $Usp22^{+/-}$ and $Atxn7/3^{+/-}$ mouse lines

 $Usp22^{+/-}$  and  $Atxn7l3^{+/-}$  mouse lines were generated at the Institut Clinique de la Souris (ICS, Illkirch, France) using mESCs containing the targeting constructs ordered from the International Knockout Mose Consortium (IKMC), including the Knockout Mouse Programme (KOMP) repository (UC, Davis). In the Usp22 targeting construct (Usp22<sup>tm1a(KOMP)Wtsi</sup>) a LacZ and Neo cassette were located in intron 1, flanked by FRT sequences, and loxP sequences were flanking exon 2 (Supplementary Fig. S1A). In the Atxn7l3 targeting construct (Atxn713<sup>tm1.1</sup>(KOMP)Wtsi) a LacZ and Neo cassette were located in intron 2, flanked by FRT sequences, and the LoxP sequences were flanking exon 2 to exon 12 (Supplementary Fig. 1C). Chimeras were generated by injecting the C57BL/6 mESCs containing the targeting constructs into BALB/c blastocysts. Mice heterozygous for the targeting allele were crossed to a Cre-, or FLP-recombinase deleter strains, in order to generate the null alleles Usp22 and  $Atxn7l3^{-}$ , respectively. Then mice heterozygous for the null allele  $(Usp22^{+/-}, \text{ or } Atxn7l3^{+/-})$  were intercrossed to generate homozygous mutant embryos  $(Usp22^{-/-}$  or  $Atxn7l3^{-/-}$ ) as shown in Supplementary Fig. S1A, C. Investigators were not blinded for animal experimentation and no randomization was used as all the conditions were processed at the same time. Genotyping primers are shown in Supplementary Table 1, and examples of genotyping gels are shown in Supplementary Fig. S1B, D. Atxn7l3<sup>+/-</sup> mice were maintained on a mixed B6D2 background. Animal experimentation was carried out according to animal welfare regulations and guidelines of the French Ministry of Agriculture and French Ministry of Higher Education, Research and Innovation.

### Generation and maintenance of $Atxn7I3^{-/-}$ mESCs and $Atxn7I3^{-/-}$ MEFs

To generate  $Usp22^{-/-}$ ,  $Atxn7l3^{-/-}$ , and wild-type (WT) mESCs, timed matings between heterozygous mice were conducted, then at E3.5, pregnant females were sacrificed, uteri were flushed with M2 medium (Sigma-Aldrich), and individual blastocysts were transferred to wells of a 96-well plates pre-coated with 0.1% gelatin. Blastocysts were cultured and expanded in regular mESCs medium [DMEM (4.5 g/l glucose) with 2 mM Glutamax-I, 15% ESQ FBS (Gibco), penicillin, streptomycin, 0.1 mM nonessential amino acids, 0.1% β-mercaptoethanol, 1500 U/mL LIF and two inhibitors (2i; 3  $\mu$ M CHIR99021 and 1  $\mu$ M PD0325901, Axon MedChem)]. After expansion, mESCs were genotyped and frozen.

To generate  $Atxn7l3^{-/-}$  and WT MEFs, timed matings between heterozygous mice were conducted, then at E10.5, pregnant females were sacrificed, and embryos were collected. The embryo yolk sacs were collected for genotyping, and the head and gastrointestinal tract were carefully dissected away from embryos. The remaining carcasses were transferred to individual 1.5 ml Eppendorf tubes, and 50 µl of 0.25% trypsin-EDTA (Gibco) was added and gently triturated five times to dissociate the embryos. The dissociated embryos were incubated in trypsin for 5 min at room temperature, then the trypsin was guenched with 500 µl of FCS. Cells were transferred to individual wells of a 6-well plate pre-coated with 0.1% gelatin and cultured in MEF medium (DMEM, 10% FCS, penicillin, and streptomycin). Cells were visualized with an EVOS XL Core Cell Imaging System (#AMEX-1100, Thermo Fisher Scientific) using a LPlan PH2 10x / 0.25 objective.

#### Protein extraction and Western blot assays

To extract histone proteins, embryos dissected at the indicated embryonic days, or about  $5 \times 10^6$  cells (were lysed with 100 µl acidic extraction buffer (10 mM Hepes, pH 7.9, 1.5 mM MgCl<sub>2</sub>, 10 mM KCl, 0.5 mM DTT and 0.2 M HCl) freshly complemented with 1 × Proteinase Inhibitor Cocktail (Roche) and 10 mM N-ethylmaleimide (Sigma-Aldrich). HCl was added to a final concentration of 0.2 M and incubated on an end-to-end rotator for 2 h at 4 °C. Following the incubation, cell extract was centrifuged at  $20,800 \times g$  for 10 min at 4 °C, to pellet the acid-insoluble material. A solution of 2 M Tris-HCl (pH 8.8) was added to neutralize the supernatant of the acidic extraction. Ten µl of the supernatant, containing histone proteins, were run on 4-12% gels (Bis-tris NuPAGE Novex, Life Technologies), then proteins were transferred and western blot assays were carried out by using standard methods. The following antibodies were used: anti-H3 (Abcam #ab1791) anti-H4 (Invitrogen 3HH4–4G8), anti-H2Bub1 (Cell Signaling Technology, #5546), anti-H3K4me3 (Abcam ab8580), anti-H3K9ac (Merck-Millipore #07-352), Peroxidase AffiniPure F(ab')<sub>2</sub> Fragment Goat Anti-Mouse IgG, Fc $\gamma$  fragment specific (Jackson ImmunoResearch #115-036-071) and Peroxidase AffiniPure Goat Anti-Rabbit IgG (H + L) (Jackson ImmunoResearch #115-035-144). Protein levels were quantified by ImageJ.

#### Actin labeling

Cells were washed twice with 1× PBS, fixed with 4% PFA (Electron Microscopy Science) for 10 min at room temperature (RT). After fixation, cells were washed three times with  $1 \times PBS$ , permeabilized with sterile 0.1% Triton X-100 in PBS for 20 min at RT, then washed three times in 1× PBS. Cells were incubated either with phalloidin conjugated to Alexa 488 dye (Phalloidin-iFluor 488, Abcam ab176753) following the manufacturer's protocol, to label F-actin filaments, or with an anti-β-actin mouse monoclonal antibody (Sigma Aldrich, A5441) at a dilution of 1:1000 in 1× PBS with 10% FCS, overnight at 4 °C. The following day, cells were washed three times with 1× PBS, then  $\beta$ -actin labeled cells were further incubated with secondary Alexa Fluor 488 goat anti-mouse IgG (H+L) (Invitrogen #A-11001) at a dilution of 1:2000 in 1× PBS with 10% FCS for 1 h at RT. The cells were washed three times with 1× PBS, then incubated in 20 mM Hoechst 3342 (Thermo Scientific) for 10 min at RT, before being washed three times with  $1 \times PBS$ , then cells were covered with a coverslip coated in ProLong Gold mounting medium (Invitrogen). Pictures were taken using a Leica DM 4000 B upright microscope equipped with a Photometrics CoolSnap CF Color camera with a HCX PL S-APO  $20 \times /0.50$  objective.

