Title

Novel intramolecular base-pairing of the U8 snoRNA underlies a Mendelian form of cerebral small vessel disease

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Abstract

U8 snoRNA cause How mutations the non-codina the neurological disorder in leukoencephalopathy with calcification and cysts (LCC) is poorly understood. We report the first vertebrate mutant U8 animal model for interrogating LCC-associated pathology. Mutant U8 zebrafish exhibit defective central nervous system development and ribosomal RNA (rRNA) biogenesis, with tp53 activation which monitors ribosome biogenesis. Importantly, LCC patient fibroblasts demonstrate rRNA processing defects. Human precursor-U8 (pre-U8) containing a 3' extension rescued mutant U8 zebrafish, indicating conserved biological function. Analysis of LCCassociated U8 alleles in zebrafish revealed that one null and one hypomorphic, but still functional. allele combine to cause LCC. Mutations involving any one of seven nucleotides within the human pre-U8 3' extension, or 5' region of U8, alter processing of pre-U8, and identify a novel basepairing interaction between the 5' end and 3' extension of human pre-U8. Variants in these seven nucleotides, one of which is present on a single allele in almost all patients, act as hypomorphic mutations. Given that biallelic null U8 alleles are likely incompatible with human development, identification of hypomorphic mutations mediating viable embryogenesis furthers understanding of LCC molecular pathology and cerebral vascular homeostasis.

Introduction

Leukoencephalopathy with calcifications and cysts (LCC), or Labrune syndrome, is a Mendelian neurological disorder of the cerebral small blood vessels associated with increased morbidity and early mortality, presenting at any age from early infancy to late adulthood. Characterized by the radiological triad of cerebral white matter disease, intracranial calcifications and cysts, LCC was recently shown to be an autosomal recessive genetic disorder caused by biallelic mutations in the gene *SNORD118*, encoding the box C/D U8 small nucleolar RNA (snoRNA) (1).

Ribosomes, the apparatus of protein synthesis, consist of 28S, 18S, 5.8S and 5S ribosomal RNA (rRNA) and 80 core ribosomal proteins distributed into 40S and 60S subunits (2). SnoRNAs are an evolutionarily conserved group of non-protein coding RNAs required for the modification and processing of rRNA. U8, a vertebrate-specific factor, is the only known snoRNA essential for maturation of the 28S and 5.8S rRNAs, components of the 60S large subunit (3). U8 snoRNA is required for removal of the 3' external transcribed spacer (3'-ETS) sequence, the first of a series of cleavage steps that liberates the 28S, 5.8S and 18S rRNA sequences from the polycistronic precursor-rRNA (pre-rRNA) (4). Pre-U8 snoRNA contains an m⁷G cap and short 3' extension. Hypermethylation of the m⁷G cap to m₃G and removal of the 3´ extension, through a series of steps that appear to involve nucleo-cytoplasmic shuttling and concurrent ordered association and dissociation of multiple protein components (including the LSm proteins, see below), results in the production of mature U8 snoRNA (5–8). The box C/D motif of U8 is bound by four core proteins: 15.5K, NOP56, NOP58 and fibrillarin, contributing to the formation of the U8 small nucleolar ribonucleoprotein (U8 snoRNP) complex and its localization to the nucleolus, the site of pre-rRNA processing (9). The role of U8 in rRNA maturation implicates LCC as a recently discovered novel ribosomopathy, a group of disorders caused by ribosome biogenesis dysfunction that manifest as a diverse set of highly stereotyped clinical syndromes (10).

Of 33 mutation-positive families identified by Jenkinson et al., 31 probands with LCC were compound heterozygotes for two different *SNORD118* variants, implying the existence of one severe (null), and one milder (hypomorphic) mutation, with biallelic null mutations likely incompatible with life. In total, Jenkinson et al., recorded seven putatively causal mutations in the

invariant box C/D motif, three within the stem of a conserved hairpin loop which would be predicted to decrease the stability of this structure, three within the highly conserved GAUU motif of the LSm-binding site, and four mutations in the short 3' extension of the precursor (1). Promoter mutations were also found that reduced expression levels of U8. Presumably, LCC-associated variant combinations reduce U8 function below a critical level while allowing for viable embryogenesis, thus maintaining sufficient levels of production of functional ribosomes. However, the precise molecular pathology of LCC remains unknown.

Here we report the first vertebrate mutant model system to study U8 snoRNA function. Zebrafish U8 mutants were found to exhibit defective rRNA biogenesis and activation of the tumor suppressor p53 (tp53), which monitors ribosome biogenesis dysfunction in a regulatory loop known as 'nucleolar stress surveillance' (11–13). Functional assessment of LCC disease-associated U8 alleles confirmed the importance of combinatorial null and hypomorphic mutations. We show that the 3' extension of U8 is critical for U8 biological activity, an observation reflected in the fact that a mutation within the 3' extension, or in nucleotides predicted to base-pair with the 3' extension, were recorded in 29 of 33 patients. Assays using HeLa nuclear cell extracts demonstrated that these mutations alter the processing of pre-U8, revealing a novel secondary structure of human pre-U8 snoRNA. Importantly, fibroblasts from patients with LCC also exhibit rRNA processing defects and human pre-U8 snoRNA was found to rescue the zebrafish U8 mutant, indicating conserved biological function. Taken together, these data support the characterization of LCC as a ribosomopathy whose effects are restricted to the cerebral vessels, and of the utility of zebrafish to provide insight into the pathology of human disease and U8 biology.

Results

U8-3 is the predominantly expressed zebrafish U8 during embryogenesis

Zebrafish contain five copies of U8 located on chromosome 10; four copies clustered within the intron of the transcript BX324123, and the remaining copy located between the genes vamp2 and and3 (Fig. 1A). Quantitative RT-PCR analysis, exploiting the single nucleotide polymorphisms present between the five copies of U8 for specificity, identified minimal maternal deposition of U8 transcripts in zebrafish, with U8-5 most highly deposited (Fig. 1B, see fig. S1 for alignment of zebrafish U8 copies). At 24 hours post fertilization (hpf) U8-3 was the sole U8 species identified in the zebrafish embryo, with only weak expression of the clustered U8-1, U8-2, U8-4 and U8-5 induced at 48hpf (albeit increasing thereafter) (Fig. 1B). These data provided a rationale for targeted disruption of the U8-3 gene locus to interrogate U8 function during early embryogenesis. CRISPR/Cas9 was employed to disrupt the U8-3 locus. Creating insertion/deletion events in a non-coding RNA could lead to unpredictable consequences for U8-3 function. Consequently, two guides were used to excise U8-3 from the genome, producing a null allele, herein referred to as ΔU8-3 (fig. S2). By 24hpf, ΔU8-3 mutants exhibited a less defined midbrain-hindbrain boundary and reduced angiogenic sprouting from the dorsal aorta (Fig. 2A). By 48hpf, these ΔU8-3 mutants demonstrated swelling of the fourth ventricle, consistent with abnormal development of the CNS, reduced melanocyte development, smaller eye size, impaired yolk resorption, disturbed branching of the trunk vasculature and a reduction in embryo length consistent with developmental delay (Fig. 2, A and B). A time course analysis indicated that the ΔU8-3 mutant comes to a developmental standstill, reflected in a failure to resorb yolk and expand the swim bladder (fig. S3). Death was observed from 6 days post fertilization (dpf), with 100% mortality recorded by 9dpf (fig. S4). Quantitative RT-PCR analysis confirmed U8-3 expression is lost in U8-3 mutants (Fig. 2C). U8 is required for removal of the 3' external transcribed spacer (3'-ETS) sequence, and in particular the biogenesis of 28S and 5.8S rRNAs (14-16). Tapestation analysis indicated that U8-3 mutants exhibit a preferential reduction in 28S biogenesis compared to 18S (Fig. 2D). Significantly, northern blotting with a probe specific to the 3'-ETS region of the pre-rRNA demonstrated that LCC patient derived fibroblasts, and control cells in which U8 had been knocked down by antisense oligonucleotides, accumulate aberrant unprocessed 3' extended rRNA precursors required for 28S biogenesis, which are also impaired for removal of ITS2 (Fig. 2E, L for longform). Further, these pre-rRNA processing defects alter the ratios of 28S to 18S similar to the U8-3 mutant, preferentially impairing biogenesis of 28S (Fig. 2 D, E). Northern blotting analysis with a probe specific to U8 confirmed antisense oligonucleotide-mediated knockdown of U8, and that LCC patient fibroblasts have reduced levels of total U8 most likely due to reduced stability of one, or both, mutant alleles (fig. S5) (1). Taken together, these data support a conserved biological function for U8 in rRNA processing and ribosome biogenesis, and show that patient-derived cells expressing U8 mutations are indeed defective for ITS2 and 3'-ETS maturation.

A portion of the U8-3 promoter was deleted in the Δ U8-3 allele, potentially containing regulatory elements required for the function of other genes. To confirm the specificity of the Δ U8-3 mutant phenotype a complementation test with an independent U8-3 mutant allele was performed. A guide specific to U8-3, the design of which was facilitated by the absence of a relevant PAM sequence in the other zebrafish U8 gene copies, was used to delete 54bp from the U8-3 gene locus, herein referred to as Δ 54U8-3 (fig. S6, A and B). Δ 54U8-3 mutants demonstrated indistinguishable morphology defects from Δ U8-3 mutants, with both alleles displaying Mendelian autosomal recessive inheritance (fig. S6C). A failure of complementation i.e. the production of 100% wildtype progeny, was observed when a Δ U8-3 heterozygote zebrafish was crossed to a Δ 54U8-3 heterozygote zebrafish, demonstrating that the two mutants are associated with loss of function (LOF) of the same gene (fig. S6C).

Precursor zebrafish U8-3 and human U8 snoRNAs are functionally equivalent

Mutations that lie within the short 3' extension of human U8 in LCC patients imply that this region is of functional significance (1). However, it has previously been reported that exogenous mature U8 snoRNA, lacking the 3' extension sequence, localizes to the nucleolus and rescues endogenous U8 depletion in Xenopus oocytes (17). We first performed an electrophoretic mobility shift assay (EMSA) with *in vitro* transcribed mature zebrafish U8-3 and the highly conserved human 15.5K to confirm the ability of *in vitro* synthesized U8-3 to interact with a key U8 snoRNP factor.

