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A mouse model for intellectual disability caused by mutations in the X-linked 2'-O-methyltransferase Ftsj1 gene.

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Abstract

Mutations in the X chromosomal tRNA 2'-O-methyltransferase FTSJ1 cause intellectual disability (ID). Although the gene is ubiquitously expressed affected individuals present no consistent clinical features beyond ID. In order to study the pathological mechanism involved in the aetiology of FTSJ1 deficiency-related cognitive impairment, we generated and characterized an Ftsj1 deficient mouse line based on the gene trapped stem cell line RRD143. Apart from an impaired learning capacity these mice presented with several statistically significantly altered features related to behaviour, pain sensing, bone and energy
metabolism, the immune and the hormone system as well as gene expression. These findings show that Ftsj1 deficiency in mammals is not phenotypically restricted to the brain but affects various organ systems. Re-examination of ID patients with FTSJ1 mutations from two previously reported families showed that several features observed in the mouse model were recapitulated in some of the patients. Though the clinical spectrum related to Ftsj1 deficiency in mouse and man is variable, we suggest that an increased pain threshold may be more common in patients with FTSJ1 deficiency. Our findings demonstrate novel roles for Ftsj1 in maintaining proper cellular and tissue functions in a mammalian organism.

Keywords
Ftsj1, tRNA methyltransferase, intellectual disability, X-linked, mouse model

1. Introduction
Intellectual disability (ID) is defined by significant limitations in intellectual performance (intelligence quotient <70) and a reduction of conceptual, social and practical skills with an onset before the age of 18. ID is a major developmental disorder and currently ID is estimated to affect around 1% of the general population, but with large regional differences (1). Historically ID was separated into syndromic and non-syndromic forms, with the former being ID in the presence of clinically consistent features. The non-syndromic forms seem more common, but as more patients with similar gene defects are molecularly diagnosed syndromic features may become apparent.

Although ID has genetic as well as environmental causes (like excessive alcohol consumption during pregnancy), severe forms are often caused by genetic defects in single genes (2–4).

Monogenic recessive causes of ID have been excessively studied in X-linked forms and with the development of next-generation-sequencing techniques the elucidation of autosomal forms has also made a leap forward. Mutation discovery studies so far have clearly shown that ID is genetically and functionally heterogeneous but also that several ID proteins interact, are involved in the same process or target the same substrate. One emerging “ID gene group” is involved in tRNA modifications and includes NSUN2 (5–7), ADAT3 (8), TRMT10A (9,10), TRIT1 (11), TRMT1 (3), ELP2 (3,12), PUS3 (13), MTO1 (14) and FTSJ1 (15–18).

ID-causing mutations affecting FTSJ1 are mainly loss-of-function variants that prematurely truncate the protein by affecting splicing or directly introduce stop-codons, while more recently an FTSJ1 missense mutation was detected in an ID patient (19).

FTSJ1 encodes a ubiquitously expressed and evolutionarily highly conserved 2’-O-methyltransferase that uses S-adenosylmethionine as methyl donor (20). Mammalian FTSJ1
is not as well characterized as its yeast homologue, but the function is highly conserved as human FTSJ1 is able to methylate yeast tRNA\textsuperscript{Phe} at position 32 and 34 in yeast cells (21). 2'-O-methylation of tRNA\textsuperscript{Phe} also appears to be its major target (21) and yeast cells deficient in the Ftsj1 homologue Trm7 grow slowly and show reduced mRNA translation (20). Trm7 methylates tRNA\textsuperscript{Phe}, tRNA\textsuperscript{Trp} and tRNA\textsuperscript{Leu(UAA)} at position C\textsubscript{32} and N\textsubscript{34} (20) using two different protein interaction partners, TRM732 and TRM734, which are required for each modification (20–22). Both modifications are required for the subsequent conversion of 1-methylguanosine at position 37 to wyebutosine in tRNA\textsuperscript{Phe} (22).

Still, mammalian FTSJ1 is largely uncharacterized and the consequences of FTSJ1 deficiency in human seem to mainly affect cognition. Therefore, in order to study the effect of FTSJ1 deficiency not only on brain function but also on other organs, we set out to generate Ftsj1 deficient mice. Here we report the results of a thorough characterization of our mouse model, showing that in addition to alterations in learning and memory other features including nociception, metabolism, hormone levels and the immune system are affected, some of which could also be found in human carriers of deleterious FTSJ1 mutations that are affected with ID.

2. Materials and methods

2.1 Generation of Ftsj1 deficient mice

Ftsj1 gene trapped mice were generated by microinjection of gene trapped ES-cells. The mouse embryonic stem cell line RRD143 (129P2/OlaHsd) containing a gene trap insertion in intron 5 of the Ftsj1 gene was obtained from Baygenomics and expanded on a layer of cell division deficient mouse embryonic feeder cells. Blastocysts were obtained from hormonally stimulated (superovulating) female C57BL/6J mice. The round and smooth ES-cells were injected into the blastocyst cavity and surgically transferred into pseudo-pregnant recipient female mice using standard methods, essentially as described (49). Chimeric pups were identified by the presence of brown coat spots and bred to select for germline transmission. Offspring harboring gene trapped Ftsj1 were then backcrossed to a C57BL/6J genetic background for at least 10 generations before they were expanded to provide wild type and gene trapped littermates of both sexes for phenotypic characterization. In this process we also removed other gene traps that could have been integrated in the genomic DNA during the RRD143 cell line generation. For the mouse characterization at the German Mouse Clinic (www.mouseclinic.de) a cohort of 54 animals were used. 12 Ftsj1\textsuperscript{+/y} males, 15 control males, 12 Ftsj1\textsuperscript{-/-} females and 15 Ftsj1\textsuperscript{-/+} control female mice were born within one week from the mating of hemizygous Ftsj1 gene trapped males with heterozygous females.

All tests performed at the GMC were approved by the responsible authority of the district government of Upper Bavaria, Germany. The hole-board test was carried out according to
the guidelines of the Landesamt für Landwirtschaft, Lebensmittelsicherheit und Fischerei (LALLF) Mecklenburg-Vorpommern, Germany.