### Colony formation assay and alkaline phosphatase staining

Four thousand mESCs were seeded on gelatin-coated 6-well plates in regular mESC medium (see above) to form colonies at low density. The medium was exchanged every 2 days for 6 days. Alkaline phosphatase (AP) activity test was performed using Red Substrate Kit, Alkaline Phosphatase (Vector Laboratories) according to the manufacturer's instructions: mESC clones were washed with 1× cold PBS and fixed with 4% PFA for 10 min at RT. After fixation, cells were washed twice with H<sub>2</sub>O and incubated in 1 ml AP detection system for 30 min at RT in the dark. Then cells were washed twice with cold 1× PBS, and visualized with an EVOS XL Core Cell Imaging System (#AMEX-1100, Thermo Fisher Scientific) using a LPlan PH2 4×/0.13 objective.

#### **Cell proliferation analysis**

To determine cell proliferation, a total of  $1 \times 10^5$  mESCs per six-well plate were seeded in regular mESC medium and  $3 \times 10^4$  passage 2-MEFs per 24-well plate were seeded in MEF medium. The medium was exchanged every two days. Cell numbers were counted with Countess cell counting chambers (Invitrogen). Statistical analyses were determined by a Wilcoxon rank-sum test (ns p > 0.05;  $*p \le 0.05$ ;  $**p \le 0.01$ ;  $***p \le 0.001$ ).

#### Cell cycle analysis

Hundred thousand mESCs were fixed in 70% EtOH overnight at -4 °C. After fixation, cells were treated with RNase A (100 µg/ml) (Thermo Fisher Scientific, #EN0531) and stained with propidium iodide (40 µg/ml) (Sigma Aldrich, #P-4170) for 30 min at 37 °C. The acquisition of the DNA content was analyzed on FACS CALIBUR (BD Sciences) flow cytometer. Quantitative results were analyzed by FlowJo software (BD Sciences).

#### Apoptosis analysis using annexin-V staining

At the indicated incubation time, floating cells were collected in culture supernatants and adherent cells were harvested by trypsinization. After collection, cells were washed twice with cold 1× PBS, and about  $2 \times 10^5$  cells were resuspended in 100 µl binding buffer (FITC Annexin V Apoptosis Detection Kit, Biolegend). Subsequently, 5 µl FITC Annexin V (FITC Annexin V Apoptosis Detection Kit, Biolegend) and 10 µl propidium iodide was added to the cell suspension. Cells were gently vortexed and incubated in the dark for 15 min at RT. Thereafter, another 400 µl Annexin V binding buffer was added to each tube. Cells were analyzed using a FACS CALIBUR (BD Sciences) flow cytometer. Dot plots were generated using the FlowJo software.

#### Statistics

Statistical analysis of WT versus mutant samples comparison (cell proliferation and H2Bub1 density) was performed using nonparametric two-sided Wilcoxon ranksum test with continuity correction using R (version 5.3). No multiple comparisons were performed therefore no multiple correction were applied. Statistical results were expressed as *P* value or *P* value ranges (ns, p > 0.05; \*,  $p \le 0.05$ ; \*\*,  $p \le 0.01$ ; \*\*\*,  $p \le 0.001$ ). Bar plot graphical data were represented as mean ± SD and individual data points. Proliferation graphical data were represented as mean ± SD. H2Bub1 density data were represented as box plots. Statistical analysis of the RNA-seq datasets was performed using a Wald test (DESeq2, see below).

#### Enrichment of ubiquitylated peptides and quantification of H2Bub1 peptides

Forty million MEFs were harvested, washed three times with PBS and lysed in 1% SDS 0.1 M Tris pH 8 DTT 50 mM, 1× Proteinase Inhibitor Cocktail (Roche) and 50 mM PR-619 DUB inhibitor (UBPBio). The whole-cell lysate was precipitated with TCA and the pellet was washed twice with cold acetone then dried with Speed Vacuum system and weighted. Ten mg of the protein pellet was dissolved in 8 M urea, 5 mM TCEP then alkylated with 10 mM iodoacetamide. Sample was first digested with endoproteinase Lys-C (Wako) at a 1/500 enzyme/protein ratio (w/w) for 4 h, then diluted four times before overnight trypsin digestion (Promega) at a 1:100 ratio. The resulting peptide mixture was desalted on C18 spin-column, quantified with Quantitative Colorimetric Peptide Assay (Thermo Fischer Scientific) and dried on Speed-Vacuum before enrichment with Ubiquitin Remnant Motif (K-E-GG) Kit (Cell Signaling). After Sep Pak desalting, peptides were analyzed in triplicate using an Ultimate 3000 nano coupled in line with an Orbitrap ELITE (Thermo Scientific, San Jose California). Briefly, peptides were separated on a C18 nano-column with a 90 min linear gradient of acetonitrile and analyzed with Top 20 CID (Collision-induced dissociation) data-dependent acquisition method. Data were processed by database searching against Mus musculus Uniprot Proteome database (www.uniprot.org) using Proteome Discoverer 2.2 software (Thermo Fisher Scientific). Precursor and fragment mass tolerance were set at 7 ppm and 0.6 Da respectively. Trypsin was set as an enzyme, and up to two missed cleavages were allowed. Oxidation (M, +15.995), GG (K, +114.043) were set as variable modification and Carbamidomethylation (C) as a fixed modification. Proteins and peptides were filtered with False Discovery Rate <1% (high confidence). Lastly, quantitative values were obtained from Extracted Ion Chromatogram (XIC) and exported in Perseus for statistical analysis [45].

## RNA-seq and chromatin immunoprecipitation coupled high throughput sequencing (ChIP-seq) analyses

For RNA-seq, total RNA was extracted from mESCs and MEFs (3 biological replicates of WT and  $Atxn7l3^{-/-}$  for each cell types) using the NucleoSpin RNA isolation kit (Macherey-Nagel), according to manufacturer's instructions. Libraries were generated from the purified RNA using TruSeq Stranded mRNA (Illumina) protocol. After

checking the quality of the libraries with the Bioanalyser (Agilent), libraries were sequenced on the Illumina HiSeq 4000 at the GenomEast sequencing platform of IGBMC. The raw sequencing data generated reads were preprocessed in order to remove adapter, poly(A) and lowquality sequences (Phred quality score below 20), then were mapped to the mouse mm10 genome using STAR [46]. After PCA analysis, one WT ESCs dataset was excluded because it did not cluster with the other biological replicates. Differentially expressed genes were measured using the DESeq2 package [47]. For the downstream analysis, only the transcripts which base median over 10 normalized reads (DESeq2 normalized reads divided by the median of the transcript length in kb) were considered. Using these criteria 16269 transcripts were expressed in mESCs, and 15084 transcripts were expressed in MEFs.