Addition of 15.5K was found to shift U8-3 migration, and also zebrafish U8-1, 2, 4 and 5, demonstrating that these zebrafish U8 species bind 15.5K (fig. S7). Alignment of human *SNORD118* and zebrafish *U8-3* gene loci identified a putative 3′ extension and 3′ BOX in zebrafish U8 (fig. S8). A transient rescue assay was performed, comparing the capacity of exogenous *in vitro* transcribed mature U8-3 and the putative pre-U8-3 snoRNA to rescue the gross morphological abnormalities observed in the ΔU8-3 mutant. Mature or pre-U8-3 snoRNAs were co-injected into the yolk of 1-cell stage zebrafish ΔU8-3 mutants or wildtype siblings with an mRNA encoding a fluorescent protein acting as a tracer, enabling ubiquitous expression of the transcripts throughout the embryo over the first two days of development (Fig. 3A). Mature U8-3 snoRNA rescued the yolk extension and hindbrain swelling of the ΔU8-3 mutant, but not the effect on embryo length (Fig. 3, B and C). The addition of 14 nucleotides 3′ of the mature U8-3 sequence resulted in a rescue of the hindbrain swelling, yolk extension and embryo length of ΔU83 mutants, thereby demonstrating the biological importance of this 3′ extension sequence (Fig. 3).

The above rescue assay provided the means to examine whether human U8 snoRNA is functional in the zebrafish. As for zebrafish pre-U8-3, pre-U8 human snoRNA rescued Δ U8-3 associated gross morphological abnormalities, whereas mature human U8 snoRNA failed to do so (Fig. 3B), indicating that zebrafish and human U8 snoRNA are functionally equivalent. However, although a number of the Δ U8-3 mutant features were rescued by exogenous pre-U8, the eye and head size did not recover to wildtype sibling levels. This lack of complete rescue likely reflects the technical challenge of introducing enough non-coding RNA at the 1-cell stage to sustain a rapidly growing zebrafish embryo over two days of development.

Functional testing identifies one null and one hypomorphic allele in LCC patients

Having demonstrated that human pre-U8 snoRNA is functional in the zebrafish, we next wanted to test the effect of LCC disease associated mutations in U8-3 null embryos. Six alleles were chosen where a defect in functionality had previously been shown *in vitro*: specifically, either a complete (n57G>A and n58A>G) or reduced (n61A>G) ability to interact with 15.5K compared to wildtype, or disrupted 3´ end processing in HeLa nuclear extracts (n*1C>T, n*5C>G and n*9C>T)

(1). Despite molecular evidence suggesting that the n61A>G variant might be hypomorphic, in each case variants affecting nucleotides required for binding to 15.5K were found to act as functional null alleles in that they failed to alter the Δ 54U8-3 mutant phenotype, demonstrating the essential nature of this domain for U8 function (Fig. 4). In contrast, all of the patient mutations in the 3´ extension rescued the morphological abnormalities observed in the Δ 54U8-3 mutant, including the embryo length defect (Fig. 4 B and C), indicating that their function in ribosome biogenesis is preserved. These data likely reflect the null and hypomorphic status of distinct alleles.

Seven distinct hypomorphic U8 alleles are present in most LCC patients

Two thirds of LCC probands in our cohort have been observed to harbor one of four mutations located within the 3' extension of pre-U8 (n*1C>T, n*5C>G, n*9C>T, n*10G>T). This observation, and the ability of these mutations to rescue Δ54U8-3 mutant zebrafish reported in Fig. 4, led us to hypothesize that these alleles confer some degree of preserved ribosomal function, and thereby support viable embryogenesis, in the presence of a second, null, mutation on the other allele. Secondary structure predictions of human pre-U8 using RNAfold (18) identified a novel putative base-paring interaction between the 3' extension of the human pre-U8 snoRNA and the 5' end of the human pre-U8 snoRNA (Fig 5A). Mapping of all the LCC patient variants on this structure revealed that another three disease-associated-mutations, located at the 5´ end of U8, lie precisely within the proposed base-paired region (Fig. 5A). Strikingly, one of the seven mutated nucleotides found within this novel human pre-U8 base paring interaction was observed in 29 of 33 LCC patients overall (Table S1). Interestingly, this predicted secondary structure of the human pre-U8 snoRNA does not appear to contain the canonical kink-turn found in mature U8 (Fig. 5A). We have previously reported that the n*1C>T, n*5C>G, n*9C>T and n*10G>T mutants demonstrate defective 3' end processing in HeLa nuclear extracts (1). The predicted secondary structure of human pre-U8, indicating base-pairing between the 5' end and 3' extension, combined with the knowledge that n*1C>T, n*5C>G, n*9C>T and n*10G>T mutants cause disrupted processing. suggested that the n2T>C, n3C>T and n8G>C mutations might also affect U8 precursor processing. The human pre-U8 snoRNA is processed to the mature U8 snoRNA in HeLa nuclear extracts after 60 minutes (fig. S9). When examining U8 processing intermediates in HeLa nuclear extracts at 30 minutes, a time point before mature U8 snoRNA is produced, we observed that each of the n2T>C, n3C>T and n8G>C mutants conferred an aberrant, increased, rate of 3′ end processing when compared to wildtype (Fig. 5B).

If the proposed interaction of the 5´ and 3´ ends of human pre-U8 is correct, then restoring base-pairing complementarity for the n8G>C mutation would be predicted to return processing to wildtype. Thus, the n8G>C and n*5C>G mutations were combined to test this hypothesis. n*5C>G alone appeared to exhibit slowed processing when compared to wildtype at the early 30 minute timepoint, whereas n8G>C was again associated with an increased rate of processing and production of mature U8 (U8-140) after 30 minutes (Fig. 5C). As predicted by the model, the n8G>C/n*5C>G double mutant pre-U8 was found to confer almost identical processing when compared to wildtype, providing functional evidence for the proposed base-pairing between the 5´ end and 3´ extension of the human pre-U8 snoRNA (Fig. 5C).

The very survival of LCC patients beyond embryogenesis suggests that, like the 3′ extension mutations tested in Fig. 4, the n2T>C, n3C>T, n8G>C and n*10G>T mutant pre-U8 snoRNAs also retain some degree of functional competence necessary for ribosome biogenesis. To test this, rescue experiments using the n2T>C, n3C>T, n8G>C and n*10G>T mutants were performed in the zebrafish Δ54U8-3 mutant. Consistent with our hypothesis, n2T>C, n8G>C and n*10G>T pre-U8 all rescued Δ54U8-3 mutant morphology, including embryo length (fig. S10, A and B). In contrast, the n3C>T pre-U8 did not salvage the yolk extension defect, or embryo length (fig. S10, A and B). However, it should be noted that almost half the nucleotides required for 28S binding in human differ from zebrafish (fig. S10C), and such a lack of conservation might adversely affect the capacity of human U8 to completely substitute for U8-3 function in the zebrafish when single base pair changes are introduced within the 5′ end of human U8. As a consequence, caution must be exercised when interpreting a failure to rescue in these circumstances.

tp53 is activated in response to loss of U8-3 in zebrafish

Perturbation of ribosome biogenesis activates the transcription factor TP53, a tumor suppressor with a role in a wide range of biological processes, including DNA damage, mitochondrial stress, autophagy and oncogenesis (19). Trans-activated tp53 increases expression of different effectors depending on the biological context, including $\Delta 113tp53$, transcribed from intron 4 of the tp53 gene in zebrafish. $\Delta 113tp53$ was found to be upregulated 50-fold in $\Delta U8-3$ mutants compared to wildtype siblings at 24hpf, as were the tp53 target genes mdm2, cyclinG1, p21 and bax (Fig. 6A, fig. S11A). To determine which tissues were affected by loss of U8-3 in zebrafish, a spatio-temporal reporter of tp53 activity was generated that utilized the $\Delta 113tp53$ promoter containing two tp53 binding sites (Fig. 6B) (20). At 17hpf, prior to any observable morphological abnormalities, tp53 trans-activation activity can be observed already in ΔU8-3 mutants (fig. S11B). By 24hpf, tp53 trans-activation activity was detected in the eye, CNS and somites of ΔU8-3 mutants (Fig. 6C, fig. S11C), and by 48hpf, the CNS and somites were the most highly fluorescent tissues (Fig. S11D). Importantly, somite derived vegf expression is critical for correct patterning of the dorsal aorta and for arterial development (21). As such, the delayed sprouting and abnormal branching of the trunk vasculature is unlikely to be a cell autonomous effect, but rather to be secondary to the impairment of somitogenesis. The tissues in which tp53 is activated all display obviously impaired development in the ΔU8-3 mutant as evidence by the reduced eye size, shortened body length and fourth ventricular swelling (fig. S11D).

Inactivation of tp53 signaling partially rescues the ΔU8-3 mutant

To characterize the consequences of tp53 signaling in the Δ U8-3 mutant, Δ U8-3 mutant embryos were crossed onto a tp53 mutant background (22). Notably, genetic inactivation of tp53 prevented induction of Δ 113tp53, mdm2, cyclinG1, p21, and bax expression in Δ U8-3 mutants that is typically observed at 48hpf (Fig. 6D), confirming that the increased expression of these mRNAs is tp53-dependent. Genetic inactivation of tp53 was found to partially restore angiogenic sprouting in Δ U8-3 mutants (Fig. 6E, F), and to rescue ventricular swelling 48hpf (fig. S12A). Although tp53 clearly contributes to the gross morphological abnormalities of Δ U8-3 mutants, inactivation of tp53

signaling would not be predicted to restore the defect in rRNA processing. Indeed, tapestation analysis demonstrated that the preferential inhibition of 28S biogenesis is not rescued in tp53 mutant Δ U8-3 mutant embryos (fig. S12B), and these embryos remain significantly shorter than their U8-3 wildtype sibling counterparts (fig. S12C). Taken together, these data indicate that the reduced embryo length of Δ U8-3 mutants results from impaired ribosome function.