2.2 RT-PCR and PCR
Total RNA was isolated from cultured cells or mouse tissues using TRIzol (Life Technologies, Carlsbad, CA, USA) according to the manufacturer’s instructions. RNA integrity was verified on ethidiumbromide stained agarose gels. For cDNA synthesis 1 µg total RNA was reverse transcribed using random priming and Superscript III (Life Technologies, Carlsbad, CA, USA) according to the manufacturer’s instructions. cDNA corresponding to 20 ng total RNA was amplified using the GoTaq G2 Colorless Master Mix (Promega, Madison, WI, USA). For RT-PCR we used the primers: FTSJ1_5UTR_U 5’-ATACGTTGTGCTCCGGTCTG-3’ and FTSJ1_ex7-8L 5’-ATATCTTGGCCACAGAAGCAG-3’ for detecting a 705 bp wild type Ftsj1 PCR product and the primers mm_FTSJ1_GT1u1 5’-GTGCTGAGCCAGAAGGTGGT-3’ and mm_FTSJ1_GT1l1 5’-GGCGATTAAGTTGGGTAACG-3’ for detecting a 352 bp PCR product containing a part of Ftsj1 and the gene trap sequence. As ladder the pUC Mix Marker 8 (MBI Fermentas #SM 0302, St Leon-Rot, Germany) was used.

2.3 Genotyping by PCR
Mouse genomic DNA was isolated from tail or ear tissue using the NucleoSpin Tissue kit (Macherey & Nagel, Düren, Germany) following the manufacturer’s instructions. The DNA concentration was measured with a QuBit fluorometer (Life Technologies, Carlsbad, CA, USA). 20 ng DNA was amplified by PCR with primer combinations specific for the wild type Ftsj1 or for the gene trapped Ftsj1, respectively. One oligonucleotide was used for both PCRs mm_FTSJ1_GT1u3 5’-TCTGACCACACGTGTTGTTG-3’ in combination with Ftsj1_in5_1l3 5’-AAGAAGAGTGTACGATCTCCTG-3’ to obtain a 531 bp product for the wild type sequence, and oligonucleotide pGT1lx_4l1 5’-AGGGAAAGGTAAAGTGTTG-3’ for detecting the gene trap in a PCR product of 240 bp.

2.4 Phenotyping
At the German Mouse Clinic (GMC) the mice were subjected to an extensive phenotypic screening, including standardized phenotyping in the areas of dysmorphology, behaviour, neurology, nociception, energy metabolism, clinical chemistry, cardiovascular features, eye, lung function, molecular features, immunology, steroid metabolism and pathology (50,51) (see also www.mouseclinic.de). The phenotypic tests were part of the GMC screen, which were performed according to standardized protocols as described (52,53). The experiments were not in general blinded. However, the preparation and organization of the experimental procedures at the German Mouse Clinic are adapted in a tailor-made way for each single test according to the specific requirements. When applicable, tests and analyses were carried out
by an automated system where the experimenter had no influence on the outcome of the test (e.g. acoustic startle and prepulse inhibition, open field test, blood based parameters and DEXA analysis). When tests were carried out in parallel with a subset of the animals (e.g. indirect calorimetry, open field test, acoustic startle and prepulse inhibition), mutant and control animals were distributed either in a random or in a balanced way between the different apparatuses. Meta data for experimental conditions were recorded and analyzed for deviations.

The hole-board test was performed as described (54) using mice from three litters in total consisting of 6 gene trapped and 7 control male mice.

2.5 Statistical analyses
About 500 parameters belonging to 14 different disease areas were measured during the primary GMC screen. As level of significance for statistical analyses of GMC-screening results we used a p-value <0.05 (without correction for multiple testing). If not stated otherwise, data generated by the German Mouse Clinic were analyzed using R. Tests for genotype effects were made by using t-test, Wilcoxon rank sum test, linear models, or ANOVA depending on the assumed distribution of the parameter and the questions addressed to the data.

2.6 mRNA Expression profiling
Brain tissue was obtained from mice from 3 litter pairs that were 7, 9 and 14 months old, each consisting of one wt male and one hemizygous gene trapped male. We chose to use littermates of different age because the effects of FTSJ1 deficiency seem to be constant over time in human patients as no improvement in cognition has been reported. By sequencing and analyzing data from the mice separately we obtain information on expression differences between gene trapped and wildtype mice at different age, while still having the option to pool data from gene trapped and control mice, respectively. The tissue was homogenized using a polytron device (VWR) in the presence of TRIzol reagent (Life Technologies, Carlsbad, CA, USA) according to the instructions of the manufacturer. Next, we removed the rRNA from the RNA samples using a RiboZero Kit (Epicentre, Madison, WI, USA) according to the manufacturers’ instructions and spiked the RNA with ERCC RNA Spike-in Mix (Life Technologies, Carlsbad, CA, USA). The RNA was fragmented by chemical hydrolysis, phosphorylated and purified. Adaptors were then ligated to the RNA fragments and reverse-transcribed into cDNA. The cDNA was then further processed to sequencing libraries using recommended protocols and chemicals, which were then paired-end sequenced (75 nucleotides in the forward direction and 35 nucleotides in the reverse direction) using a SOLiD 5500xl Genetic Analyzer (Life Technologies, Carlsbad, CA, USA). For mapping of the
sequence reads we used the GRCm38/mm10 assembly sequence build (ftp://hgdownload.cse.ucsc.edu/goldenPath/mm10/) as a reference and followed the workflow 'whole.transcriptome.pe' of LifeScope-v2.5.1-r0 (Life Technologies, Carlsbad, USA). Of these we considered sequence reads that mapped to RefSeq coding exons (http://hgdownload.cse.ucsc.edu/goldenPath/mm10/database/refGene.txt.gz) and matched the coding strand to be coding RNAs. All other reads were considered to be non-coding. Differential expression analysis was performed as previously described (55). In brief, reads were normalized for GC-content using EDASeq’s full quantile normalization (56) and the differential expression analysis was carried out using DESeq (57). All data were treated as replicates of a single condition. The significance of the differential expression values was determined using Benjamini-Hochberg corrected p-values between two conditions (control or gene trapped mice) with the significance threshold set to 0.05.