ChIP-seq experiments were performed using the protocol described in [48], with some minor modifications, including the use of 10 mM N-ethylmaleimide (NEM, Sigma-Aldrich) in all buffers and the use of either the anti-H2Bub1 antibody (MediMabs, NRO3), the anti-RPB1 C-terminal domain (CTD) Pol II antibody (1PB 7G5 [49]) from control and Atxn7l3<sup>-/-</sup> mESCs and MEFs (n = 1, each) or the anti-Ser2P CTD Pol II antibody (3E10 [50]) from control and  $Atxn7l3^{-/-}$  mESCs (n = 1, each). Briefly, mESCs or MEFs were fixed in 1% PFA for 10 min at RT, then the PFA was quenched with glycine at a final concentration of 125 mM for 5 min at RT. Cells were washed two times in 1× cold PBS, scraped, and pelleted. Nuclei were isolated by incubating cells with nuclear isolation buffer [50 mM Tris-HCl pH 8.0, 2 mM EDTA pH 8.0, 0.5% Nonidet P-40, 10% glycerol, Proteinase Inhibitor Cocktail (Roche), 10 mM NEM (and 1× Phos-STOP only in Pol II Ser2P ChIP samples)] for 10 min at 4 °C with gentle agitation, followed by centrifugation at maximum speed to pellet the nuclei. Nuclei were resuspended in sonication buffer [0.1% SDS, 10 mM EDTA, 50 mM Tris-HCl pH 8.0, 1× Proteinase Inhibitor Cocktail, 10 mM NEM (and 1× PhosSTOP only in Pol II Ser2P ChIP samples)]. Then chromatin was sheared with the E220 sonicator (Covaris) and chromatin concentration was measured with the Qubit 3.0 (Thermo Fischer Scientific). Approximately 50 µg of chromatin was used for Pol II or H2Bub1 ChIP, and 240 µg of chromatin was used for Pol II Ser2P ChIP which was diluted in ChIP dilution buffer [0.5% Nonidet P-40, 16.7 mM Tris-HCl pH 8.0, 1.2 mM EDTA, 167 mM NaCl, 1× Proteinase Inhibitor Cocktail, 10 mM NEM (and 1× PhosSTOP in Pol II Ser2P ChIP samples)]. Antibodies were incubated with the chromatin overnight with gentle agitation at 4 °C. The next day, Dynabeads protein G magnetic beads (Invitrogen) were added for 1 h, then were isolated and washed for 5 min at 4 °C, three times with low salt wash buffer (0.1% SDS, 0.5% Nonidet P-40, 2 mM EDTA, 150 mM NaCl, 20 mM and Tris-HCl pH 8.0), three times with high salt wash buffer (0.1% SDS, 0.5% Nonidet P-40, 2 mM EDTA, 500 mM NaCl, 20 mM and Tris-HCl pH 8.0), and once with LiCl wash buffer (0.2 M LiCl, 0.5% Nonidet P-40, 0.5% sodium deoxycholate, 1 mM EDTA, 10 mM Tris-HCl pH8.0), then washed three times with TE buffer. then the beads were incubated in elution buffer (1% SDS, 0.1 M NaHCO<sub>3</sub>) at 65 °C with shaking to elute complexes. Crosslinks were reversed with by adding NaCl at a final concentration of 0.2 M as well as 50 µg/ml RNase A at 65 °C overnight and the following day the samples were treated with 20 µg Proteinase K, 26.6 µl of 1 M Tris-HCl pH 7.9, and 13.3 µl of 0.5 M EDTA incubated at 45 °C for 1 hr, and DNA was phenol/chloroform purified and precipitated. The precipitated DNA was used to generate libraries with the MicroPlex Library Preparation kit v2 (Diagenode) for ChIP-seq according to the manufacturer's instructions. The samples were then sequenced on HiSeq 4000 with read lengths of 50 bp single end, reads were mapped to the mouse mm10 genome by the software Bowtie1. Samples were normalized (see below) and peak calling was performed using the MACS2 software.

#### **Bioinformatics tools and data-analysis methods**

#### Normalization between ChIP-seq datasets

For H2Bub1 ChIP-seq samples to correct for the bias introduced by the differences in the sequencing depth among samples, the total reads present at 3115 intergenic regions far away from genes and larger than 100 kb were selected as described previously [19]. These intergenic reads were used for the normalization of H2Bub1 ChIP-seq samples. We calculated the size factor of these intergenic regions for each sample using DESeq2 (version 1.16) [51]. These size factors were used to normalize the H2Bub1 ChIP-seq data.

For Pol II ChIP-seq and Pol II Ser2P ChIP-seq samples, the total number of mapped reads were used for normalization. Twenty million of total reads were used to generate. bed files for the seqMINER software analysis. The .bigwig files for the Integrative Genomics Viewer (IGV) visualization were generated with makeUCSCfile program in Homer package, the total number of reads was normalized to 10 million and the fragment length was normalized to 100 bp.

#### Calculation of density values

Density values were defined as follows: density = [(number of aligned reads in a region of interest)/(length of the region of interest in bp)]/(size factor  $\times 10^{-8}$ ). For H2Bub1 datasets,

**Table 1** Offsprings from  $Usp22^{+/-}$  intercrosses.

Stage	$Usp22^{+/+}$	$Usp22^{+/-}$	Usp22 <sup>-/-</sup>	Total	Number of litters
E9.5	3 (16.7%)	9 (50%)	6 (33.3%)	18	2
E10.5	5 (23.8%)	11 (52.4%)	5 (23.8%)	21	3
E12.5	8 (19.5%)	21 (51.2%)	12 (29.3%)	41	5
E13.5	4 (28.6%)	7 (50%)	3 (21.4%)	14	2
E14.5	6 (27.3%)	10 (45.4%)	6* (27.3%)	22	3
Weaning	93 (37.6%)	154 (62.4%)	0 (0%)	247	37

\* dead embryo (no beating heart).

Table 2 Of	fsprings from
Atxn7l3 <sup>+/-</sup>	intercrosses.

Stage	Atxn7l3 <sup>+/+</sup>	Atxn7l3 <sup>+/-</sup>	Atxn7l3 <sup>-/-</sup>	Total	Number of litters
E7.5	10 (47.6%)	5 (23.8%)	6 (28.6%)	21	2
E8.5	20 (31.2%)	35 (54.7%)	9 (14.1%)	64	7
E9.5	13 (25.5%)	26 (51%)	12 (23.5%)	51	6
E10.5	53 (28.8%)	83 (45.1%)	48 (26.1%)	184	21
E11.5	7 (28%)	12 (48%)	6 (24%)	25	3
E12.5	9 (47.4%)	10 (52.6%)	0 (0%)	19	3
Weaning	138 (44.7%)	171 (55.3%)	0 (0%)	309	47

we considered only the gene bodies of expressed genes containing at least 1 ChIP-seq read. Out of 16269 expressed genes in mESCs, 15467 contain at least 1 ChIP-seq read. Out of 15084 expressed genes in MEF cells, 14500 contain at least 1 ChIP-seq read (Supplementary Table S4).

#### Generation of average profiles and heat maps

Average profiles and k-means clustering were generated with the seqMINER program [52]. The end of each aligned read was extended to 200 bp in the direction of the read. For the analyses around promoters, the tag density was extracted in a 2 kb window centered on each TSS. For average gene profiles, each gene body was divided into 160 equal bins (the absolute size depending on the gene length), 5 kb upstream and downstream were added. Moreover, 20 equally sized bins (250 bp/bin) were created upstream and downstream of genes. Densities were collected for each dataset in each bin.

#### Calculation of Pol II traveling ratio

Pol II pausing was based on the "Pausing Index" which is also referred to as "Traveling Ratio" [53]. We estimated the "Traveling Ratio" as the ratio of normalized Pol II ChIP-seq reads within the TSS region (-100 to +300 bp around TSS) to that in the gene body (TSS + 300 bp to TSS + 2 kb), for genes expressed more than 100 normalized reads to the median size of transcript in kb and >1 kb in length. Maximal distance and Kolmogorov & Smirnov test were analyzed using R (version 3.5).