Discussion

Informed by the genetic architecture of the Mendelian disease LCC, here we describe the use of a novel vertebrate mutant animal model of U8 dysfunction to shed new light on the processing of the snoRNA U8, its role in ribosome biogenesis, and the pathology of LCC.

Our data indicate that both zebrafish U8 mutant animals and fibroblasts from patients with LCC exhibit rRNA processing defects. The majority of human syndromes linked to defective ribosome biogenesis are thought to result from haploinsufficiency or partial loss of function of ribosomal proteins or ribosome assembly factors (23–25), and LCC patient-derived fibroblasts express reduced amounts of U8. In our previously published series, 31 of 33 probands with LCC were compound heterozygous for two different *SNORD118* mutant alleles (1). Given that most rare autosomal recessive traits demonstrate enrichment for consanguinity and allelic homozygosity, these molecular data indicate that biallelic null mutations in *SNORD118* are likely incompatible with development. In keeping with this hypothesis, we show that the U8-3 zebrafish mutant is embryonic lethal. Furthermore, functional rescue assays in mutant zebrafish confirm the importance of combinatorial null and hypomorphic mutations in LCC pathology, with partial loss-of-function alleles mediating viable embryogenesis.

The biogenesis and processing of essential, independently transcribed snoRNAs is dependent on highly organized secondary structures and their sequential association with core snoRNP proteins that halt advancing exonucleases, with nucleolytic trimming of a precursor RNA necessary to achieve functionality and to provide metabolic stability and nucleolar localization (7,8,26). Using our zebrafish model we were able to shed new light on the processing of human U8, identifying a novel base-pairing interaction between the 5′ end and 3′ extension of human pre-U8. Twenty one of 33 patients identified by Jenkinson et al. harbored at least one mutation within the 3′ extension, whilst a further eight patients were positive for a mutation in a 5′ situated nucleotide predicted to base-pair with this 3′ extension (1). Human pre-U8 snoRNA containing a 3′ extension rescued the zebrafish U8 mutant, indicating a conserved biological function. To our knowledge, the requirement for such nucleotide base-pairing within the human pre-U8 snoRNA has not been described previously. Close examination of the five zebrafish pre-U8 snoRNAs does not reveal an extensive

potential for base-pairing between the 5' end and the 3' extension as proposed for the human pre-U8 snoRNA. Any base-pairing potential of the zebrafish pre-U8 snoRNAs would require exact mapping of the 5' and 3' ends of the zebrafish pre-U8 snoRNAs and experimental validation. We hypothesize that in human cells, the base-pairing between the 5' and 3' ends of the human pre-U8 snoRNA may be important to regulate the timing of final maturation of U8 by either masking its functionally important 5' end and/or to prevent premature formation of a canonical kink-turn, and associated protein binding, until the U8 snoRNP has sufficiently matured. Such regulation may be important for the correct processing of U8, its metabolic stability, its nucleo-cytoplasmic trafficking (7,27), its final targeting to the nucleolus, or even possibly to facilitate extra-ribosomal functions of U8 independent of its interaction with the pre-rRNAs. Previous work from the Lührmann laboratory identified four U8 snoRNA precursor processing intermediates, leading them to hypothesize that either the U8 pre-snoRNA is processed by more than one exonuclease activity, or that the sequence of the 3' extension may determine the kinetics of 3' processing (7). Our data, as presented in Figure 5, are in agreement with the latter hypothesis, as LCC-associated single nucleotides changes located within the 3´ extension, or that pair with this region, appear to alter the kinetics of processing of the U8 pre-snoRNA. The assembly, processing and export of small nuclear RNPs (snRNPs) also follows a well-defined pathway like that described for snoRNPs (28-30). Precursor forms of the spliceosomal U2 small nuclear RNA (snRNA) pair the 3' extension with an internal sequence, this base-pairing being essential for correct processing into a mature snRNP (31). It is possible that base-pairing of 3' extension sequences may be a more general characteristic of snRNA/snoRNA maturation.

In cell culture models, tp53 is required for activation of cell cycle arrest within 24 hours from the onset of ribosomal stress before, over time, impaired ribosome function becomes rate limiting for cellular growth and division (11,32–34). Depletion of U8 has recently been shown to result in potent induction of TP53 in human cells (13,16), and tp53 signaling in the U8-3 mutant would be predicted to induce an earlier and more complete inhibition of the cell cycle. In keeping with this, the angiogenic sprouting defect observed in the U8-3 mutant at 24hpf was largely, but not completely, rescued by genetic inactivation of tp53; with the partial rescue probably explained by a preferential

reduction of 28S in U8-3 mutants indicating that ribosomal dysfunction is already manifested by this time. By 48hpf, when ribosome dysfunction is likely more pronounced, the morphology of the U8-3 mutant was only minimally salvaged. TP53 is normally constitutively degraded by the proteasome following Hdm2-mediated ubiquitination. When ribosome biogenesis is compromised, unassembled ribosomal components accumulate; this is notably the case for a trimeric complex consisting of the two ribosomal proteins uL5 and uL18 and the 5S rRNA which captures Hdm2, titrating it away from TP53, resulting in a net stabilization of TP53 and the activation of a cell death program (35,36). Although knockdown of U8 in human cells would, therefore, indirectly activate TP53, it remains to be determined whether such activation occurs in the affected neurological tissue of patients with LCC, and whether this plays a mechanistic role in disease pathology.

Alterations in ribosomal components underlie a heterogeneous class of diseases referred to as the ribosomopathies, with a diversity in associated clinical phenotype providing an indication of the multiple specialized roles of the ribosome in normal physiology. Despite ubiquitous expression of the U8 snoRNA, germline mutations in *SNORD118* cause a progressive microangiopathy apparently limited to the cerebral vasculature. Although the phenotype of the exclusively neurological disease LCC is highly distinctive, it is not pathognomonic, as a remarkably similar radiological association is seen in the context of the multisystem disorder Coats plus. Coats plus is caused by mutations in *CTC1* (37) and *STN1* (38), both components of the conserved heterotrimeric telomeric capping complex, and in the telomeric protein POT1 (39). Interestingly then, an unbiased enChip-RNAseq approach identified U8 as a telomere associated RNA (40). As such, the precise link between U8 and cerebral vascular homeostasis awaits elucidation, and may conceivably involve both ribosomal defects, that activate TP53 and/or induce selective impairment of translation of distinct mRNAs in a cell lineage specific context, or currently undefined non-ribosomal functions of U8.

Materials and Methods

Zebrafish strains and husbandry. Establishment and characterization of the *tp53*^{M214K/M214K} and $Tg(kdrl:GFP)^{S843}$ strains have been described elsewhere (22,41). Embryos and adults were maintained under standard laboratory conditions as described previously (42) and experiments were approved by the University of Manchester Ethical Review Board and performed according to UK Home Office regulations. No statistical method was used to predetermine sample size for experimental groups.

Real-time quantitative PCR. RNA was isolated from homogenised zebrafish embryos using TRIzol (Thermo Fisher Scientific) and genomic DNA removed with the TURBO DNA-free Kit (Thermo Fisher Scientific). Reverse transcriptase was performed with the ProtoScript II First Strand cDNA synthesis kit (New England BioLabs) using 1µg of total RNA with random hexamer primers. qRT–PCR analysis was performed with the primers described in table S2 using a 60°C annealing temperature, primers with efficiencies between 95-105%, and the SensiFAST SYBR No-ROX kit (Bioline) and the Mx3000P system (Stratagene).

Genome editing. Capped nls-zCas9-nls, mKate2 or H2B-mCerulean3 mRNA was synthesized using a mMESSAGE mMACHINE SP6 kit (Life Technologies) from a linearized pCS2 construct and purified using a RNeasy mini kit (Qiagen). Zebrafish guides were designed using the Harvard chopchop program (https:// chopchop.rc.fas.harvard.edu). Guide RNA (gRNA) incorporating this target sequence was generated from a PCR amplification product (see table S2 for primer sequences) including the remaining sequence of *S. pyogenes* chimeric single guide RNA through *in vitro* transcription using a HiScribe T7 Quick kit (New England Biolabs). The gRNA was then precipitated in a 1/10 volume of 3M Sodium Acetate and two volumes of 100% ethanol by chilling the reaction at −20 °C for 15 min, then spinning in a microcentrifuge (sigma) at 13K for 15 min, and finally the RNA pellet was resuspended in 15 μl of RNase free water. Cas9 mRNA (250 pg) and gRNA (30 pg) and mKate2 (100pg) or H2B-Cerulean3 (100pg) were injected into the yolk of 1-cell stage embryos and fluorescence used to identify successfully injected embryos. Working

guides were identified by PCR amplifying the target region and running the PCR product on a 3%

agarose gel to identify INDEL events that produce visible shifts or smearing of the amplification

product.

Genotyping. Embryos or fin-clips were placed in PCR tubes with 50 µl of 50 mM NaOH and

denatured for 20 min at 95 °C. A volume of 20 µl of Tris-HCl pH 8 was added to each tube and 1

μl of the genomic DNA used for PCR amplification

Polymerase chain reaction conditions. Polymerase chain reaction (PCR) was performed in a

25–50 µl reaction mix containing DNA template (0.1–100 ng DNA), sense and antisense primer

0.8µm each, 0.25 mM dNTPs (Bioline), 1X HF buffer (New England Biolabs), 1U Phusion Tag

polymerase (New England Biolabs), 0.5 mM MgCl (New England Biolabs) and 1.5 µl DMSO (New

England Biolabs) per 50 µl reaction. PCR was performed in a Techne TC-PLUS or Alpha Thermal

Cycler PCRMAX machine with an initial denaturing step at 98 °C for 3 min followed by 35 cycles

of denaturing at 98 °C for 10 s, annealing at 60 °C for 30 s, and amplification at 72 °C for 45 s/1

kb. A final 5-10 min cycle at 72 °C was routinely performed to allow the complete extension phase

to occur

Imaging and embryo measurement. Zebrafish embryos were anesthetized using MS-222

(Sigma-Aldrich) and imaged on an M165FC fluorescent stereomicroscope (Leica) with a DFC310

FX camera (Leica). For measuring the length of embryos, images were taken at 2.5x magnification

and embryo length quantified in the CorelDRAW graphic suite. For confocal microscopy, images

were taken from anesthetized embryos using a Leica TCS SP8 AOBS upright confocal using a

20x 0.50 Plan Flurotar objective and processed using LAS X (Leica version 3.5.2.18963).