2.7 Patient reexamination
Our studies of the molecular background of cognitive disorders are approved by the ethical committee of the University Medicine Greifswald. Clinical examinations were performed with the consent of the affected Individuals and/or their legal guardians. All methods were carried out in accordance with relevant guidelines and regulations.

3. Results

3.1 Validation of the gene trap in cells and mice
By PCR amplification of genomic DNA extracted from the mouse embryonic stem cell line RRD143 and subsequent Sanger sequencing of the PCR products we identified the exact location of the gene trap in intron 5 of the Ftsj1 gene (ChrX: 7,825,518 and 7,825,539) corresponding to position c.361+566 to c.361+587 at the cDNA level (mouse NCBI37/mm9). After generating the gene trapped mice we extracted RNA from brain, spleen and kidney and could show by RT-PCR that the gene trap is part of the Ftsj1 transcript, effectively generating an Ftsj1 deficient mouse line as no wild-type transcript was visible in these tissues (Fig. 1). As human female heterozygous FTSJ1 mutation carriers are affected to a much lesser extent than male hemizygous carriers, we chose to compare homozygously gene trapped female mice with heterozygous female control mice.
Figure 1

3.2 Morphological features, body composition and energy metabolism

As expected for a mouse model for non-syndromic intellectual disability we did not observe genotype specific anatomical differences or differences in organ weight when normalized to body weight. Also no genotype effect on tibia length was observed and there was no association between genotype and the occurrence of anterior segment abnormalities, axial eye length, or altered visual properties. However, our preliminary observation that *Ftsj1* deficient mice are lighter (p<0.001 for males and females) and shorter (p<0.001 for males and p=0.017 for females) was confirmed (Table 1). In addition, we found that fat mass and
lean mass were significantly decreased in male gene trapped mice even though food intake was comparable for all genotypes. Using Dual-energy X-ray absorptiometry (DEXA) we found that relative body fat content was increased in gene trapped male mice whereas the lean mass was lower (Table 1), a difference that was attenuated over time (data not shown). Bone mineral content (BMC) and bone content (% of total weight) was significantly reduced in the gene trapped mice (p<0.001) but without affecting bone mineral density (BMD, Table 1). Energy turnover and body temperature was not significantly changed and lactate levels and lactate dehydrogenase activity were also within the normal range (Supplementary Fig. 1).

Table 1. Gene trapped mice are small and have reduced body weight and bone content.

|                           | female control N=14 | female mutant N=8 | male control N=16 | male mutant N=13 | pairwise (Tukey) females mutant-control adj. p-value | pairwise (Tukey) males mutant-control adj. p-value | ANOVA genotype p-value | ANOVA sex p-value | ANOVA sex: genotype p-value |
|---------------------------|---------------------|-------------------|-------------------|------------------|------------------------------------------------------|---------------------------------------------------|------------------------|--------------------------|-----------------------------|
| BMD* [mg/cm^2]           | 47 ± 4              | 49 ± 5            | 51 ± 4            | 50 ± 4           | 0.457                                                | 0.882                                             | 0.698                  | 0.013                    | 0.115                       |
| BMC** [mg]               | 308 ± 53            | 156 ± 48          | 378 ± 125         | 155 ± 70         | 0.001                                                | < 0.001                                           | < 0.001                | < 0.292                   | 0.158                       |
| Bone content [%]         | 1.29 ± 0.18         | 0.78 ± 0.23       | 1.25 ± 0.38       | 0.70 ± 0.30      | 0.001                                                | < 0.001                                           | < 0.001                | < 0.216                   | 0.777                       |
| Body length [cm]         | 9.41 ± 0.21         | 9.11 ± 0.31       | 9.61 ± 0.19       | 9.22 ± 0.18      | 0.017                                                | < 0.001                                           | < 0.001                | 0.027                    | 0.449                       |
| Body weight [g]          | 23.83 ± 1.49        | 19.93 ± 1.62      | 30.7 ± 2.22       | 22.17 ± 1.44     | < 0.001                                              | < 0.001                                           | < 0.001                | < 0.001                   | < 0.001                     |
| Fat Mass [g]             | 1.87 ± 1.08         | 0.55 ± 0.94       | 4.73 ± 2.68       | 0.40 ± 0.98      | 0.326                                                | < 0.001                                           | < 0.001                | < 0.006                   | 0.004                       |
| Lean Mass [g]            | 17.24 ± 1.43        | 15.06 ± 1.84      | 20.14 ± 2.27      | 17.59 ± 1.66     | 0.05                                                 | 0.003                                             | < 0.001                | < 0.001                   | 0.732                       |

*BMD: Bone mass density
**BMC: Bone mass content

3.3 Altered behaviour and nociception in Ftsj1 deficient mice

To investigate behavioural changes in Ftsj1 gene trapped mice, we subjected the animals to the following tests: Open field test, Y-maze, social discrimination, prepulse inhibition/acoustic startle, place learning/reversal learning, grip strength, rotarod, hole-board and hot plate test.
In the open field test we observed a slightly decreased rearing activity (p<0.05) among the homozygous mutant females when compared to the heterozygous controls, but neither the Y-maze nor the social discrimination test showed consistent differences between mutant and control mice (Supplementary Table 1). Only Ftsj1 gene trapped males showed decreased acoustic startle activity (after body weight normalization, \( F(7,196) = 4.584, p < 0.001 \)), but no change in prepulse inhibition (Supplementary Table 1). Place and reversal learning as evaluated by using the IntelliCage system, however, was significantly impaired in male mutants (Fig. 2).

**Figure 2**

(A) Place Learning error rate for male mutant and control mice on four different consecutive days. (B) Ftsj1 gene trapped male and female mice have elevated pain threshold as measured by the (prolonged) response time to a thermal stimulus. sd: Standard deviation, ANOVA: Analysis of variance results.

**Fig. 2 Place learning, reversal learning and nociception is impaired in Ftsj1 gene trapped mice.** (A) Place learning and (B) reversal learning error rate for male mutant and control mice on four different consecutive days. (C) Ftsj1 gene trapped male and female mice have elevated pain threshold as measured by the (prolonged) response time to a thermal stimulus. sd: Standard deviation, ANOVA: Analysis of variance results.