#### Results

### In vivo loss of ATXN7L3 results in a more severe phenotype than the loss of USP22

To compare the deubiquitylation requirement for USP22 and ATXN7L3 in vivo, we generated  $Usp22^{+/-}$  and Atxn7l3<sup>+/-</sup> mice (Supplementary Fig. S1A–D). As heterozygous Usp22<sup>+/-</sup>, or Atxn7l3<sup>+/-</sup> mice were phenotypically indistinguishable from their wild type littermates (Tables 1 and 2),  $Usp22^{+/-}$ , or  $Atxn7l3^{+/-}$  mice were intercrossed to obtain  $Atxn7l3^{-/-}$  and  $Usp22^{-/-}$  homozygous mutants.  $Usp22^{-/-}$  embryos started to resorb at E13.5 (Fig. 1Ah) and could not be observed after E14.5. similarly to what has been previously published [40, 44, 54] (Table 1, Fig. 1A). Similarly, no Atxn7l3<sup>-/-</sup> pups could be retrieved at weaning (Table 2). Analysis of  $Atxn7l3^{+/-}$  intercross litters collected at different stages of development revealed a more severe phenotype than  $Usp22^{-/-}$  mutants. A growth delay was already observed as early as E7.5 in  $Atxn7l3^{-/-}$  embryos (Fig. 1B), which did not turn at E9.5 (Fig. 1Bf). From E10.5 onwards, two classes of phenotype were observed: a severe and a mild, corresponding to 2/3 and 1/3 of the  $Atxn7l3^{-/-}$  embryos, respectively. No  $Atxn7l3^{-/-}$  embryos could be retrieved after E11.5 (Table 2). The mild class embryos were growth delayed (Fig. 1Bi and 1Bl) and in some instances blood pooling could be observed (Fig. 1Bh-Bl). The severe class embryos were smaller, failed to turn and displayed shortened trunk, abnormal head development, blood in the heart and enlarged pericardium Fig. 1 Loss of the SAGA DUB adaptor ATXN7L3 results in a more severe phenotype than loss of the DUB enzyme USP22. A. Comparison of Usp22<sup>+/+</sup> and Usp22<sup>-/-</sup> littermates from E9.5 to E14.5. **B**. Comparison of *Atxn7l3*<sup>+/+</sup> and  $Atxn7l3^{-/-}$  littermates from E7.5 to E11.5. From E10.5 onwards, the Atxn7l3<sup>-/-</sup> embryos can be categorized in 2 phenotypic classes; severe (h, k) and mild (i, l). Western blot analyses of E10.5 and E11.5  $Usp22^{+/+}$  and  $Usp22^{-/-}$  (C), as well as E9.5 and E10.5  $Atxn7l3^{+/+}$  and  $Atxn7l3^{-/-}$  (**D**) whole embryo lysates using anti-H2Bub1, anti-H3K4me3, anti-H3K9ac and anti-H4 (C) or anti-H2Bub1, anti-H3K4me3, anti-H3K9ac and anti-H3 (D) antibodies. A Ponceau staining view is displayed at the bottom of each panel. M: molecular weight marker (in kDa). The dotted line in D indicates where the blot was cut. Each lane represents a biological replicate. E, F. H2Bub1 western blot analyses shown in C, D were scanned and analyzed densitometrically with ImageJ and the Ponceau normalized results are represented for each genotype.



(Fig. 1Bh and 1Bk). Our in vivo data demonstrate that loss of the DUB adaptor protein ATXN7L3 has a more severe effect on embryonic development than the USP22 enzyme, in agreement with published in vitro data [34].

## *Atxn7l3<sup>-/-</sup>* embryos show strong increase in global H2Bub1 levels

To investigate the importance of USP22 and ATXN7L3 on H2Bub1 deubiquitylation in vivo, we analyzed global

H2Bub1 levels from WT and  $Usp22^{-/-}$  embryos at E10.5, E11.5, and E12.5 stages (Fig. 1C, and data not shown), as well as from WT and  $Atxn7l3^{-l-}$  embryos at E9.5, or E10.5 stages (Fig. 1D). While almost no changes were observed between controls and  $Usp22^{-/-}$ embryos lysates (Figs. 1C and 1E), a 4-5-fold increase in global H2Bub1 levels were observed in  $Atxn7l3^{-/-}$ embryo extracts (Figs. 1D, F), confirming similar observations made in human cancer cell lines [32, 34]. Histone H3K4 trimethylation, and H3K9 acetylation were not affected in  $Usp22^{-/-}$ , or  $Atxn7l3^{-/-}$  embryos (Figs. 1C, D). Thus, ATXN7L3 is required for the full activity of the three related DUB modules to regulate global H2Bub1 levels, whereas USP22-containing DUB module is less involved in genome-wide deubiquitylation of H2Bub1.

### Atxn713<sup>-/-</sup> mESCs and Atxn713<sup>-/-</sup> MEF-like cells show abnormal proliferation and phenotypes

As  $Usp22^{-/-}$  embryonic phenotypes have been already described [40, 44], we concentrated our analyses on  $Atxn7l3^{-/-}$  mutants. To determine the mechanistic outcome of perturbed DUB function(s), we derived mESCs and MEF-like cells from  $Atxn7l3^{-/-}$  embryos. As in embryos, ATXN7L3 protein levels were undetectable in these  $Atxn7l3^{-/-}$  cellular systems (Supplementary Fig. S2A) and global H2Bub1 levels were significantly upregulated, by almost 4-5-fold in mESCs and about 7.5-8-fold in MEFs (Figs. 2A, 2B and Supplementary Fig. S2B).

Alkaline phosphatase staining and expression of pluripotency markers, such as Pou5f1, Sox2, Klf4, Nanog, *Esrrb*, and *Tfcp2l1* [55], were similar between  $Atxn7l3^{-/-}$ and control mESCs (Fig. 2C, and Supplementary Fig. S2C), indicating that the pluripotency of these cells was not significantly affected in absence of ATXN7L3. Similarly, when apoptotic cell death and non-synchronized cell cycle phase distribution were measured, no significant differences were detected when comparing WT and  $Atxn7l3^{-/-}$  mESCs (Supplementary Fig. S2D, E). However, we observed that  $Atxn7l3^{-/-}$  mESCs colonies were more irregular (Fig. 2C) and proliferated slower (Fig. 2D) compared to WT mESCs. Thus, ATXN7L3-regulated DUB activity may be necessary to facilitate efficient cell cycle progression and consequent cell proliferation, similarly to USP22 that is critical for progressing through G1 phase of the cell cycle [37].

In  $Atxn7l3^{-/-}$  MEFs, many cells had an abnormal round morphology (Fig. 2E, right panel) originating from clusters of cells that proliferated faster than elongated  $Atxn7l3^{+/+}$ MEFs (Fig. 2E). The round  $Atxn7l3^{-/-}$  cells were present in all MEFs generated from E10.5  $Atxn7l3^{-/-}$  embryos (n > 12embryos). The proportion of round cells relative to elongated cells appeared to correlate with the severity of the phenotype. No significant differences were detected when comparing WT and  $Atxn7l3^{-/-}$  MEFs cell cycle phase distribution and apoptotic cell death (Supplementary Fig. S2F, G). However,  $Atxn7l3^{-/-}$  MEFs from passage 2 tended to proliferate somewhat slower for the first 3 days compared to WT MEFs, but then started to grow faster than WT MEFs (Fig. 2F).

Thus, ATXN7L3-linked DUB activity loss, and the resulting increased H2Bub1 levels do not result in severe phenotypic changes in  $Atxn7l3^{-/-}$  mESCs, but cause profound morphological changes and proliferation alterations in  $Atxn7l3^{-/-}$  MEF-like cells.