Quantitation of 28S:18S ratios. Total RNA was run on a TapeStation 4200 (Agilent) according

to manufacturers instructions. Total RNA with integrity values in excess of 9 were selected for

16

28S:18S quantitation

ASO-knockdown of U8 and rRNA processing assays in fibroblasts. ASO-mediated depletion of U8 in control cells (HCT116, colon carcinoma) was performed as described previously (16). RNA extraction and pre-rRNA processing analysis of U8 depleted cells and patient-derived fibroblasts (LCC) was performed as described previously (16,43). The ATTCC fibroblast control PCS-201-012 (primary dermal fibroblasts normal human adult) was used. Sequence of anti-U8 ASO used in depletion: mGmGmAmUm*UATCCCACCTG*mAmCmGmAmU

N and mN are deoxynucleotide and 2´-O-methoxyethylribonucleotide, respectively. Phosphodiester backbones are phosphorothioates.

In vitro transcription of U8 RNA variants. U8 DNA templates containing a T7 consensus sequence were PCR amplified from human or zebrafish genomic DNA (see Supplementary Table 2 for primer sequences) and purified after agarose gel electrophoresis using a QIAEX II kit (Qiagen). Human and zebrafish U8 snoRNAs were generated using 400-1000ng of template DNA and a mMESSAGE mMACHINE T7 kit (Life Technologies) followed by Lithium Chloride precipitation and quantitation using a NanoDrop (Thermo Fisher Scientific)

Microinjection and transgenesis. The yolk of fertilised 1-cell stage embryos were microinjected with 2nl of synthetic mRNA/snoRNA, or for transgenesis, with 40pg of both DNA and tol2 transposase mRNA (44), using a PLI-90 pico-injector (Harvard Apparatus) and Leica MZ6 stereomicroscope

Electrophoretic mobility shift assays. For EMSAs, recombinant His-15.5K was incubated with 50,000 dpm ³²P-end-labeled U8 snoRNA in EMSA buffer (20 mM HEPES-KOH, 150 mM KCl, 1.5 mM MgCl₂, 0.2 mM EDTA and 0.1% Triton X-100) for 30–45 min on ice. Resulting RNA–protein complexes were resolved on a native 7% acrylamide gel for 8.5 h at 4 °C. Gels were dried and exposed to X-ray film for approximately 8 hr at -80 °C in the presence of an intensifying screen.

3'-processing assays. For 3'-processing assays, ³²P-end-labeled U8 snoRNA was incubated with HeLa nuclear extract (CIL Biotech) at 30°C in buffer containing 0.25 mM ATP, 10 mM phosphocreatine, 3.2 mM MgCl₂, 20 mM HEPES KOH, pH 7.9, 2.6% polyvinyl alcohol and 240 U RNasin (Promega). At 0, 30 and 60 min, 10 μl of the reaction was removed and added to a tube containing 4 μl of stop solution (1 mg/ml proteinase K, 50 mM EDTA and 1% SDS). Reactions were then incubated at 37 °C for 15 min, phenol extracted, precipitated and resolved on a 5% acrylamide/7 M urea gel. Gels were dried and exposed to X-ray film at −80 °C in the presence of an intensifying screen.

Statistics and reproducibility. All statistical analyses were performed using GraphPad Prism 7.03 or Microsoft Excel software. Results are presented as mean \pm s.d. For all analyses P < 0.05 was considered statistically significant. Statistical methods were not used to predetermine sample size, which varies between experiments. Experiments were not randomized. The investigators were not blinded to allocation during experiments and outcome assessment. For Fig. 1B significance was determined by one-way ANOVA and post-hoc Tukey's multiple comparisons test. For fig. S3 significance was determined by a Mantel-Cox test. For all other statistical analyses' significance was determined by an unpaired t-test. The number of biological replicates upon which significance was determined are specified in the figure legend

Oligonucleotides used in this study Refer to Table S2

Supplementary Materials

- Table S1. LCC cohort published in Jenkinson 2016.
- Table S2. Oligonucleotides used in this study.
- Fig. S1. Human U8 and zebrafish U8 1-5 mature sequences exhibit extensive conservation.
- Fig. S2. The Δ U8-3 mutant is homozygous for a 1.26kb deletion encompassing the *U8-3* gene locus.
- Fig. S3. The ΔU8-3 mutant exhibits a range of gross morphological abnormalities and comes to a developmental standstill.
- Fig. S4. The $\Delta U8-3$ mutant phenotype is lethal.
- Fig. S5. LCC patient fibroblasts express less total U8 compared to wildtype fibroblasts.
- Fig. S6. Non-complementation with independent mutant alleles of U8-3 demonstrates that the Δ U8-3 mutant is specific.
- Fig. S7. EMSA demonstrates all five zebrafish U8s bind 15.5K.
- Fig. S8. Alignment of human U8 and zebrafish 3' extension sequences.
- Fig. S9. Time course of wild type human pre-U8 snoRNA processing.
- Fig. S10. Functional testing of four LCC patient mutant U8 snoRNAs in zebrafish.
- Fig. S11. tp53 is trans-activated in response to loss of *U8-3* in zebrafish.
- Fig. S12. Inactivation of tp53 partially rescues the Δ U8-3 mutant phenotype though fails to rescue the embryo length or rRNA biogenesis defect.

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Investigation: A.P.B performed the majority of the experiments and analysis of data with the

exception of 2E, S5 (L.W, C.U.), 5B, 5C, S7, S9 (R.T.O.), 6C (S.C.) and Table S1 (G.I.R.)

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Writing original draft: A.P.B., Y.J.C., R.T.O.

Writing – review and editing: all authors

Competing interests

The authors declare no competing interests.

Figures and Legends

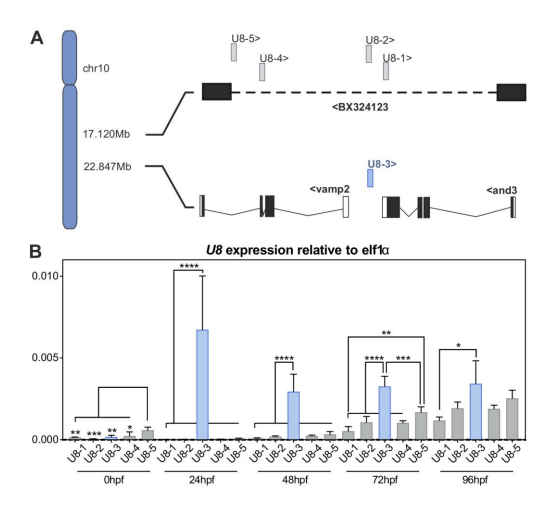


Fig. 1. U8-3 is the predominantly expressed U8 snoRNA during zebrafish embryogenesis. **A**) schematic depicting the five U8 zebrafish snoRNA gene copies located on chromosome 10. *U8-3* (blue) is located between the *vamp2* and *and3* genes, whereas the other four copies (grey) are clustered within the intron (dashed line) of the non-coding transcript BX324123. < or > indicates direction of transcription. **B**) quantitative RT-PCR to *U8-1*, *U8-2*, *U8-3*, *U8-4* and *U8-5* snoRNA transcripts at the indicated developmental time points. n=4 biological replicates per time point. hpf – hours post fertilization.

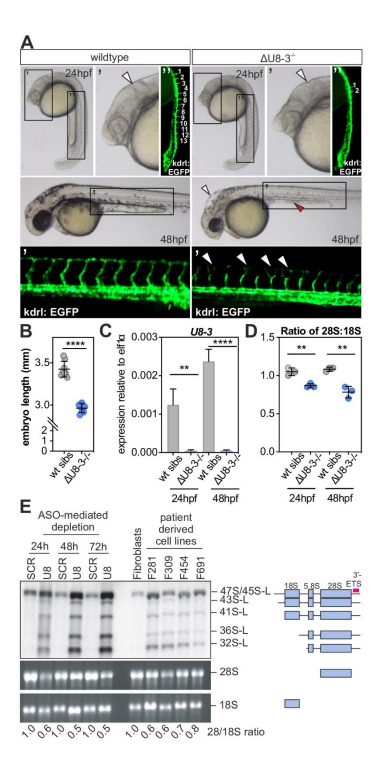


Fig. 2. U8-3 is required for correct development of the central nervous system and vasculature in the zebrafish and 28S biogenesis in fish and patient-derived cells.

A) Δ U8-3 mutants exhibit a less defined midbrain-hindbrain boundary (arrowheads) and reduced angiogenic sprouting (numbered in white, visualized with the kdrl:EGFP transgene) from the dorsal aorta compared to wildtype siblings at 24hpf. By 48hpf Δ U8-3 mutant embryos exhibit hindbrain swelling (arrowhead), an underdeveloped yolk extension (red arrowhead), reduced eye size, impaired melanocyte development and disorganised trunk vasculature compared to wildtype siblings. hpf, hours post fertilization. **B**) quantitation of embryo length of the indicated genotype 48hpf. n=8 embryos per genotype. **C**) qRT-PCR of *U8-3* snoRNA in the indicated genotype and developmental time point. n=4 biological replicates per genotype. **D**) Tapestation analysis demonstrates Δ U8-3 mutant embryos display a preferential reduction in 28S biogenesis consistent with defective 3´-ETS processing. n=3 biological replicates per genotype. **E**) ASO-mediated

depletion of human U8 snoRNA results in accumulation of 3´ extended forms of pre-rRNA intermediates (identified by a 3´-ETS specific probe, pink bar) which are required for biogenesis of 28S rRNA. Likewise, these same 3´ extended intermediates accumulate in LCC patient fibroblast cell lines F281, F306, F454 and F691 when compared to wildtype fibroblasts (ATCC). ASO, antisense oligonucleotide. SCR, non-targeting ASO control. The mature 28S:18S ratio, established by densitometry, are indicated.