Interestingly, heterozygous females showed a significantly increased error rate in place learning during the first and the third day, but the significance level was not reached any more during the following sessions and reversal learning did not show differences between female heterozygotes and female homozygous wild type mice (not shown). Muscle function was
assessed through grip strength tests, which showed reduced strength for the male gene trapped mice. However, after normalizing for the lower body weight observed for the gene trapped mice, only the forelimb grip strength was statistically significant (Fig. 3H). The reduced muscle strength did not affect motor coordination as the rotarod test did not show genotype specific differences (Supplementary Table 1). Using the hole-board test we observed that male gene trapped mice dipped the head through the opening more often than control mice (p=0.005) suggesting a more explorative/less anxious behaviour as compared to controls. The total distance traveled and rearing behaviour was not statistically different between the two groups (Supplementary Table 1).

Figure 3

Fig. 3 Blood glucose, metabolite levels and grip strength are reduced in Ftsj1 deficient male mice. Blood Glucose concentration is lower in gene trapped mice as compared to wild type male mice (A). Injected glucose is cleared faster from the blood in gene trapped male mice as compared to wild type male mice. Shown are area under curve values for the first 30 minutes of the test (AUC 0-30 min) (B). Alpha-amylase (C), creatinine (D), triglycerides (E), cholesterol (F), total protein concentration (G) and grip strength (H) are reduced in male gene trapped mice. Female mice show the same trend as male mice, but less pronounced. The data are presented as box plot with strip charts showing the individual animals split by sex and genotype. Female mice are shown as filled circles and males as diamonds. The box represents the 25% and 75% quartile and the line within the boxplot represents the median. The asterisk represents the mean. Circled points or diamonds denote values outside 1.5 x the interquartile range. f.con: female Ftsj1+/− mice (n=15); f.mut: female Ftsj1−/− mice (n=12); m.con: male Ftsj1+/− mice (n=16); m. mut: male Ftsj1−/− mice (n=14). Data were analyzed using a linear regression model.
We used the hot plate test to investigate the response to a thermal pain stimulus and found that gene trapped male mice as well as homozygous gene trapped females show hypoalgesia. The latency time was significantly longer for both the first (p=0.007) and the second reaction (p<0.001). There were no differences in the type of reaction to the stimuli or in hind paw shaking or hind paw licking (Fig 2).

3.4 Altered metabolism in Ftsj1 deficient mice:
We observed that the loss in body weight after overnight fasting was relatively larger for the Ftsj1 deficient males as compared to the male controls (p<0.001). The heterozygous females showed the same trend, but not as pronounced (p<0.05). We observed lower glucose levels both in ad libitum fed and fasted male gene trapped mice. In the intraperitoneal Glucose Tolerance Test (IpGTT) we furthermore found lower AUC 0-30 (Fig. 3) and AUC 30-120 values in male gene trapped mice suggesting that they were significantly better at clearing intraperitoneally injected glucose from the bloodstream (p<0.001). This indicates an improved glucose tolerance in the male gene trapped mice. Further clinical chemistry analysis revealed decreased cholesterol and triglyceride levels in fasted and ad libitum fed mutant male mice. Both HDL and non-HDL cholesterol was similarly affected in the fasted animals (not shown). In the ad libitum fed state we also found decreased total protein and creatinine levels and decreased alpha-amylase activity in gene trapped male mice (Fig. 3). We also observed a trend towards lower iron levels and higher alkaline phosphatase (ALP) activity (p=0.008, Supplementary Fig 1). Only in male gene trap carriers we saw slightly decreased sodium (p=0.04) and calcium (p=0.011, Supplementary Fig 1) levels.

3.5 Hematology, Immune System and steroid metabolism
Through a hematological analysis we observed slightly more red blood cells in gene trapped male mice (p=0.084) than in the controls. The hematocrit values were not affected but the mean corpuscular volume was significantly decreased (p<0.001) suggesting mild microcytosis. The red blood cell distribution width (RDW) was slightly higher in mutant mice (p=0.005) suggesting increased anisocytosis in these animals. The effects were more pronounced in the male mice. The mean corpuscular hemoglobin (MCH) was significantly decreased in mutant animals (p<0.001) but the cellular hemoglobin concentration (MCHC) was not significantly affected.
The platelet count (PLT) showed a significant reduction in the gene trapped mice (p=0.003) and the mean platelet volume was also reduced in these animals (p=0.019) whereas the platelet distribution width (PDW) was not affected. We also found a higher white blood cell count in gene trap carriers (p=0.01) of both sexes and measured the leukocyte subpopulation frequencies. We observed a reduced proportion of CD25 expressing CD4+ T cells in male and female mutant mice (p<0.001). Within the B cell compartment, we observed lower frequencies of IgD (female: p=0.021, male p=0.013) and MHC class II expressing cells (female: p=0.04, male: p<0.001). In male mutants we furthermore found a slight increase in granulocytes (p=0.021) and a slight decrease in B cell frequency (p=0.027). However, in the allergy screen, we found no differences in IgE levels between mutant and control mice, suggesting an absence of allergy or autoimmune responses as the level of autoantibodies was in the zero-range (not shown). Measuring IgG in blood plasma we found less IgG1 in male (p<0.001), but not in female mutant animals and less IgG3 in female (p=0.023) but not in male mutant mice. Finally, we measured the levels of three steroids in blood plasma: corticosterone, testosterone and androstenedione. Testosterone and androstenedione were within normal range but corticosterone levels were significantly increased in male and female gene trap carriers (p<0.001, Fig 4).