### Deregulation of gene expression is more severe in $Atxn7l3^{-/-}$ MEFs than in $Atxn7l3^{-/-}$ mESCs

To further characterize ATXN7L3-dependent DUB activity, we measured changes in steady-state mRNA levels between  $Atxn7l3^{+/+}$  and  $Atxn7l3^{-/-}$  mESCs, as well as between  $Atxn7l3^{+/+}$  and  $Atxn7l3^{-/-}$  MEFs by carrying out RNA-seq analyses. We first verified whether the  $Atxn7l3^{-/-}$  MEF-like cells still belong to the MEF lineage in spite of their unusual morphology by comparing our MEFs RNA-seq results with 921 RNA-seq datasets from 272 distinct mouse cell types or tissues [56]. This analysis indicated that the  $Atxn7l3^{-/-}$  MEF-like cells clustered together with  $Atxn7l3^{+/+}$  MEFs or fibroblasts (Supplementary Fig. S3C), suggesting that the  $Atxn7l3^{-/-}$  MEF-like cells belong to the fibroblast lineage.

Differential gene expression analysis between  $Atxn7l3^{-/-}$ and WT mESCs, or Atxn7l3<sup>-/-</sup> and WT MEFs, showed that in both  $Atxn7l3^{-/-}$  samples there are significant numbers of genes which expression was up- or down-regulated (Fig. 3A, B, and Supplementary Fig. S4A, B). When compared to control cells, 1116 upregulated and 810 downregulated transcripts were identified in  $Atxn7l3^{-/-}$ mESCs, while 1185 upregulated and 1555 downregulated transcripts were found in the Atxn7l3<sup>-/-</sup> MEFs (Fig. 3A, B). These observations suggest that out of ~15000 Pol II transcribed genes in mESCs (16269 transcripts), or in MEFs (15084 transcripts), ATXN7L3-linked DUB function regulates the transcription of only a subset of genes. In both cellular systems, downregulated, upregulated, and unchanged gene sets were validated using RT-qPCR (Supplementary Figs. S2C, S3D, E). The fold change in differentially expressed gene was much more pronounced in Atxn7l3<sup>-/-</sup> MEFs than in Atxn7l3<sup>-/-</sup> mESCs (Figs. 3A and 3B), as in  $Atxn7l3^{-/-}$  MEFs about 151 transcripts changed their expression 32-fold or more (up and down), while in  $Atxn7l3^{-/-}$  mESCs only 2 genes changed their expression 32-fold (Fig. 3C). Moreover, when comparing the down- or upregulated genes between Atxn7l3<sup>-/-</sup> mESCs and  $Atxn7l3^{-/-}$  MEFs, only very few transcripts were similarly affected in the two cellular systems (Fig. 3D, E), suggesting



Fig. 2 Primary  $Atxn713^{-/-}$  mESCs and  $Atxn713^{-/-}$  MEF-like cells show strong increase in H2Bub1 levels, abnormal proliferation, and phenotypes. A Western blot analysis of H2Bub1 levels in acidic histone extracts obtained from  $Atxn713^{+/+}$  or  $Atxn713^{-/-}$  mESC clones and  $Atxn713^{+/+}$  or  $Atxn713^{-/-}$  MEFs. Histone H3 western blot and ponceau stained membranes are shown as loading controls. Each lane represents a biological replicate. B Quantification of H2Bub1 levels from A by using ImageJ. The y axis represents the fold change compared with WT cells. Histone H2Bub1 quantification was carried out with H3 normalization. Error bars indicate mean  $\pm$  SD based on two biological replicates (represented by gray dots). C  $Atxn713^{+/+}$  or  $Atxn713^{-/-}$  mESCs (3 biological replicates for each genotype) cultured in serum/LIF plus 2i medium for 6 days were either observed by phase contrast microscopy (left panels) or visualized by alkaline phosphatase staining (right panels). Scale bar, 200 µm. D  $Atxn713^{+/+}$  or  $Atxn713^{-/-}$  mESCs cell proliferation was determined by cell counting at the indicated time points (two biological and three technical replicates for each genotype). Error bars indicate mean  $\pm$  SD based on two biological replicates for each. Statistical significance was calculated using two-sided Wilcoxon rank sum test with continuity correction (ns, p > 0.05; \*,  $p \le 0.01$ ; \*\*\*,  $p \le 0.001$ ). E Morphology of  $Atxn713^{+/+}$  and  $Atxn713^{-/-}$  MEFs derived from E10.5 embryos. Scale bar, 200 µm (>12 biological replicates). F MEF cell number was determined by cell counting at the indicated time points (two biological and three technical replicates for each genotype). Error bars indicate mean  $\pm$  SD based on two biological and three technical replicates for each genotype). Error bars indicate mean  $\pm$  SD based on two biological replicates for each genotype). Error bars indicate mean  $\pm$  SD based on two biological replicates for each genotype). Error bars indicate mean  $\pm$  SD based on t

that ATXN7L3-linked DUB activity regulates different subset of genes in the two cellular environments.

Gene ontology (GO) analyses of genes down-regulated in  $Atxn7l3^{-/-}$  mESCs revealed enrichment of GO categories linked to regulation of transcription, as well as cell differentiation, while in the up-regulated genes the GO categories "Metabolic processes", and "Cell adhesion" were enriched

(Supplementary Fig. S4C, D). Analyses of *Atxn7l3*<sup>-/-</sup> MEFs indicated that genes involved in "Multicellular organism development", and "Cell adhesion" were down-regulated, while genes belonging in "Metabolic" and "Immune system" processes were up-regulated (Supplementary Fig. S4E, 4F). Thus, ATXN7L3-related DUB activities regulate different subsets of genes in the two cellular systems.

Fig. 3 Atxn7l3<sup>-/-</sup> mESCs and **MEF-like cells show** significant deregulation of transcription. MA-plots of RNA-seq data carried out on  $poly(A)^+$  RNA isolated from  $Atxn7l3^{+/+}$  and  $Atxn7l3^{-/}$ mESCs (A two and three biological replicates, respectively), or from Atxn7l3<sup>+/+</sup> and Atxn7l3<sup>-/-</sup> MEFs (B three biological replicates, each). Log<sub>2</sub> fold changes are shown versus Log<sub>2</sub> mean expression signal. Differentially expressed genes were selected using the following thresholds: adjusted p value  $\leq 0.05$ , absolute value of  $Log_2$  fold change  $\geq 1$  and base median expression over 10 normalized reads to the median size of transcript in kb. Orange dots indicate up-regulated genes and blue dots indicates downregulated genes. C The number of significantly affected genes for Atxn7l3<sup>-/-</sup> (KO)/  $Atxn7l3^{+/+}$  (WT) are represented for either mESCs or MEFs: adjusted *p*-value  $\leq 0.05$  and absolute value of fold change ≥2, 4, 8, 32, 64, 128, 256, separately. Venn diagrams indicate the overlap of downregulated (D) and upregulated (E) genes between mESCs and MEFs.



### Cell adhesion and extracellular matrix genes are downregulated in $Atxn7l3^{-/-}$ MEFs

We further investigated the expression changes observed in the "Cell adhesion" GO category, since they could account for the unusual shape of the  $Atxn7l3^{-/-}$  MEFs. RNA-seq analyses indicated that a majority of genes coding for proteins belonging to this GO category: such as cadherins, catenins, collagens, and other cell adhesion molecules, were massively down-regulated in  $Atxn7l3^{-/-}$  MEFs compared to control MEFs (Fig. 4A). The deregulation of several of these genes was confirmed (Supplementary Fig. S3E).

We next analyzed actin cytoskeletal proteins by fluorescence imaging. Using phalloidin staining, labeling F-actin filaments, and anti- $\beta$ -actin immunofluorescence, we observed a massively reduced abundance of F-actin filaments and  $\beta$ -actin in *Atxn7l3<sup>-/-</sup>* MEFs compared to WT MEFs (Fig. 4B), suggesting that loss of ATXN7L3 results in a down-regulation of cell adhesion complexes, which in turn would disrupt actin cytoskeleton in MEFs.