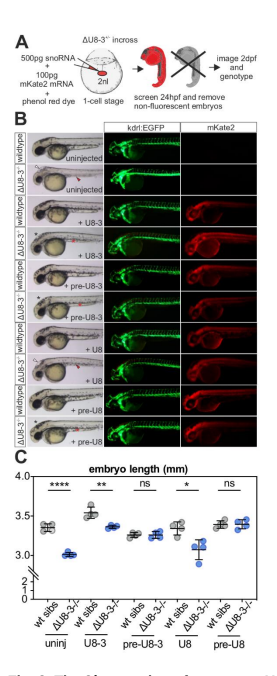


Fig. 3. The 3' extension of precursor U8 is required for optimal U8 biological activity.

A) schematic depicting the experimental design for transient ubiquitous expression of U8 snoRNA variants and mKate2 mRNA. Bolus size is monitored by a phenol red dye and successful uptake of the microinjected solution into the animal pole traced by red fluorescence. B) representative brightfield and fluorescent images of the genotype, exogenous snoRNA, transgenic or fluorescent protein indicated, taken at 48hpf. U8 denotes human sequence whereas U8-3 denotes zebrafish. White arrowheads denotes hindbrain swelling, red arrowheads denotes aberrant yolk extension, black asterisks denotes rescued hindbrain and red asterisks denotes rescued yolk extension. C) quantitation of embryo length for the genotypes and introduced snoRNAs indicated. uninj, uninjected. n=4 embryos per genotype.

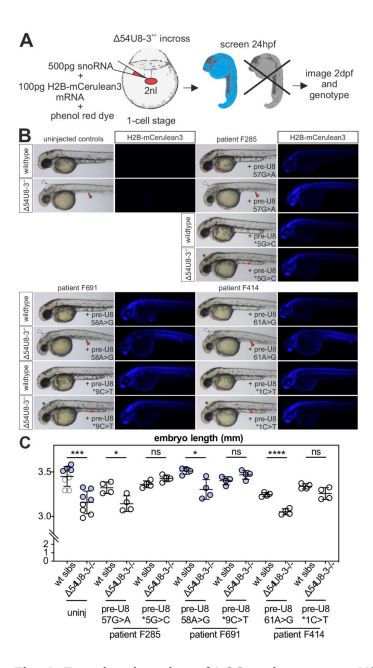


Fig. 4. Functional testing of LCC patient mutant U8 snoRNAs identifies one null and one functional allele.

A) schematic depicting the experimental design for transient ubiquitous expression of U8 snoRNA mutants and H2B-mCerulean3 mRNA. Bolus size is monitored by phenol red dye and successful uptake of the microinjected solution into the animal pole traced by fluorescence. B) LCC mutant snoRNAs specific to the region required for 15.5K binding (n57, n58 and n61) fail to rescue the hindbrain swelling (white arrowheads) and yolk extension (red arrowheads) of Δ 54U8-3 mutants, whereas LCC mutants specific to the 3´ extension (n*1, n*5, n*9) rescue hindbrain swelling (black asterisks), yolk abnormalities (red asterisks) and pigmentation defects. C) LCC mutant snoRNAs specific to the region required for 15.5K binding (n57, n58 and n61) do not rescue the reduced embryo length of Δ 54U8-3 mutants, whereas LCC mutants specific to the 3´ extension restore Δ 54U8-3 mutant embryo length to that of wildtype siblings. n=4-8 embryos per genotype. Blue and black data points represent embryos collected from two different pairs of heterozygote Δ 54U8-3 adults.

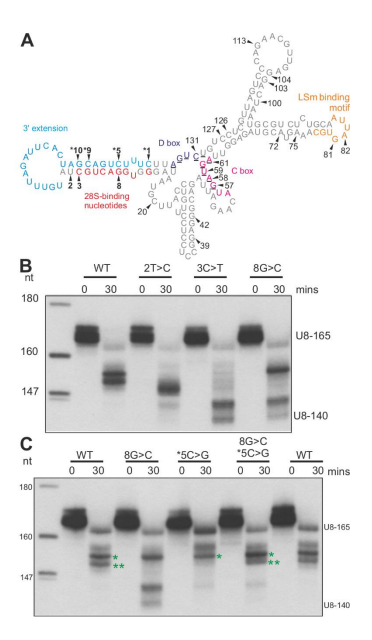


Fig. 5. Aberrantly processed mutant U8 snoRNA alleles are present in 29 of 33 LCC patients. A) schematic depicting the secondary structure of human precursor U8 predicted by RNAfold software (18). Black arrowheads/bold numbering identify seven mutated nucleotides in LCC patients that lie within a base-pairing region encompassing the 3' extension. The remaining mutated nucleotides recorded in LCC patients are numbered (refer to Supplementary Table S1 for a complete list of LCC-associated mutant alleles). The novel 5' end/3' extension base-pairing may preclude premature kink-turn formation (normally involving residues: 57-61 and 130-134, underlined) until U8 has matured sufficiently. B) processing of 5' end-radiolabeled in vitro transcribed precursor U8 wildtype/mutant snoRNA (U8-165) was assessed in HeLa nuclear extracts. At 30 min, 2T>C, 3C>T, and 8G>C mutant U8 snoRNAs exhibit an enhanced rate of processing when compared to wildtype as demonstrated by the presence of mature U8 (U8-140). C) processing of 5' end-radiolabeled in vitro transcribed precursor U8 wildtype/mutant snoRNA (U8-165) was assessed in HeLa nuclear extracts. At 30 min, the n*5C>G mutant U8 is blocked in processing when compared to wildtype (** band is absent), in contrast to 8G>C mutant U8 snoRNA that exhibits production of mature U8 (U8-140). When base-pairing complementarity is restored by combining the two mutations the pattern of processing is restored to wildtype.

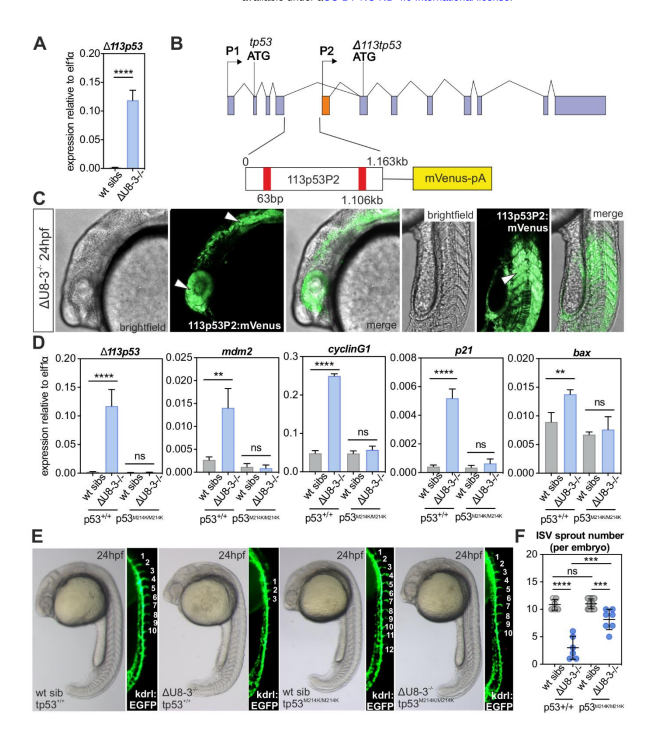


Fig. 6. tp53 is trans-activated in response to loss of *U8-3* in zebrafish.

A) quantitative RT-PCR of Δ113tp53 transcripts 24hpf. B) schematic depicting 1.163kb of intron 4 of the tp53 gene that contains two tp53 binding sites (red bars) used to drive expression of mVenus when tp53 is trans-activated. C) tp53 activity is observed in numerous tissues that develop abnormally in ΔU8-3 mutants, including the eye, hind-brain and somites (arrowheads) at 24hpf. D) quantitative RT-PCR demonstrates significant upregulation of the tp53 target genes Δ113tp53, mdm2, cyclinG1, p21 and bax in ΔU8-3 mutants when compared to wildtype siblings at 48hpf, that is abrogated by genetic inactivation of tp53. hpf – hours post fertilization. n=4 biological replicates per genotype. E) representative brightfield and corresponding fluorescent images of the indicated genotype and transgenic reporter. Angiogenic sprout number indicated by white numbering. F) quantitation of angiogenic sprouts in the indicated genotype. n=6-13 embryos per genotype.

