Article

Mid-Gestation lethality of Atxn2l-Ablated Mice

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Abstract: Depletion of yeast/fly Ataxin-2 rescues TDP-43 overexpression toxicity. In mouse models of Amyotrophic Lateral Sclerosis via TDP-43 overexpression, depletion of its ortholog ATXN2 mitigated motor neuron degeneration and extended lifespan from 25 days to >300 days. There is another ortholog in mammals, named ATXN2L (Ataxin-2-like), which is almost uncharacterized but also functions in RNA surveillance at stress granules. We generated mice with Crispr/Cas9-mediated deletion of Atxn2l exons 5-8, studying homozygotes prenatally and heterozygotes during aging. Our novel findings indicate that ATXN2L absence triggers mid-gestational embryonic lethality, affecting female animals more strongly. Weight and development stages of homozygous mutants were reduced. Placenta phenotypes were not apparent, but brain histology showed lamination defects and apoptosis. Aged heterozygotes showed no locomotor deficits or weight loss over 12 months. Null mutants in vivo displayed compensatory efforts to maximize Atxn2l expression, which were prevented upon nutrient abundance in vitro. Mouse embryonal fibroblast cells revealed more multinucleated giant cells upon ATXN2L deficiency. In addition, in human neural cells, transcript levels of ATXN2L were induced upon starvation and glucose and amino acids exposure, but this induction was partially prevented by serum or low cholesterol administration. Neither ATXN2L depletion triggered dysregulation of ATXN2, nor a converse effect was observed. Overall, this essential role of ATXN2L for embryogenesis raises questions about its role in neurodegenerative diseases and neuroprotective therapies.

Keywords: poly(A)-tail; RNA chaperone; Spinocerebellar ataxia type 2; SCA2; tauopathy; fronto-temporal lobar dementia; tyrosine kinase receptor signaling; nutrient endocytosis

1. Introduction

In all eukaryotic organisms, at least one copy of the Ataxin-2 gene (gene symbol ATXN2 in humans) is phylogenetically conserved and serves roles during nutrient stress for RNA surveillance [1]. A conserved Lsm and LsmAD motif enables direct interaction with RNAs, and a PAM2 motif mediates association with the poly(A)-binding protein PABPC1 [2,3]. Thus, most Ataxin-2 protein localizes with mRNAs at the rough endoplasmic reticulum with marker Ribosomal Protein S6 (RPS6 aka S6R) during cell growth periods [4], where its absence leads to expression adaptations of the associated ribosomal translation machinery [5] and modulates the phosphorylation control of translation [6]. During cell stress, e.g., from nutrient deprivation, Ataxin-2 is transcriptionally induced [6] and relocates with the small ribosomal subunit and PABPC1 to stress granules (SG) [7]. The RNA surveillance
function of Ataxin-2 seems to be relevant to protect against the translation of viral RNAs, given that poliovirus is optimized to cleave Ataxin-2 [8]. In mammals, all these protein structure domains are also found in its paralog Ataxin-2-like (gene symbol ATXN2L in humans). Both ATXN2 and ATXN2L mRNAs also conserved an alternatively spliced exon, which encodes a proline-rich domain (PRD) that mediates its direct association with SH3 motifs in components of the growth factor receptor endocytosis apparatus [9–11]. Abnormal ATXN2 splicing and alternative polyadenylation were documented in diseases with RNA toxicity, such as amyotrophic lateral sclerosis (ALS) [12]. The common ancestor of both proteins in yeast and Caenorhabditis elegans was observed to suppress growth signaling via mTORC1, modulating cell size, and lipid stores [13–15]. This reprogramming of nutrient metabolism is accompanied by an important influence on the mitochondrial breakdown of fatty acids and amino acids, as well as glucose utilization [16–19], probably mediated by the direct protein interaction of ATXN2 with the cytosolic enzyme BCAT1 [20] as the rate-limiting factor in the breakdown of leucine, isoleucine, and valine.

The role of ATXN2 in neurodegenerative diseases has triggered intense research over the past 25 years. Exclusively in human ATXN2, an N-terminal domain with 22 consecutive glutamines (polyQ) exists, which can undergo expansion mutations across generations. Large expansions beyond the size of 32Q trigger the multi-system nervous tissue atrophy Spinocerebellar ataxia type 2 (SCA2), while intermediate expansions of sizes 27Q-32Q increase the risk to be affected by motor neuron diseases such as ALS, fronto-temporal lobar atrophy (FTLD) [24,27–29]. In addition, in yeast, depletion of the ATXN2/ATXN2L ortholog PBP1 rescues the lethal effect of poly(A)-binding protein deletions [30]. The constitutive knock-out of Atxn2 in mice leads to progressive weight gain with excessive storage of lipid droplets and glycogen in the liver, elevated cholesterol and other lipids in the blood, beta-cell hyperplasia in the pancreas with hyperinsulinemia and insulin resistance, increased ganglioside and sulfatide lipids in the brain myelin, locomotor hyperactivity, and mild infertility with gender-dependent impairment of embryogenesis [31,32].