### H2Bub1 levels increase in the gene bodies of $Atxn7l3^{-/-}$ mESCs and $Atxn7l3^{-/-}$ MEFs

To evaluate the changes in the genome-wide distribution of H2Bub1 in  $Atxn7l3^{-/-}$  mESCs, or  $Atxn7l3^{-/-}$  MEFs versus WT controls, ChIP-seq was performed using an anti-H2Bub1 antibody. The genomic distribution of H2Bub1 on several housekeeping genes was visualized using IGV. H2Bub1 levels in both WT cell populations are relatively low, but highly increase in coding regions of both  $Atxn7l3^{-/-}$  mESCs and  $Atxn7l3^{-/-}$  MEFs, often showing a H2Bub1 enrichment peak downstream of the transcription start site (TSS) (Fig. 5A, B).

To analyze quantitatively how the loss of the ATXN7L3linked deubiquitylation activity changes H2Bub1 levels Fig. 4 Cell adhesion genes are down-regulated in  $Atxn7l3^{-/-}$ MEFs. A Heat map showing transcript levels belonging to the cell adhesion GO category from the three biological replicates of  $Atxn7l3^{+/+}$  and  $Atxn7l3^{-/-}$ MEFs for transcripts that are differentially expressed. Log<sub>2</sub> of normalized expression is shown on the vertical column on the left. **B** DAPI and immunofluorescence (IF) images of  $Atxn7l3^{+/+}$  and  $Atxn7l3^{-/-}$ 

MEFs stained with anti- $\beta$ -actin antibody (left) and phalloidin (right) in MEF cells (n = 1). The merge of DAPI and IF images is also shown. Scale bar: 100 µm.



genome-wide, the presence of H2Bub1 over coding sequences of all annotated genes was normalized to intergenic regions and calculated. These analyses indicated that in  $Atxn7l3^{-/-}$  mESCs and in  $Atxn7l3^{-/-}$  MEFs the levels of H2Bub1 increase significantly over the gene body regions of either all expressed genes (Fig. 5C, D), or non-

neighboring expressed genes (after removing overlapping gene units within a 5 kb window 5' and 3'; Supplementary Fig. S5A, B). In transcribed genes, we observed a 1.8-fold increase in H2Bub1 levels in  $Atxn7l3^{-/-}$  mESCs compared to WT controls, and a 6.5-fold increase in  $Atxn7l3^{-/-}$  MEFs (Fig. 5C, D, Supplementary Fig. S5C, D).

Fig. 5 Histone H2Bub1 levels increase strongly in the gene bodies of both Atxn7l3<sup>-</sup> mESCs and Atxn7l3<sup>-/-</sup> MEFs. A, B IGV genomic snapshots of H2Bub1 binding profiles (n = 1)at three selected genes (Pgk1, Klhl11, and Acly). Direction of the transcription is indicated by arrows. Group scaled tag densities on each gene either in mESCs, or in MEFs, are indicated on the left. C, D. Boxplots showing the Log<sub>10</sub> (H2Bub1 density) on the gene bodies of expressed transcripts or intergenic regions. Two-sided Wilcoxon rank sum test with continuity correction (\*\*\*: *p* value < 2.2e-16). Average metagene profiles showing H2Bub1 distribution on the bodies of expressed genes. 16269 expressed genes in mESCs (E) and 15084 expressed genes in MEFs cells (F) were chosen. TSS: transcription start site. TES: transcription end site. -5 kb region upstream of the TSS and +5 kb region downstream of the TES were also included in the average profile analyses.



Next, metagene profiles of H2Bub1 spanning the entire transcribed regions and extending 5 kb upstream from TSSs and 5 kb downstream of the transcription end site (TES) were generated in Atxn7l3<sup>-/-</sup> versus WT mESCs or MEFs (Fig. 5E, F). These profiles revealed a global increase of H2Bub1 over the whole transcribed region with an important enrichment in the region downstream from the TSS in  $Atxn7l3^{-/-}$  compared to WT mESCs (Fig. 5E). Similar results were obtained when we compared WT and  $Atxn7l3^{-/-}$  MEFs, however with a much stronger increase in H2Bub1 levels on the gene-body regions of Atxn7l3<sup>-/-</sup> MEFs than in Atxn7l3<sup>-/-</sup> mESCs (Fig. 5E, F). Moreover, our analyses on all expressed nonoverlapping genes, up- or down-regulated in Atxn7l3<sup>-/-</sup> cells (Supplementary Fig. S5E-J) show that H2Bub1 levels increase strongly in all categories, except at genes downregulated in Atxn7l3<sup>-/-</sup>mESCs (Supplementary Fig. S5F). These results show that ATXN7L3-linked DUB activity is responsible for the genome-wide deubiquitylation over the coding regions of expressed genes in mESCs and MEFs and suggest no general link between genome-wide H2Bub1 erasure defects and Pol II transcription.

### Pol II occupancy does not correlate with the H2Bub1 increase in $Atxn7/3^{-/-}$ cells

To test whether the strong increase in H2Bub1 over the transcribed regions observed in the  $Atxn7l3^{-/-}$  cells would influence Pol II occupancy at promoters and/or in gene bodies,  $Atxn7l3^{-/-}$  mESCs and MEFs, as well as control cells, were subjected to ChIP-seq, using an antibody recognizing the non-modified CTD of the largest subunit of Pol II (RPB1). Data obtained from Atxn7l3<sup>-/-</sup> mESCs, and  $Atxn7l3^{-/-}$  MEFs at selected genes (Fig. 6A, B), or genome-wide (k-means clustering, Fig. 6C, D; and meta-gene plots, Fig. 6E, F) indicated that Pol II occupancy did not change dramatically, compared with the corresponding WT cells. Pol II occupancy at expressed genes was almost not affected at the TSS regions and slightly decreased in the gene body regions in both  $Atxn7l3^{-/-}$  cell types compared to WT cells (Fig. 6E–F). In contrast, at most of Pol II occupied regions, H2Bub1 levels were highly increased in Atxn7l3<sup>-/-</sup> mESCs and MEFs, compared to control cells (Fig. 5E, F). Next, we

Fig. 6 The modest genomewide Pol II occupancy changes do not correlate with the strong H2Bub1 increases observed in the Atxn713<sup>-/-</sup> mESCs or MEFs. A, B IGV genomic snapshots of H2Bub1 and Pol II binding profiles (n = 1) at four selected genes (Zpr1, Bud13, Gan and Cmip). Direction of the transcription is indicated by arrows. Group scaled tag densities on each gene either in mESCs, or in MEFs, are indicated on the left. Kmeans clustering showing the distribution of Pol II and H2Bub1 on genes expressed in mESCs (C, 16269 transcripts) and in MEFs (D, 15084 transcripts) (from -5 kb upstream from the TSS to +5kb downstream of the TES) in control and Atxn713-/- mESC (C) and MEF (D). Average metagene profiles showing Pol II distribution on bodies of expressed genes (from -5 kb upstream from the TSS to +5kb downstream of the TES) in control and Atxn7l3-/- mESCs (E) and MEFs (F). Average profiles depicting Pol II and H2Bub1 distribution around the TSS (TSS -1 kb / +1 kb) of expressed genes in control and Atxn7l3<sup>-/-</sup> mESCs (G) and MEFs (H).



tested Pol II occupancy on genes, which were either down- or up-regulated by the loss of ATXN7L3-linked DUB activity (Fig. 3). Our metagene analyses on the upand down-regulated gene categories in both  $Atxn7l3^{-/-}$ cell types suggest that Pol II occupancy changes tend to correlate with the observed changes in transcript levels, but not with changes in H2Bub1 levels (Supplementary Fig. S6A, B, E and F). As expected, we observed a complete loss of Pol II occupancy at highly downregulated genes, or a strong increase in Pol II occupancy at highly upregulated genes in  $Atxn7l3^{-/-}$  MEFs, compared to control cells (Supplementary Fig. S7). However, these totally opposite Pol II occupancy changes were often accompanied by a strong increase in H2Bub1 levels at these genes (Supplementary Fig. S7). These results together suggest that a strong global H2Bub1 increase in  $Atxn7l3^{-/-}$  cells do not globally deregulate Pol II occupancy at transcribed genes.