Supplementary Table S1 LCC cohort published in Jenkinson 2016

Family	Family members tested	Ethnicity	Chromosomal position (Hg19)	Variants detected	Zygosity	gnomAD frequency
F172 VS	1A (child)	White European	g.8076761C>A	n.10*G>T	Het	0.002495
		(Swedish)	g.8076851dup	n.56dup	Het	0.000008617
F278 Pel#	1A, 3U, M, F	White European	g.8076832T>C	n.75A>G	Het	0.00001723
	(child)	(North American)	g.8076899C>G	n.8G>C	Het	0.00001523
F281 TR	1A, M, F (child)	White European	g.8076826C>G	n.81G>C	Het	Novel
		(Estonian)	g.8076904G>A	n.3C>T	Het	0.001482
F285 JW	1A (12y)	White European	g.8076766G>C	n.*5C>G	Het	0.0006056
		(British)	g.8076850C>T	n.57G>A	Het	0.00002657
F309	1A (child)	White European	g.8076776G>C	n.131C>G	Het	0.00001520
TO'N		(North American)	g.8076904G>A	n.3C>T	Het	0.001482
F330 DW	1A (child)	White European	g.8076825T>C	n.82A>G	Het	0.00006832
		(British)	g.8076899C>T	n.8G>A	Het	0.002756
F331	2A, M, F (child)	White European	g.8076761C>A	n.*10G>T	Het	0.002495
		(German)	g.8076835T>C	n.72A>G	Het	0.00006894
F334 EH	1A (child)	White European	g.8076825T>C	n.82A>G	Het	0.00006832
		(British)	g.8076761C>A	n.*10G>T	Het	0.002495
F337 LP	1A, M, F (child)	White European	g.8076905A>G	n.2T>C	Het	0.00002167
		(Italian)	g.8076849dup	n.58dup	Het	Novel
F343	1A (child)	White European	g.8076766G>C	n.*5C>G	Het	0.0006056
VDH		(Dutch)	g.8076885_8076913dup	n7_22dup	Het	Novel
F344 FAH	1A (child)	East African	g.8076794G>A	n.113C>T	Hom	0.00004177
		(Somalian)	g.8076899C>G	n.8G>C	Hom	0.00001523
F362 JB	2A, 1U, M, F	White European	g.8076766G>C	n.*5C>G	Het	0.0006056
	(child)	(British)	g.8076887G>A	n.20C>T	Het	0.00003021

F414 RPJ	1A (25y)	White European	g.8076770G>A	n.*1C>T	Het	0.005054
	• • • • • • • • • • • • • • • • • • • •	(Finnish)	g.8076846T>C	n.61A>G	Het	0.00006073
F426 TrF	2A, M, F (child)	White European	g.8076766G>C	n.*5C>G	Het	0.0006056
		(North American)	g.8076826C>T	n.81G>A	Het	0.00004740
F433 AL	1A (50y)	White European	g.8076766G>C	n.*5C>G	Het	0.0006056
		(British)	g.8076887G>A	n.20C>T	Het	0.00003021
F445 ED-	1A (child)	White European	g.8076766G>C	n.*5C>G	Het	0.0006056
P		(British)	g.8076780G>C	n.127C>G	Het	0.00004316
F446 GL	1A (child)	White European (British)	g.8076766G>C	n.*5C>G	Hom	0.0006056
F454 AA	2A (child)	White European	g.8076766G>C	n.*5C>G	Het	0.0006056
		(Canadian)	g.8076955_8076960del	n5449del	Het	Novel
<mark>F465</mark> DJ	1A, M, F (child)	White European	g.8076825T>C	n.82A>G	Het	0.00006832
		(German)	g.8076846_8076847insA	n.60_61insT	Het	0.000007591
F521 HW	2A (child)	White European	g.8076803C>T	n.104G>A	Het	0.0004176
		(North American)	g.8076776G>C	n.131C>G	Het	0.00001520
F551 EDP	1A (child)	White European	g.8076762G>A	n.*9C>T	Het	0.001923
		(Belgium)	g.8076780G>C	n.127C>G	Het	0.00004316
F564 LDR	1A (child)	White European	g.8076770G>A	n.*1C>T	Het	0.005054
		(Belgium)	g.8076781G>A	n.126C>T	Het	0.0001178
F691	1A, M, F (child)	White European	g.8076762G>A	n.*9C>T	Het	0.001923
THWF		(British)	g.8076849T>C	n.58A>G	Het	0.00002277
F730 PTR	1A, M (child)	White European	g.8076761C>A	n.*10G>T	Het	0.002495
		(Australian)	g.8076826C>T	n.81G>A	Het	0.00004740
F766 EW	1A, M, F (12y)	White European	g.8076865C>T	n.42G>A	Het	0.001070
		(British)	g.8076904G>T	n.3C>A	Het	0.000004327
F780 KP	2A, 1U, M, F	White European	g.8076770G>A	n.*1C>T	Het	0.005054
	(child)	(North American)	g.8076776G>C	n.131C>G	Het	0.00001520

F819 JAM	2A, M (adolescent)	White European (British)	g.8076762G>A g.8076770G>A g.8076696_8076977del§	n.*9C>T n.*1C>T	Het Het Het	0.001923 0.005054 Novel
<mark>F906</mark> GG	1A, M, F (child)	White European	g.8076804C>T	n.103G>A (de novo)	Het	0.00001724
		(Italian)	g.8076868C>G	n.39G>C	Het	0.0001253
F1127 PG	1A, M, F (child)	White European	g.8076762G>A	n.*9C>T	Het	0.001923
		(French)	g.8076807A>C	n.100T>G	Het	Novel
F1172	1A (50y)	White European	g.8076762G>A	n.*9C>T	Het	0.001923
VVE		(German)	g.8076776G>C	n.131C>G	Het	0.00001520
F1288 LD_0807	1A, M, F (child)	White European (North American)	g.8076848A>C g.8076762G>A	n.59T>G n.*9C>T	Het Het	Novel 0.001923
F1424	1A, M, F (child)	Mixed white European (German) / north African	g.8076777A>G g.8076912C>T	n.130T>C n6G>A	Het Het	0.000008628 0.0008539
F1445	1A (child)	White European (British)	g.8076904G>A g.8076826C>T	n.3C>T n.81G>A	Het Het	0.001482 0.00004740

[#]Also reported in McNeill N et al. Neurol Genet 14: e162 (2017)

Bolded mutations proposed to be hypomorphic – individuals highlighted in green lack mutations in any of the affected seven nucleotides located in the novel base-pairing region between the 5´ end and 3´ extension.

Nomenclature in gnomAD does not correspond to nomenclature in Alamut

Het = heterozygous; Hom = homozygous. gnomAD = Genome Aggregation Database

[§] Deletion extends beyond these boundaries, but boundaries have not been fully defined. One sib carries g.8076762G>A plus g.8076696_8076977del. The other sib carries g.8076770G>A plus g.8076696_8076977del. Mum carries g.8076762G>A plus g.8076770G>A.

Supplementary Table S2 Oligonucleotides used in this study

PURPOSE	OLIGONUCLEOTIDE SEQUENCE 5' TO 3'
Sequence zebrafish U81 from genomic DNA	S ACGATACCACAATGCTGTAC
	AS GTAAGGATACACAGAATACC
Sequence zebrafish U82 from genomic DNA	S CTCCACCAATAGTATCGCAG
	AS CTGTGGTGAGAAGTAACTTTC
Sequence zebrafish U83 from genomic DNA	See Genotype Δ54U8-3 allele
Sequence zebrafish U84 from genomic DNA	S CCAATAGGATCACAGGAGAC
	AS GCATAGTAAGCTATAGTGCTC
Sequence zebrafish U85 from genomic DNA	S GGAGCCATTAGCATGCTAAC
	AS TTTCTAGCACACCCTGCTG
Quantitative RT-PCR to U8-1	S GGTTTATCCTTACCTGTTGTC
	AS CAGACAGGATTCATAAAGGTC
Quantitative RT-PCR to U8-2	S ACATCAGTGAGGTTTGTC
	AS CACATAGGATCCAGAGTTGTG
Quantitative RT-PCR to U8-3	S GAGGTACATCCTTACCTGTTAC
	AS GATTCGTAAAGGGGTTGCAG
Quantitative RT-PCR to U8-4	S GGTTTATCCTTACCTGTTGTC
	AS TAAAGAGGTTGCAAGTCATG
Quantitative RT-PCR to U8-5	S GCATCAGTGAGGTTTATCCTTG
	AS CAAGCCCTATATGAACACAGAG
DNA template for production of guide RNA to	S TAATACGACTCACTATAGGCGTCGTTTAAAAAGCATGTGGTTTTAGAGC
cleave 5' of U8-3	AS AAAAGCACCGACTCGGTGCCACTTTTTCAAG
DNA template for production of guide RNA to	S TAATACGACTCACTATAGGCGCCGGCGTAGAGACGAGGGTTTTAGAGC
cleave 3' of U8-3	AS AAAAGCACCGACTCGGTGCCACTTTTTCAAG
DNA template for production of guide RNA specific	S TAATACGACTCACTATAGATCTGTAACCTTATTGCTAGTTTTAGAGC
to U8-3	AS AAAAGCACCGACTCGGTGCCACTTTTTCAAG
Genotype ΔU8-3 allele	S1 ACCCTTGACAGAGGAGTTTG
	S2 TCCAATCCAACCAATGAGAC
	AS CATGCAGAAATGTCTCACTATC
Genotype Δ54U8-3 allele	S TAAACGATCGTCCAC
	AS GGAATGGAGTCACAGACTTAC
DNA template for production of zebrafish U8-3	S TAATACGACTCACTATAGGGGACATCAGTGAGGTACATCC
snoRNA	AS GTCAGACAGGATTCGTAAAG
DNA template for production of zebrafish pre-U8-3	S TAATACGACTCACTATAGGGGACATCAGTGAGGTACATCC
snoRNA	AS GGTTATTAAAATGTGTCAGACAGGATTC