Fig. 4 Hematological, immunological and hormonal alterations in Ftsj1 gene trapped mice. The red blood cell mean corpuscular volume was reduced in mutant mice (A) and less equal in size (B) whereas the white blood cell count was increased in mutant mice (C). We also observed reduced proportions of CD45+CD3-CD4-CD25 expressing T-cells (D) and IgD
positive cells in gene trapped mice (E). In addition, we observed fewer B cells positive for B220 and MHC class II in the gene trapped mice (F). The plasma IgG3 levels were reduced in female homozygously gene trapped mice (G) and IgG1 levels were reduced in male gene trapped mice (H). In contrast we found elevated corticosterone levels in male and female gene trapped mice (I). The data are presented as box plot with strip charts showing the individual animals split by sex and genotype. Female mice are shown as filled circles and males as diamonds. The box represents the 25% and 75% quartile and the line within the boxplot represents the median. The asterix represents the mean. Circled points or diamonds denote values outside 1.5 x the interquartile range. F.con: female Ftsj1 +/− mice (n=15); f.mut: female Ftsj1 −/− mice (n=10); m.con: male Ftsj1 +/y mice (n=14); m. mut: male Ftsj1 −/y mice (n=12). Data were analyzed using a linear regression model.

3.6 RNA expression profiling

We extracted RNA from the whole brain of 3 male wt and 3 male gene trapped mice and prepared libraries that were sequenced on a SOLiD 5500xl machine. After read-mapping to the mouse genome (GRCm38/mm10 assembly), expression profiles were generated. We found that Ftsj1 expression in gene trapped mice was reduced to 15 to 21% of the levels observed in unaffected littermates. However, almost all of the reads in the gene trapped mice were located between the transcription start site and the gene trap (located between exon 5 and 6) whereas almost no reads were found between the gene trap and the 3’ end of the gene. This result is in line with the RT-PCR analysis (see above) and confirms that gene trapped mice do not express detectable full length Ftsj1.

We then compared expression profiles of the three gene trapped mice with that of the three control mice and obtained 50 transcripts with an adjusted p-value < 0.05 (Table 2). Of these genes 39 code for proteins, which were functionally analyzed using the DAVID 6.8 software (23,24). Based on the Gene Ontology (GO) annotation we found enrichment for genes belonging to the category “Biological processes” (BP), “Cellular compartment” (CC) and “Molecular function” (MF) (Supplementary Table 2). The two most significant entries from each category were “Plasminogen activation” (p=1.1E-6) and “Blood coagulation” (p=3.1E-5) for BP, “Blood microparticle” (p=1.3E-13) and “Extracellular space” (p=2.4E-12) for CC, and “Phospholipid binding” (1.5E-6) and “lipid binding” (2.8E-5) for MF (supplementary Table 3). A search for genes enriched in KEGG Pathway database revealed enrichment for genes belonging to the pathways “complement and coagulation cascades” (p=4.2E-2), “platelet activation” (p=9.8E-2) and “fat digestion and absorption” (p=6.7E-2) (supplementary Table 4). Of 50 differentially expressed genes 11 are not coding for proteins. Two of these encode miRNAs and eight encode snoRNAs. snoRNAs are RNAs that either mediate or are predicted to mediate site-specific 2’-O-methylation or pseudouridinylation of 18S, 28S, 5.8S
rRNAs and the U6 spliceosomal RNA (25). In addition, we found enhanced expression of Gas5, which is upregulated in nutrient-deprived cells and contains a glucocorticoid response element. Expression levels in each of the three litter mates are shown in supplementary Table 5.

Table 2. Mean RNA expression level and fold change in the brains of three Ftsj1 gene trapped and three wild type littermates

| Symbol  | Ftsj1 -/y | Ftsj1 +/y | Fold change | Adjusted p-value | Description                                                                 |
|---------|-----------|-----------|-------------|------------------|------------------------------------------------------------------------------|
| Alb     | 620.57    | 17.86     | 34.74       | 5.05E-141        | Albumin                                                                     |
| Apob    | 195.29    | 11.56     | 16.89       | 1.20E-46         | Apolipoprotein B                                                            |
| Ftsj1   | 26.96     | 152.18    | 0.18        | 7.68E-20         | Ftsj1                                                                        |
| Serpina3k| 48.65     | 0.99      | 49.21       | 1.92E-15         | Serine protease inhibitor A3K precursor                                      |
| Mup3    | 38.33     | 0.00      | Inf         | 2.04E-14         | Major urinary protein 3                                                    |
| Pzp     | 52.32     | 3.31      | 15.83       | 1.62E-12         | Pregnancy zone protein                                                      |
| Gc      | 51.31     | 3.63      | 14.12       | 9.77E-12         | Group specific component                                                    |
| Fga     | 34.65     | 0.99      | 34.86       | 5.46E-10         | Fibrinogen Gamma Chain                                                     |
| ApoA1   | 45.97     | 4.63      | 9.93        | 9.85E-09         | Apolipoprotein A-I                                                         |
| Cyp3a11 | 41.32     | 3.64      | 11.37       | 2.44E-08         | Cytochrome P450, family 3, subfamily a, polypeptide 11                       |
| Fga     | 32.65     | 1.99      | 16.44       | 1.73E-07         | Fibrinogen alpha chain                                                     |
| Gas5    | 8811.21   | 5710.96   | 1.54        | 2.00E-07         | Growth Arrest Specific 5 (Non-Protein Coding)                               |
| Snora16a| 815.61    | 474.92    | 1.72        | 2.10E-07         | Small Nucleolar RNA, H/ACA Box 16A                                         |
| Mug1    | 21.33     | 0.00      | Inf         | 2.89E-07         | Murinoglobulin 1                                                           |
| Doc2g   | 73.89     | 173.61    | 0.43        | 1.18E-06         | Double C2, gamma                                                           |
| ApoA2   | 40.99     | 5.63      | 7.28        | 2.31E-06         | Apolipoprotein A-II                                                        |
| Snord22 | 2958.36   | 1948.18   | 1.52        | 4.22E-06         | Small Nucleolar RNA, C/D Box 22                                            |
| Cps1    | 42.64     | 7.29      | 5.85        | 1.40E-05         | Carbamoyl Phosphate Synthase                                               |
| Capn11  | 30.62     | 3.31      | 9.25        | 2.69E-05         | Calpain 11                                                                  |
| Akr1c6  | 15.33     | 0.00      | Inf         | 1.09E-04         | Aldo-keto reductase family 1, member C6                                    |
| Omp     | 6.32      | 35.35     | 0.18        | 3.14E-04         | Olfactory Marker Protein                                                   |
| Slco1b2 | 26.99     | 3.30      | 8.17        | 3.99E-04         | Solute carrier organic anion transporter family, member 1b2                |
| Ahsg    | 22.66     | 1.98      | 11.43       | 4.48E-04         | Alpha-2-HS-glycoprotein                                                    |
| Fgb     | 23.99     | 2.65      | 9.05        | 8.14E-04         | Fibrinogen beta chain                                                      |
| Pigr    | 14.66     | 0.33      | 44.14       | 1.43E-03         | Polymeric Immunoglobulin Receptor                                          |
| Snord96a| 255.96    | 148.92    | 1.72        | 1.81E-03         | Small Nucleolar RNA, C/D Box 96A                                           |
| Azgp1   | 14.32     | 0.33      | 43.27       | 1.84E-03         | Alpha-2-glycoprotein 1, zinc                                               |
| Fabp1   | 21.66     | 2.32      | 9.35        | 2.00E-03         | Fatty acid binding protein 1, liver                                        |
| Snora28 | 935.71    | 644.67    | 1.45        | 2.59E-03         | Small Nucleolar RNA, H/ACA Box 28                                          |
| C3      | 67.62     | 26.50     | 2.55        | 5.56E-03         | Complement Component 3                                                    |
to cognition. Female carrier may have hematology, triglyceride and hormone alterations related to syndromic, we did not expect to find consistent alterations, but we did observe individual alterations related to cholesterol levels, muscle strength, hypoalgnesia, hematology, triglyceride and hormone levels (summarized in Table 3) in patients or the female carrier, which strongly suggest that the effects of FTSJ1 deficiency are not restricted to cognitive features in humans either.