Fig. 7 Pol II elongation rates are not changed by H2Bub1 increase in the gene-body regions of  $Atxn7l3^{-/-}$  cells. A. K-means clustering showing the distribution of Pol II and Pol II Ser2P (n = 1) on 16269 expressed genes in mESCs (from -5 kb upstream from the TSS to +5 kb downstream of the TES) in control and Atxn7l3<sup>-/-</sup> mESC. Two clusters have been separated (as indicated) based on the Pol II Ser2P signal accumulation around the TES region in WT ESCs. Average metagene profiles showing Pol II and Pol II Ser2P distribution on bodies of expressed genes (B), Cluster 1 (C) or Cluster 2 (D) (from -5 kb upstream from the TSS to + 5 kb downstream of the TES) in WT and Atxn7l3-/- mESCs. The color code is indicated on the right of the panels. Pol II (E) and Pol II Ser2P (F) traveling ratios were calculated in WT and Atxn7l3-/- mESC, and Pol II (G) traveling ratio was calculated in WT and Atxn7l3-/-MEFs for genes expressed more than 100 normalized reads to the median size of transcript in kb. Kolmogorov & Smirnov test p values and maximal distance (max dist) are indicated.



### Paused Pol II and H2Bub1 peaks downstream of the TSSs do not overlap

Next, we analyzed whether promoter proximal Pol II peaks observed at transcribed genes around the +60 bp region [57-59], would overlap with the H2Bub1 peak observed downstream of the TSSs both in WT and  $Atxn7l3^{-/-}$  cells. As expected meta-gene analyses around the TSSs showed that in both mESCs and MEFs (WT and  $Atxn7l3^{-/-}$ ) Pol II peaks gave the highest signal at around the +60 bp region (Fig. 6G, H). In contrast, H2Bub1 density in WT and Atxn7l3<sup>-/-</sup> mESCs and  $Atxn7l3^{-/-}$  MEFs is low in the +60 bp region and reaches its maximum more downstream, in the +300 bp region (Fig. 6G, H). Importantly, Pol II accumulation at the pause site was not influenced by the large increase of H2Bub1 in  $Atxn7l3^{-/-}$  cells, suggesting that H2Bub1 deubiquitylation by the ATXN7L3-dependent DUB module(s) may not regulate promoter-proximal pausing of Pol II, and/or Pol II turnover.

Pol II elongation rates are not changed by H2Bub1 increase in  $Atxn7/3^{-/-}$  cells

As transcription elongation-linked Ser2 phosphorylation (Ser2P) of Pol II peaks downstream of the TSS and then increases again gradually in the gene body region [60], we performed anti-Pol II-Ser2P ChIP-seq in the Atxn7l3<sup>-/-</sup> mESCs, compared to WT cells. Our genome-wide analyses of Pol II-Ser2P occupancy on expressed genes either by k-means clustering or by metagene profiling indicated no detectable changes around the TSS regions, but a slight (1.13-fold) increase of the Pol II-Ser2P signal toward the TES and downstream of it (Fig. 7A, B). K-means clustering analyses distinguished genes with a strong Pol II-Ser2P signal around their TESs in WT mESCs (Cluster 1, Fig. 7A) and genes devoid of Pol II-Ser2P enrichment around their TESs (Cluster 2, Fig. 7A). Metagene profiles of each cluster indicated that the increase of Pol II-Ser2P signal downstream of the TES in Atxn7l3<sup>-/-</sup> mESCs was more pronounced at genes of Cluster 1 than at Cluster 2 (1.4-fold,

Fig. 7A, C, D), suggesting potential defects in Pol II transcription, elongation rates and/or termination on genes belonging to Cluster 1.

Thus, we determined whether Pol II elongation rates in gene bodies was altered in  $Atxn7l3^{-/-}$  mESCs or MEFs, compared to WT cells. To this end we measured Pol II traveling ratios [53, 61]. Pol II and Pol II-Ser2P traveling ratios in  $Atxn7l3^{-/-}$  mESCs, or Pol II traveling ratio in  $Atxn7l3^{-/-}$  MEFs, compared to WT cells, showed only very minor changes on expressed genes (Fig. 7E, G), suggesting that Pol II elongation rates were not significantly changed in  $Atxn7l3^{-/-}$  mESCs or MEFs.

#### Discussion

# Loss of the DUB adaptor ATXN7L3 results in a more severe phenotype than the loss of the DUB enzyme of SAGA, USP22

The relative abundance and function of the various DUB complexes, their redundant activities and/or compensatory mechanisms, in different cell types, at various stages of mouse embryogenesis has not been explored. Koutelou et al. (2019) [44] revealed that USP22 is essential for placental development. Consistent with our findings, Usp22<sup>-/-</sup> embryos developed normally up to E12.5, but then die around E13.5-E14.5. It has been reported that Usp22 is expressed ubiquitously in the embryo and hyppomorphic Usp22<sup>lacZ/lacZ</sup> mice have a reduced body size and weight [54]. The absence of a strong morphological phenotype in the  $Usp22^{-/-}$  embryos before E13.5 suggests that many key early developmental processes do not require USP22, or that USP22 function can be compensated by other USPs, such as USP27X, USP51. It is however remarkable that placental development in  $Usp22^{-/-}$  embryos cannot be compensated by other USPs, suggesting a possible direct requirement of the SAGA complex in placental development.

On the other hand, no compensation is expected in  $Atxn7l3^{-/-}$  embryos as the absence of ATXN7L3 is supposed to inactivate all SAGA-related DUB complexes [34]. Indeed, the  $Atxn7l3^{-/-}$  phenotype is more severe than that of  $Usp22^{-/-}$ , occurring as early as E7.5. Although it is not known whether the H2Bub1 deubiquitylation is linked to  $Usp22^{-/-}$  or  $Atxn7l3^{-/-}$  embryos phenotypes, it is interesting to note that there is a parallel between the severity of the phenotypes and the changes in H2Bub1 levels. E10.5  $Usp22^{-/-}$  embryos are normal and their genome-wide histone H2Bub1 levels do not increase (Fig. 1C, E), while in contrast E10.5  $Atxn7l3^{-/-}$  embryos are seriously affected and their H2Bub1 levels increase 4-5-fold (Fig. 1D, F).