DNA template for production of human U8 snoRNA	S TAATACGACTCACTATAGGGGATCGTCAGGTGGGATAATCC AS AATCAGACAGGAGCAATCAGGGTGTTGCAAG
DNA template for production of human pre-U8 snoRNA	S TAATACGACTCACTATAGGGGATCGTCAGGTGGGATAATCC AS ACAAATGTAAGTGATCGTCAG
DNA template for production of human pre-U8 snoRNA 57G>A	S TAATACGACTCACTATAGGGGATCGTCAGGTGGGATAATCCTTACCTGTTCCTCCCG GAGGGCAGATTAGAACATAATGATTGGAGATGCATG AS ACAAATGTAAGTGATCGTCAG
DNA template for production of human pre-U8 snoRNA 58A>G	S TAATACGACTCACTATAGGGGATCGTCAGGTGGGATAATCCTTACCTGTTCCTCCCG GAGGGCAGATTAGAACATGGTGATTGGAGATGCATG AS ACAAATGTAAGTGATCGTCAG
DNA template for production of human pre-U8 snoRNA 61A>G	S TAATACGACTCACTATAGGGGATCGTCAGGTGGGATAATCCTTACCTGTTCCTCCCG GAGGGCAGATTAGAACATGATGGTTGGAGATGCATGAAAC AS ACAAATGTAAGTGATCGTCAG
DNA template for production of human pre-U8 snoRNA *1C>T	S TAATACGACTCACTATAGGGGATCGTCAGGTGGGATAATCC AS ACAAATGTAAGTGATCGTCAGAAAAAATCAGACAGGAG
DNA template for production of human pre-U8 snoRNA *5C>G	S TAATACGACTCACTATAGGGGATCGTCAGGTGGGATAATCC AS ACAAATGTAAGTGATCGTCACAAAGAATCAGACAGGAG
DNA template for production of human pre-U8 snoRNA *9C>T	S TAATACGACTCACTATAGGGGATCGTCAGGTGGGATAATCC AS ACAAATGTAAGTGATCATCAGAAAGAATCAGACAGGAG
DNA template for production of human pre-U8 snoRNA 2T>C	S TAATACGACTCACTATAGGGGACCGTCAGGTGGGATAATCCTTACC AS ACAAATGTAAGTGATCGTCAG
DNA template for production of human pre-U8 snoRNA 3C>T	S TAATACGACTCACTATAGGGGATTGTCAGGTGGGATAATCC AS ACAAATGTAAGTGATCGTCAG
DNA template for production of human pre-U8 snoRNA 8G>C	S TAATACGACTCACTATAGGGGATCGTCACGTGGGATAATCCTTACC AS ACAAATGTAAGTGATCGTCAG
DNA template for production of human pre-U8 snoRNA *10G>T	S TAATACGACTCACTATAGGGGATCGTCAGGTGGGATAATCC AS ACAAATGTAAGTGATAGTCAGAAAGAATCAGACAGGAG
Amplification of 113p53 promoter	S GACAGACTCGAGTAGCTCGTTGGTCTGACATC AS GACAGAACTAGTAGTCTGCAGTTTGTGCTGTG
Quantitative RT-PCR to zebrafish Δ113p53	S ATATCCTGGCGAACATTTGG AS ACGTCCACCACTTTGAAC
Quantitative RT-PCR to zebrafish mdm2	S TGACAACGAGAAACTGGTAAGA AS AAACATAACCTCCTTCATGGT
Quantitative RT-PCR to zebrafish p21	S AGCTGCATTCGTCTCGTAGC AS CGGTTGAAATAAAAACGGAATA
Quantitative RT-PCR to zebrafish cyclinG1	S GTGCGGAGACGTTTTCCTT AS AAGACAGATGCTTGGGCTGA

Quantitative RT-PCR to zebrafish bax	S GGAGGCGATACGGGCAGTG
	AS TCGGCTGAAGATTAGAGTTGTTT
Quantitative RT-PCR to zebrafish elf1a	S CTGGAGGCCAGCTCAAAC
	AS ATCAAGAAGAGTACCGCTAGCATTAC
Genotyping of the M214K substitution in p53 mutant	S1 CCATGTAGTGAAGTATAGTTGC
zebrafish. The p53 locus is amplified with S1-AS1	AS1 GTCGGGTCTTCAGTTTTATGC
for 35 cycles. 1ul of the PCR product is used as a	S2 TGTAAAACGACGCCAGTGAATTATATCAA
DNA template for amplification with S2-AS2.	AS2 AGGAAACAGCTATGACCATCCAATGGCATG
Sequencing is performed with M13	M13 CAGGAAACAGCTATGAC
Probe sequence for human 3'-ETS - LD2612	GAGGAGGCGGAACCGAAGAAGCGG

Supplementary Figures

	28S binding
SNORD118	ATCGTCAGGTGGGATAATCCTTACCTGTTCCTCCTCCGGAGGGCCAGAT
U8-1	G-CATCAGTGAGGTTTATCCTTACCTGTTG-TCTT-A-GAAATAAGACTA
U8-2	A-CATCAGTGAGGTTTGTCC-TACCTGTTTT-A-GAAATAAGACTT
U8-3	A-CATCAGTGAGGTATATCCTTACCTGTTA-CCTT-A-GCAATAAGGTTA
U8-4	GT-ATCAGTGAGGTTTATCCTTACCTGTTG-TCTT-A-GAAATAAGACTG
U8-5	G-CATCAGTGAGGTTTATCCTTGCCTGTTG-TCTT-A-GAATTAAGACTT
	*** ** ** **** * * *
	C box LSm binding motif
SNORD118	TAGAACATGATGATTGGAGATGCATGAA-ACGTGATTAACGTCTCT
U8-1	CAGCACATGATGATTGGAT-TCTATGAATGCGTGATTATATTCTGT
U8-2	CAGCACATGATGATTGGAT-TCTATGAATATGTGATTATATTCTGT
U8-3	CAGATCATGATGATTGGAT-TCTATGAATACGTGATTATGTTCCGT
U8-4	CAGTATATGATGATTGGAT-TCTATGAATATGTGACTATGTTTGTACTGT
U8-5	CAGCACATGATGATTGGAT-TCTATGAATACATGATTATACTCTGT
	** ******* * **** * * * *
	D box
SNORD118	GCGTAATCAGGACTTGCAACACCCTGATTGCTCCTGTCTGATT
U8-1	ATTCTTAAAGGGTTTGCAACACCTTTATGAATCCTGTCTGA-T
U8-2	ATTCGTATAGGGTTCACAACTCTGGATCCTATGTGA-T
U8-3	ATTTGTTTAGGGACTGCAACCCCTTTACGAATCCTGTCTGA-C
U8-4	ATTTGTACATGACTTGCAACCTCTTTATAAATCCTATCTGA-T
U8-5	GTTCATATAGGGCTTGTCACCCATTTATGAATCCTGT <mark>CTGA=</mark> T
	* * * * * * * * * * * * * * * * * * * *

Fig. S1. Human U8 and zebrafish U8 1-5 mature sequences exhibit extensive conservation Mature human U8 and zebrafish U8 1-5 sequences were aligned using ClustalW (https://www.ebi.ac.uk/Tools/msa/tcoffee/). * denotes conserved nucleotides.

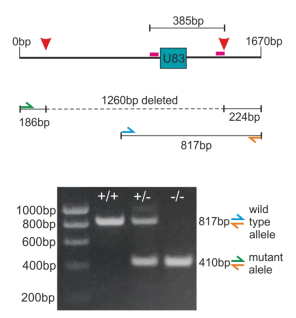


Fig. S2. The $\Delta \text{U8-3}$ mutant is homozygous for a 1.26kb deletion encompassing the U8-3 gene locus

Schematic depicting the genotyping strategy for the zebrafish U8-3 mutant. Two sgRNAs predicted to excise 385bp encompassing the U8-3 gene are represented by pink bars. Red arrowheads depict actual excision sites that delete 1260bp (dashed line) encompassing the U8-3 gene locus. A three primer PCR was performed to genotype whereby 817bp (blue, orange primer) is amplified in the presence of a wildtype copy of U8-3 and 410bp (green, orange primer) amplified in the presence of a deleted $\Delta U8-3$ allele. bp, base pairs.

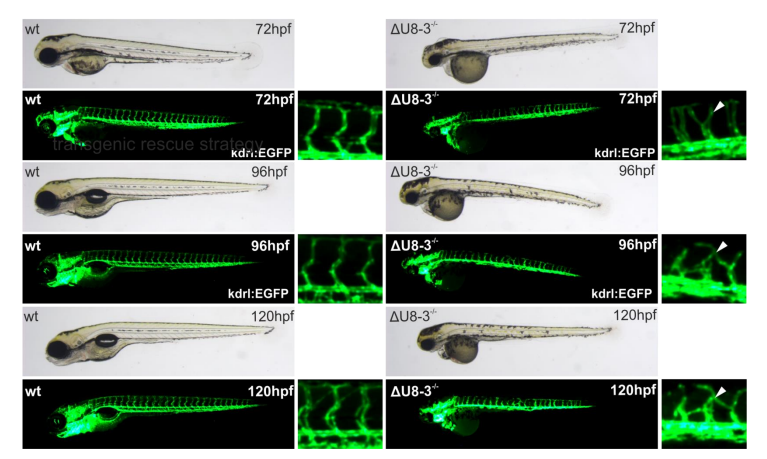


Fig. S3. The $\Delta \text{U8-3}$ mutant exhibits a range of gross morphological abnormalities and comes to a developmental standstill

From 72hpf Δ U8-3 mutants display impaired yolk resorption, reduced craniofacial structures and hindbrain swelling. By 96hpf Δ U8-3 mutants fail to inflate their swim bladders, display an underdeveloped intestine, and by 120hpf cardiac oedema. At all stages disorganized trunk vasculature is observed (white arrowheads) when compared to wildtype.

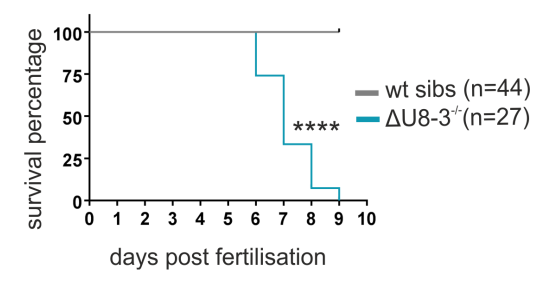


Fig. S4. The Δ U8-3 mutant phenotype is lethal Kaplain-Meier plot demonstrating that Δ U8-3 mutants begin to die from 6dpf and exhibit 100% mortality by 9dpf. dpf- days post fertilization.