Expression values are given in reads per kilobase per million mapped reads (RPKM)

### 3.7 Patient reexamination

Based on the phenotypic features found in Ftsj1 deficient mice the mutation carriers from two previously published families A003 and MRX9 were re-examined. As FTSJ1 related ID is categorized as non-syndromic, we did not expect to find consistent alterations, but we did observe individual alterations related to cholesterol levels, muscle strength, hypoalgnesia, hematology, triglyceride and hormone levels (summarized in Table 3) in patients or the female carrier, which strongly suggest that the effects of FTSJ1 deficiency are not restricted to cognitive features in humans either.

| Table 3. Clinical investigation of ID patients with mutation in FTSJ1. |
|-----------------|-----------------|-----------------|-----------------|
| **Family A003** | **Affected older brother 1894** | **Mother carrier 1751** | **MRX9** |
| **Index patient 1848** | **Affected uncle 1894** | **Mother carrier 1751** |
| **Moderate** | **Mild** | **None** | **Moderate** | **Mild** | **None** |
| **Cholesterol** | 4.6 (3.5-5.5) | 4.3 (3.5-5.5) | 7.2 (3.5-5.5) | 134 (150-190) | 197 (<190) | 176 (<190) |
| **LDL-Cholesterol** | 2.9 (<3.5) | 2.3 (<3.5) | 4.9 (<3.5) | normal | 117 (<115) | normal |
| **HDL-Cholesterol** | 1.46 (<1) | 1 (<1) | 1.63 (<1) | normal | normal | normal |
| **Impaired muscle strength** | yes | yes | not investigated | no | not investigated | no |
| **Hypoalgnesia** | yes | yes | no | not investigated | yes | yes |
| **Granulocytes elevated** | no | no | no | no | no | no |
| **Neutrophil** | 2.4 (2.0-7.5) | 2.4 (2.0-7.5) | 1.9 (2.0-7.5) | 45% (50-70) | 40.1 (38.9-74.9) | 45% (50-70) |
| **Eosinophil** | not investigated | not investigated | not investigated | 17% (0-5%) | 5.30-5% | 3.2% (0-5%) |
| **Triglyceride (S Trig)** | 0.6 (<1.5) | 1.6 (<1.5) | 1.5 (<1.5) | normal | normal | normal |
| **Estimated glomerular filtration rate (egfr)** | normal | normal | normal | normal | normal | normal |
| **Glucose levels (Fasting)** | 5.5 (3.6-6.0) | 5.3 (3.6-6.0) | 6.8 (3.6-6.0) | 72 (55-115) | 84 (82-109) | 85 (70-100) |
| **Bicarbonate** | 21 (20-32) | 33 (20-32) | 29 (20-32) | not investigated | 28 (22-33) | 25 |
| **Creatine** | 50 (60-110) | 58 (60-110) | 61 (60-110) | 0.7 (0.70-1.20) | 0.65 (0.5-1.2) | 0.9 (<1.10) |
| **Aspartate aminotransferase** | 16 (10-40) | 9 (10-40) | 21 (10-40) | 22 (3-35) | 16 (<40) | 29 (<31) |
| **Amylase** | not investigated | 19 (20-100) | 56 (20-100) | not investigated | 52 (29-100) | 71 (<125) |
| **Testosterone** | not investigated | not investigated | not investigated | not investigated | not investigated | not investigated |
| **Sexual hormone binding globulin (SHBG)** | not investigated | not investigated | not investigated | not investigated | not investigated | not investigated |