We observed two categories of  $Atxn7l3^{-/-}$  mutants: i) the severely affected embryos (2/3rd) which are growth retarded, fail to turn and display shortened trunk and abnormal head development; ii) the mildly affected embryos  $(1/3^{rd})$ , which do turn and only display mild growth delay. It is conceivable that ATXN7L3 is involved in embryo patterning as for example, Nodal signaling mutant embryos, which are defective in early patterning of the primitive streak, also fail to turn [62]. Nevertheless, the fact that some  $Atxn7l3^{-/-}$  embryos escape the severe phenotype suggests that ATXN7L3 and the corresponding DUB module(s) could be involved in a developmental checkpoint control at the time of embryo turning. Remarkably, all Atxn7l3<sup>-/-</sup> embryos die around E11.5, and in addition to placental defects, they exhibit cardiovascular defects, as enlarged pericardium and blood pooling in the heart are observed in the severely affected  $Atxn7l3^{-/-}$  embryos. Thus, the comparison of the  $Usp22^{-/-}$  and  $Atxn7l3^{-/-}$  embryo phenotypes suggest that the defects observed in  $Usp22^{-/-}$  embryos could be compensated until E13.5 in the absence of USP22 by the activity of USP27X- and/or USP51-containing DUBs, which would require ATXN7L3 and ENY2 cofactors. Such compensation would not happen in  $Atxn7l3^{-/-}$ embryo, as in the absence of ATXN7L3 all three related DUBs would be inactive.

The comparisons of *Atxn7l3* regulated genes (RNA-seq from our study) with SAGA-bound genes (anti-TAF6L ChIP-seq [63]) in mESCs suggest that only a minority (< 8%) of SAGA-bound genes are regulated (up or down) by the loss of ATXN7L3-linked DUBs. This further suggests that the USP27x- and/or USP51-containing DUBs may have also SAGA-independent gene regulatory functions.

In conclusion, our results showing that  $Usp22^{-l-}$  embryo phenotypes are less severe agree with the biochemical findings suggesting that in  $Usp22^{-l-}$  cells the activity of only one of the three related DUB modules is eliminated. In contrast, in the  $Atxn7l3^{-l-}$  embryos the activities of all three related DUB modules are eliminated, thus, causing a more severe phenotype. It is also possible that ATXN7L3 loss may influence the epithelial-mesenchymal transition during gastrulation. The fact that  $Atxn7l3^{-l-}$  embryos survive until E11.5, suggests that none of these three related DUBs would play an essential role before this embryonic stage, and that also H2Bub1 deubiquitylation is not essential for Pol II transcription before this developmental stage.

### Histone H2Bub1 deubiquitylation is not linked to global Pol II transcription

Although H2B monoubiquitylation has been linked to increased transcription, transcription elongation, DNA replication, mitosis, and meiosis [64], how this histone modification and the erasing of this mark function is not well understood. In *Saccharomyces cerevisiae* H2B mutant, K123R (which cannot be ubiquitylated) the expression of only a low number of genes (about 300–500) is affected [65], suggesting that ubiquitylation and deubiquitylation may not have global transcriptional regulatory functions in yeast. Nevertheless, it has been suggested that H2Bub1 stimulates FACT-mediated displacement of a H2A/H2B dimer which in turn would facilitate the passage of Pol II through the nucleosome [16] and that H2Bub1 would be required for efficient reassembly of nucleosomes behind the elongating Pol II [66, 67]. On the contrary, it was reported that the effect of H2Bub1 on nucleosome stability is relatively modest [68].

If H2Bub1 deposition (by RNF20/RNF40), or H2Bub1 erasure (by the ATXN7L3-containing DUB modules) would carry out opposite genome-wide actions, their gene regulatory actions would result in mirroring effects. However, when comparing RNA-seq data from *Rnf40* knock-out MEFs [12] with our RNA-seq obtained from *Atxn7l3<sup>-/-</sup>* MEFs, we did not observe any anti-correlation between the regulated genes in these two datasets (the Pearson correlation coefficient is -0.042, Supplementary Fig. 8), suggesting that the H2Bub1 deposition and erasure are not (only) carrying out opposite functions on the transcribed genome.

Contrary to H2B monoubiquitylation, it is much less well understood whether H2Bub1 deubiquitylation would be a process significantly impacting Pol II transcription. Previously, by using an ATXN7L3 knock-down strategy in HeLa cells we showed that the ATXN7L3-related DUB activities are directed toward the transcribed region of almost all expressed genes, but correlated only poorly with gene expression [19]. Our present results indicate that impairment of H2Bub1 deubiquitylation does not directly impact transcription initiation and/or elongation, because while we observe a massive H2Bub1 retention at almost every expressed gene in both Atxn7l3<sup>-/-</sup> mESCs and Atxn7l3<sup>-/-</sup> MEFs, Pol II and Pol II-Ser2P occupancy were only slightly impacted and only limited subsets of genes changed expression in both cellular systems (Figs. 3, 5-7). In addition, when comparing nascent and steady state mRNA changes of selected transcripts (Supplementary Fig. 3D) in WT ESCs and  $Atxn7l3^{-/-}$  ESCs, we did not observe any significant differences (data not shown). Moreover, in both cellular systems the lack of correlation between global H2Bub1 increase and consequent inhibition of global transcription suggests that H2Bub1 deubiquitylation does not directly regulate Pol II transcription. In agreement, the H3K4me3 chromatin mark present at the TSSs of active genes in eukaryotes, did not change in  $Usp22^{-/-}$  or in Atxn7l3<sup>-/-</sup> embryos, in spite of the fact that in  $Atxn7l3^{-/-}$ embryos the H2Bub1 levels were increased by 4-5-fold (Fig. 1C, D). Similarly, global H3K9ac levels do not change in  $Usp22^{-/-}$  or in  $Atxn7l3^{-/-}$  embryos (Figs. 1C and 1D). Thus, our study corroborates other recent studies demonstrating catalytic-independent functions of chromatin modifying complexes in mouse ES cells [69–71].

Our results also suggest that the dynamic erasure of the H2Bub1 mark does not seem to influence global Pol II recruitment, pre-initiation complex formation at promoters, promoter proximal pausing or Pol II elongation rates (Figs. 6G, H and 7E–G). However, Ser2 phosphorylation of Pol II in Cluster 1 downstream of the TES regions in *Atxn713<sup>-/-</sup>* ESCs was more pronounced, compared to WT cells (about 1.4-fold, Fig. 7A–C), suggesting a possible compensatory «sensing» mechanism by the kinases regulating Pol II transcription elongation.

Whether the observed embryo and cellular phenotypes in the  $Atxn7l3^{-/-}$  embryos can be directly linked to increased H2Bub1 levels in specific transcribed regions, and/or to deubiquitylation failures of other ubiquitylated protein targets, will need to be further investigated in the future.

#### **Ethics declarations**

#### Animal experimentations

Animal experimentations were carried out according to animal welfare regulations and guidelines of the French Ministry of Agriculture.

#### Data availability

All the datasets generated during the current study are available together in Gene Expression Omnibus (GEO) database under the accession number GSE153587. Individual RNA-seq data can be accessed at GSE153578 and ChIP-seq data at GSE153584.

#### **Code availability**

Data and figures were generated using R (version 3.5). All custom code is available upon request.

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Author contributions FW, FES, DD, SDV, and LT designed the study; LT supervised the project; FW, FES performed and analyzed the molecular and cellular experiments; LN carried out the proteomic analyses; FW, FES, MD, MS, and VF generated the cell lines; FES, FW and MS performed the mouse experiments; FES and SDV analyzed the embryo data. FW, TY, SDV, and LT carried

out bioinformatic analyses; FES, DD, SDV, and LT wrote the paper with input from FW. All the authors gave final approval for publication.

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#### **Compliance with ethical standards**

Conflict of interest The authors declare no competing interests.

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