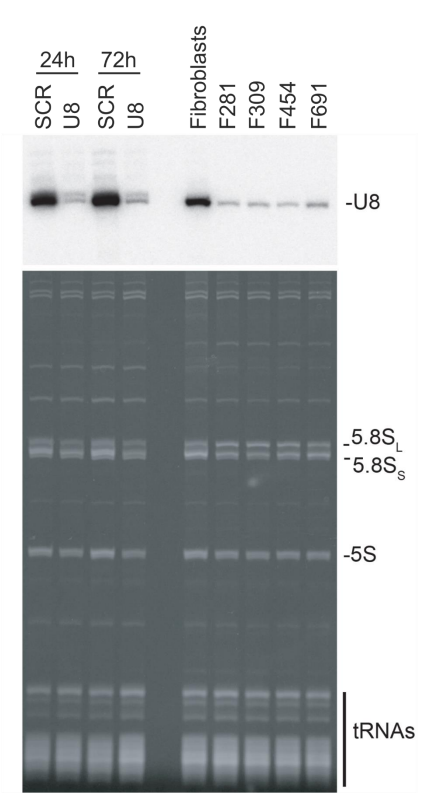


Fig. S5. LCC patient fibroblasts express less total U8 compared to wildtype fibroblasts Northern blot analysis with a probe specific to U8 snoRNA. U8 expression is shown after ASO-mediated depletion of human U8 snoRNA from HCT116 cells in which rRNA processing was analyzed (see Figure 2E), or in LCC patient fibroblast cell lines F281, F306, F454 and F691. ASO – antisense oligonucleotide. The same amount of total RNA (OD₂₆₀) was loaded in each lane of the gel. Ethidium bromide staining of the acrylamide gel demonstrates equal loading, as determined by tRNAs which are produced independently.

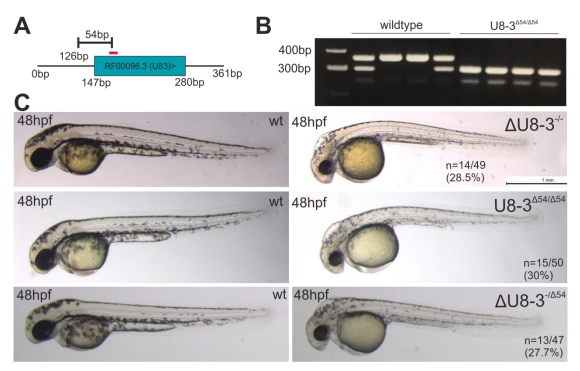


Fig. S6. Non-complementation with independent mutant alleles of $\it U8-3$ demonstrates that the $\it \Delta U8-3$ mutant is specific

- **A**) Schematic depicting the 54bp deletion allele (Δ 54U8-3) generated with a sgRNA (pink bar) from the sgRNAs used to excise *U8-3* shown in Figure 2A.
- **B**) Genotyping demonstrates the 54bp deletion (307bp band) segregates with the U8-3 mutant phenotype.
- **C**) Both ΔU8-3 and Δ54U8-3 alleles display mendelian recessive inheritance i.e. approximately 25% of mutant progeny are produced from heterozygous carriers. When heterozygous carriers of each allele were crossed to each other non-complementation (failure to produce 100% wildtype progeny) was observed, demonstrating the alleles result from loss-of-function in the same gene.

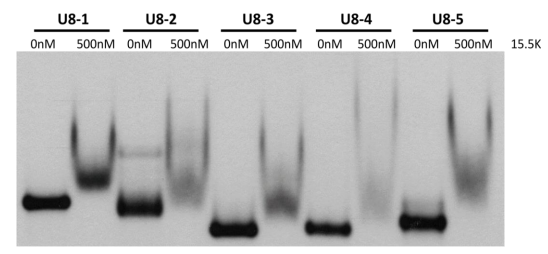


Fig. S7. EMSAs demonstrates all five zebrafish U8s bind 15.5K

Protein binding of zebrafish U8 variants. Electrophoretic mobility shift assays (EMSAs) using 5'-

end-radiolabeled *in vitro* transcribed mature zebrafish U8-1, U8-2, U8-3, U8-4 and U8-5 with 500nM of His₆-tagged 15.5K human protein (15.5K). Binding of 15.5K to zebrafish U8 is indicated by a mobility shift.

3'BOX
3' processed region GTYY (N2-5) AAARRYAGA
human U8CTTTCTGACGATCACTTACATTTGTGTTATGCTGATTAGCAGA
zebrafish U8-3ACAT TTT AATAACC GTTT CAAAATCAGA

Fig. S8. Alignment of human U8 and zebrafish 3´ extension sequencesHuman and putative zebrafish 3´ extension sequences (blue) are divergent. A 3´ BOX is predicted for zebrafish U8-3 (pink) according to the 3´ BOX consensus sequence (black).

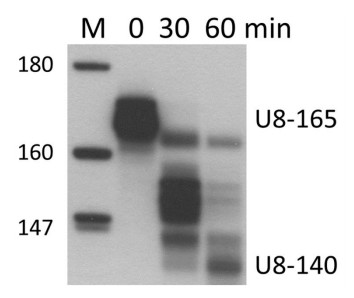


Fig. S9. Time course of wild type human pre-U8 snoRNA processing.Processing of 5´ end-radiolabelled *in vitro* transcribed precursor U8 wildtype snoRNA (U8-165) was assessed in HeLa nuclear extracts at 0, 30 and 60 minutes (min) to monitor production of mature U8 (U8-140). Mature U8 is clearly present after 60 min of processing.

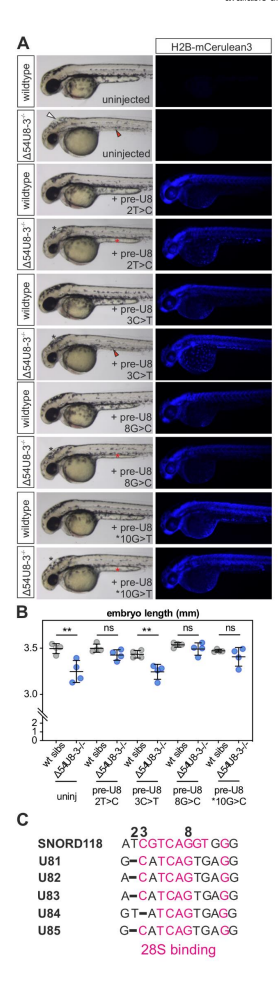


Fig. S10. Functional testing of four LCC patient mutant U8 snoRNAs in zebrafish

- A) Representative images of rescue experiments conducted with four putatively hypomorphic U8 mutations detected in LCC patients. White arrowhead indicates hindbrain swelling and red arrowheads indicate abnormal yolk extension. Black and red asterisks indicate rescued hindbrain swelling and yolk extension respectively
- **B**) pre- $U8^{3C>T}$ does not rescue the reduced embryo length of $\Delta U8-3$ mutants, whereas pre- $U8^{2T>C}$, pre- $U8^{8G>C}$ and pre- $U8^{*10G>T}$ restores $\Delta U8-3$ mutant embryo length to that of wildtype siblings. n=4 embryos per genotype
- **C**) alignment of human U8 and zebrafish U8 28S interaction region. Shared identity between zebrafish and human 28S-binding nucleotides is shown in pink

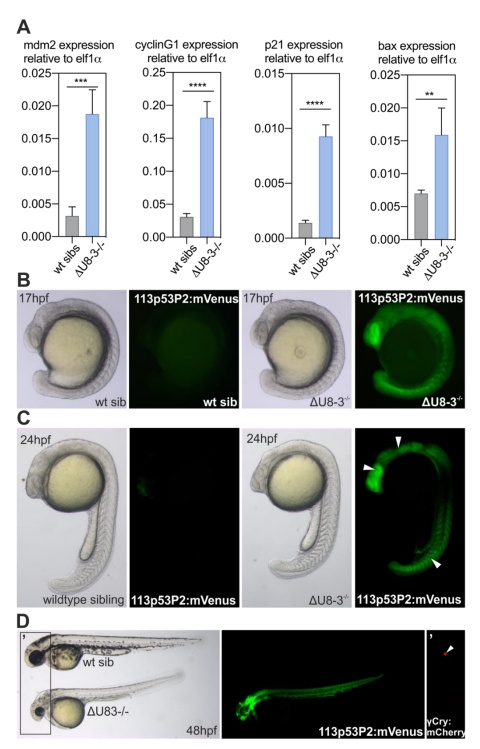


Fig. S11. tp53 is trans-activated in response to loss of U8-3 in zebrafish

- A) quantitative RT-PCR demonstrates significant upregulation of the tp53 target genes *mdm*2, *cyclinG1*, *p21* and *bax* in ΔU8-3 mutants when compared to wildtype siblings at 24hpf.
- **B**) tp53 activity is observed throughout the U8-3 mutant at 17hpf, prior to the onset of observable gross morphological abnormalities, particularly in the eye.
- \mathbf{C}) tp53 activity is observed in numerous tissues that develop abnormally in $\Delta U8-3$ mutants, including the eye, hind-brain and somites (arrowheads) at 24hpf.
- **D**) tp53 activity is observed throughout the Δ U8-3 mutant at 48hpf. Integrated 113p53P2:mVenus transgene is reported by γ -crystallin:mCherry in the backbone of the plasmid which drives lens specific mCherry expression from approximately 36hpf. The developmental delay in Δ U8-3 mutants delays expression of this reporter.

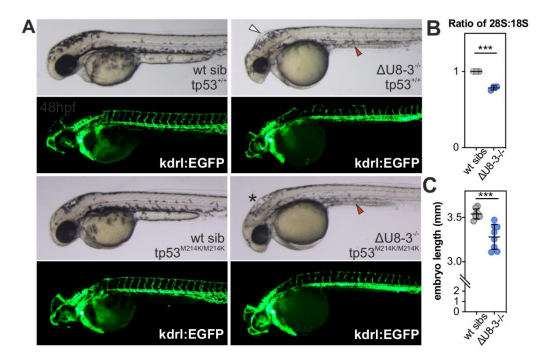


Fig. S12. Inactivation of tp53 partially rescues the Δ U8-3 mutant phenotype though fails to rescue the embryo length or rRNA biogenesis defect

- **A**) representative images showing inactivation of tp53 signaling rescues the hindbrain swelling of Δ U8-3 mutants (white arrowhead compared to black asterisk) and the trunk vasculature, but not the yolk extension (red arrowhead) abnormality.
- **B**) Tapestation analysis demonstrates tp53 mutant Δ U8-3 mutant embryos display a preferential reduction in 28S biogenesis at 48hpf. n=3 biological replicates per genotype.
- **C**) tp53 mutant U83 mutant embryos are significantly shorter than tp53 mutant U8-3 wildtype sibling embryos at 48hpf. n=8 biological replicates per genotype.