| **Intellectual Disability** | **Family A003** | **Affected older brother 1894** | **Mother carrier 1751** | **MRX9** |
|-----------------|-----------------|-----------------|-----------------|-----------------|
| **Hemicopenin** | 22.99 | 3.64 | 6.32 | 9,33E-03 | Hemopexin |
| **Mir145a** | 0.33 | 12.25 | 0.03 | 1,07E-02 | MicroRNA 145a |
| **Aldob** | 52.29 | 18.52 | 2.82 | 1,07E-02 | Aldolase B, fructose-bisphosphatase |
| **Eps81** | 68.92 | 28.46 | 2.42 | 1,07E-02 | EPS8-like 1 |
| **Ugt2b1** | 17.33 | 1.98 | 8.75 | 2,05E-02 | UDP glucuronosyltransferase 2 family, polypeptide B1 |
| **Tor2a** | 116.15 | 194.21 | 0.60 | 2,16E-02 | Torsin Family 2, Member A |
| **Car3** | 31.32 | 8.26 | 3.79 | 2,76E-02 | Carbonic anhydrase 3 |
| **Mup20** | 9.33 | 0.00 | Inf | 2,89E-02 | Major urinary protein 20 |
| **Snr66** | 189.74 | 115.19 | 1.65 | 3,57E-02 | Small Nucleolar RNA, C/D Box 66 |
| **Snr67** | 634.48 | 451.26 | 1.41 | 3,64E-02 | Small Nucleolar RNA, C/D Box 67 |
| **Serp[hal]h1** | 302.56 | 199.10 | 1.52 | 3,69E-02 | Serine Peptidase Inhibitor H1 |
| **Snr87** | 1003.32 | 735.87 | 1.36 | 3,75E-02 | Small Nucleolar RNA, C/D Box 87 |
| **Gab[hal]a6** | 729.60 | 526.65 | 1.39 | 3,88E-02 | Gamma-aminobutyric acid (GABA) A receptor, subunit alpha 6 |
| **Lhx1** | 60.60 | 26.16 | 2.32 | 4,81E-02 | LIM Homeobox Protein 1 |
| **Apoh** | 15.00 | 1.66 | 9.04 | 4,91E-02 | Apolipoprotein H |
4. Discussion

We report here on the generation and characterization of a mouse model for Ftsj1 deficiency and show that Ftsj1 deficiency is not restricted to cognitive features but affects the entire organism. The Ftsj1 genetrapped mice show a mild learning/memory phenotype only, whereas human male patients are also moderately or severely affected. Since the male mice were almost a year old at the time of testing more experiments using mice of different age are required to investigate this impairment in detail. We observed pronounced differences between hemizygous male mutation carriers and male controls. We also saw differences between heterozygous and homozygous females, albeit less pronounced, suggesting that heterozygous female mice are not entirely unaffected. This is surprising, as human heterozygous females seem largely unaffected by FTSJ1 mutations (18). Still, as the Ftsj1 expression pattern in female heterozygous mice is not clear we focused on the effects in male mice where the differences are more pronounced. The gene trapped mice are smaller than wild type littermates but have no organ abnormalities when correcting for the lower body weight. However, the reduced bone mass observed in the gene trapped mice suggests an altered bone metabolism. This is probably related to the reduced calcium levels and the increased alkaline phosphatase activity we observed in the blood of gene trapped mice.

In red blood cells of gene trap carriers, we observed mild microcytosis and increased anisocytosis, which could be related to the lower iron levels we also observed. The effect is significant but not very pronounced as the hemoglobin levels were similar between gene trapped mice and controls. The platelet number was slightly reduced, which could indicate an impaired thrombopoiesis. We also found an increased number of leukocytes, but no signs of allergy or an autoimmune response.

Although we did not observe altered energy metabolism or body temperature, the mutant mice have reduced blood glucose levels and clear intraperitoneally injected glucose faster than controls. Interestingly, patients with mutations in the tRNA modifying enzyme TRMT10A (MIM616033) present with ID, short stature and an abnormal glucose metabolism related to abnormal insulin levels (9,10). These patients also have reduced blood sugar levels but, in contrast to Ftsj1 gene trapped mice, clear externally given glucose inefficiently, even though they have normal cortisol levels (10).

The Ftsj1 gene trapped mice also have decreased levels of alpha-amylase, cholesterol, triglycerides and creatinine, which is indicative of an altered energy metabolism. We also observed a relative increase in body fat content and reduction in lean mass. However, we found no alterations in lactate levels or in oxygen consumption, suggesting normal mitochondrial oxidative capacity.
Of the three investigated hormones (testosterone, aldosterone and corticosterone) corticosterone showed significant upregulation. Corticosterone belongs to the glucocorticoids, which are important regulators of bone metabolism, energy metabolism and the immune system. It is the major glucocorticoid in mice, whereas the major glucocorticoid in humans is cortisol. It is therefore tempting to speculate that the reduced body size, bone phenotype, altered glucose and fat metabolism as well as the immune system phenotype are related to the elevated corticosterone levels we observed in the gene trapped mice. Still, further investigations are required to elucidate such putatively specific effects in gene trapped mice as the phenotype varies between individuals. Corticosterone is also a stress related hormone that is known to influence murine learning and memory (26). In a Rett syndrome (MIM 312750) mouse model a stronger corticosterone response to a stress stimulus was observed when compared to controls (27). However, as Ftsj1 gene trapped mice show more explorative behaviour as compared to controls, there is no suggestion of Ftsj1-deficiency related anxiety or any other stress related phenotype. The hypoalgesic phenotype we observed in mutant animals suggests that also the peripheral nervous system is affected by the absence of functional Ftsj1. Interestingly, an increased pain threshold has also been observed in ID patients with Prader-Willi syndrome (MIM 176270) (28) or Rett syndrome (29). In Children with Prader-Willi syndrome, for example, the higher pain threshold has been related to the increased serum neurotensin levels (28) and in a zebrafish model for Rett syndrome knock-down of mecp2 leads to an impaired sensory system due to defects in peripheral innervation of trigeminal sensory neurons (30). Moreover, reduced pain sensitivity is also observed in familial dysautonomia (FD, MIM 223900) patients that harbor a splice-site mutation in IKBKAP/ELP1 (31,32). IKBKAP/ELP1 encodes a subunit of the Elongator complex, which is required for modifying uridine at the wobble position in certain tRNAs, highlighting the importance for proper tRNA modifications in nervous system cells. The mouse model expresses no detectable wild type transcript and truncated forms are found at a reduced level. A similarly reduced expression level has also been reported for most of the ID patients where truncating mutations have been found (15,16,18). Although the presence of a truncated Ftsj1 protein in our mice cannot be entirely excluded, such a putative protein would still lack half of the amino acids and one third of the conserved methyltransferase domain and thus still be dysfunctional. Two seemingly less severe mutations have been reported in ID patients with FTSJ1 mutations. One mutation (c.655G>A) leads to skipping of exon 9 without affecting the FTSJ1 mRNA expression level (15) and a splice-site mutation in intron 8 (c.571+1 G>A), leading to a frameshift, was shown to cause premature termination in exon 9 as well as reduced FTSJ1 mRNA levels (18). Moreover, the ExAC or gnomAD databases contain no loss-of-function entries for this gene.
These observations further corroborate our conclusion that premature termination of the protein abolishes its enzymatic activity.

The RNA expression profile of our model showed significantly altered expression levels of 50 genes, most of them being upregulated in the gene trapped mouse brain tissue. By applying the DAVID functional annotation software (23,23) to the 50 genes, we found enrichment for genes included in the GO term “blood microparticle” (GO:0072562). Blood microparticles are phospholipid microvesicles that contain receptors and other protein characteristics of the parental cell. Elevated microparticle levels have been found in the cerebrospinal fluid of Alzheimer disease patients together with increased Calpain activity (33) which is interesting as we found Capn11 significantly upregulated in Ftsj1 deficient mouse brain tissue.

Differentially expressed genes were also enriched in the KEGG pathways “Complement and coagulation cascade” and “platelet activation” supporting an effect on microparticles in Ftsj1 deficient mice. However, further investigations are necessary to show if Ftsj1<sup>−/−</sup> mice actually contain more blood microparticles than controls.

We also found modest upregulation of non-coding RNAs including the long non-coding RNA gene Gas5 (Growth arrest-specific 5), and several snoRNAs. Gas5 shows higher expression in growth-arrested cells (34) as well as in lysine deprived cells (35). This suggests that Ftsj1 deficient mouse brain cells activate an amino acid starvation response as it has been shown for Trm7 deficient yeast cells (37). The Gas5 transcript contains a glucocorticoid-binding motif in the 3’ end, which is able to mimic the glucocorticoid response element. The binding of Gas5 transcript to the glucocorticoid receptor (GR) leads to inactivation of GR, which is then no longer able to activate transcription (36). The elevated corticosterone levels observed in the blood of gene trapped mice furthermore suggest that the active GR levels are balanced with the elevated Gas5 levels because we did not observe differential expression of other GR regulated genes than Gas5 (38).

Eight of 50 deregulated genes encode snoRNAs involved in site specific modifications of ribosomal RNA molecules (5.8S, 18S and 28S rRNA) and the U6 spliceosomal RNA. snoRNAs containing an H/ACA box are involved in the conversion of uridine to pseudouridine (39) and CD box containing snoRNAs guide 2′-O-methylation (40). Yeast Trm7 and human FTSJ1 deficient cells contain hypo-methylated tRNA<sup>Phe</sup> (19), which is most likely also the case for the gene trapped mouse brain cells. Upregulation of snoRNA genes in the gene trapped cells could suggest the presence of a feedback mechanism that is activated in response to the presence of hypomethylated tRNAs. Two tRNA quality control mechanisms have been described that eventually either degrade pre-tRNAs lacking 1-methyladenosine (41) or degrade mature hypomodified tRNAs through the rapid tRNA decay pathway (42). Cleavage of tRNAs in response to cellular stress has also been reported for
Nsun2 deficient mice (43). However, further experiments are required to determine the fate of hypomethylated tRNA<sup>the</sup> in Ftsj1 deficient nerve cells.

Mutations in FTSJ1 mainly give rise to intellectual disability but a re-investigation of patients from two families (A003 and MRX9) showed that several features present in individual patients also could be observed in some of the gene trapped mice. Phenotypic variability seems to be pronounced in both human and mice with mutated FTSJ1/Ftsj1. However, the phenotypic variability is not caused by differences in the genetic background or from different treatment of the mice as the mice are essentially identical, born within the same week and fed and treated the same way.

That impaired protein synthesis probably is the key to understand the effects of FTSJ1 deficiency in mouse and man is in line with recent findings of mutations in aminoacyl-tRNA synthetase genes in ID patients. Mutations have been reported for SARS and WARS2 (44), PARS2 and NARS2 (45), IARS (46) and RARS2 (47) and in AIMP1, which encodes a part of a complex that ligates amino acids to their respective tRNA (48). Furthermore, yeast cells deficient for the Ftsj1 homolog Trm7 show a severe growth defect and impaired translation (20). This is also in line with the lower total protein levels we observed in blood from the gene trapped mice.

FTSJ1 is ubiquitously expressed, but only causing non-syndromic ID when mutated as other clinical manifestations are inconsistent. However, an increased pain threshold was found in the gene trapped mice and in affected individuals from both investigated ID families and may be a feature that is more common in patients with FTSJ1 deficiency. Why mutations in this evolutionarily old protein lead to a restricted phenotype in human and mice is still an open question. As FTSJ1 deficiency is not lethal it is possible that evolutionarily comparatively young features like human cognition may be more severely affected than other tissues because the brain has not had the time to develop proper back-up mechanisms to compensate for FTSJ1 deficiency. The Ftsj1 gene trapped mice will be a suitable model for addressing the different effects of disturbed protein synthesis in different mammalian tissues and answering the question as to why the central and peripheral nervous systems are more sensitive than other organs.

In conclusion, we have generated and thoroughly characterized a mouse model for non-syndromic intellectual disability. The Ftsj1 deficient mice are small and have cognitive deficits and altered bone and energy metabolism. Muscle strength and pain sensing is reduced and the immune system and the hormone system is also affected. Some of these features were found upon re-examination of ID patients with FTSJ1 mutations, though too few ID families with FTSJ1 mutations have been identified to define a syndrome. Functional enrichment of differentially expressed mRNAs suggests involvement of blood microparticles and coagulation factors in brains of mice with defective Ftsj1.
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**Databases**

DAVID: https://david.ncifcrf.gov/

ExAC Browser: http://exac.broadinstitute.org/

gnomAD: http://gnomad.broadinstitute.org/
Highlights for
A mouse model for intellectual disability caused by mutations in the X-linked 2'-O-methyltransferase $Ftsj1$ gene:

- $FTSJ1$ encode a tRNA methyltransferase
- Mutations in $FTSJ1$ causes non-syndromic intellectual disability (ID)
- Ftsj1 genetrapped mice are affected by ID and show other syndromic features
- Some of these features were identified in patients with $FTSJ1$ mutations