In view of the importance of ATXN2 orthologs for stress response, redundancy occurred in land plants and in vertebrates (except birds) by the co-existence of two homologous genes, named CID3-CID4 in Arabidopsis thaliana weed and ATXN2-ATXN2L in humans and rodents [33]. ATXN2L protein dimerizes with ATXN2 in yeast-two-hybrid tests, and is also a regulator of SGs and mRNA processing during starvation periods, but shows more co-localization with the nuclear splice apparatus than ATXN2 due to an arginine-dimethylation [34,35]. Similar to ATXN2, ATXN2L associates with plasma membrane receptors in dependence on their phosphorylation status, is involved in epidermal-growth-factor (EGF)-receptor signaling, and exists in several isoforms [36–38]. Little more is known at present about ATXN2L. Database mining at the STRING web platform for Protein-Protein Interaction Networks and Functional Enrichment Analysis, available online at: https://string-db.org/ [39] confirms that human and mouse ATXN2L show direct protein-protein-interaction with the poly(A)-binding-proteins PABPC1/2, PABPC1L/PABPC3/PABPC4/PABPC6, with further stress-granule-components such as G3BP2, DDX6, and LSM12 [40,41], as well as nuclear RNA-binding protein NUFIP2 and nuclear transcription modulator NFATC2IP. The RNA helicase activity of DDX6 enhances the expression of growth factor receptors such as HER2 and FGFR2 [42]. Possibly by this mechanism, ATXN2 was found to regulate surface receptors [43]. A recent global protein interactome study via FLAG-tag affinity purification and mass spectrometry in HEK293 cells also showed ATXN2L to associate with DDX6 and the proline-rich-domain-containing RNA-binding Ras-GTPase G3BP2, but in addition, demonstrated ATXN2L binding to KLHL20 [20], which is responsible for the degradation of the autophagy/growth modulators DAPK1 and ULK1, as well as of the Rho Guanine nucleotide exchange factor and glutamate-transport/neurite-growth-modulator ARHGEF11. STRING also lists a potential interaction (based on association in lower species) with SLC9A3R2/NHERF2 that is known to act as
PSD-95 scaffold to control albumin endocytosis, the glutamate transporter GLAST, and the metabotropic glutamate receptor mGluR5, in addition to activating Src phosphorylation and modulating the high density lipoprotein receptor SR-B1 that is crucial for cholesterol metabolism [44–48].

Further data mining at the Allen brain atlas, available online at: https://mouse.brain-map.org/ [49] retrieves evidence that *Atxn2l* mRNA has a particularly strong expression in cerebellar Purkinje neurons. Protein data in different cell types from the Human Protein Atlas, available online at: https://www.proteinatlas.org/ [50] demonstrate its prominent abundance in testis. Its moderate abundance in postmitotic neurons from the cortex, cerebellum, hippocampus, and caudate nucleus, as well as glia and muscle cells were detected by the Atlas antibody HPA041506 targeting ATXN2L at aa 456-547 (in the Uniprot Q8WWM7-1 sequence), which exhibits a single immunoblot band of predicted size and has siRNA-controlled specificity. However, it was not detected by Atlas antibody HPA043391, which targets human ATXN2L aa 572-644, but does not exhibit immunoblot signals of correct size and was not tested by siRNA, as can be seen within the Human Protein Atlas online at: https://www.proteinatlas.org/ENSG00000168488-ATXN2L/tissue/primary+data [50]. Subcellular analyses with both antibodies showed a diffuse cytosolic signal. Among cell lines, a particularly strong expression is detected under lymphoid differentiation, in good agreement with the importance of RNA surveillance for the innate and adaptive immune system. The upregulation of ATXN2L in cancer tissue is an unfavorable marker, in agreement with its published induction by EGF growth signaling [38]. A co-expression survey at the Coexpedia database, online available at: www.coexpedia.org [51] shows that mouse Atxn2l mRNA is regulated together with pathways of thin myocardium, open neural tube, improved glucose tolerance, immune defense, and cell cycle. The RNA binding spliceosome component Srrm2 and the proline-rich inflammatory splicing factor Prrc2a show the highest co-expression score.

In a recently generated mouse with the aggregation of ATXN2 due to a polyglutamine (Q100) expansion knock-in [52], we observed a significant accumulation of ATXN2L peptides in cerebellar global proteome profiles from old animals. It was difficult to know if the impact of this anomaly has pathogenic or compensatory roles, and it was impossible to study it further in view of the unavailability of antibodies shown to be specific, of tagged recombinant constructs and mutant cells. We decided to start an exploratory project into the physiological functions of ATXN2L. Working at organism level, we generated novel Atxn2l-ablated mice to assess their phenotypes in growth, histology, and behavior, in comparison to our previous study of Atxn2-ablated mice [31]. Working at cell culture level, we examined the cellular phenotype of Atxn2l−/− mouse embryonal fibroblasts (MEF), and whether the regulation of human ATXN2L and ATXN2 mRNA levels is similar in response to stressors for neural cells. Overall, the data indicate that ATXN2L is much more important than ATXN2 for growth at early embryonal age, particularly in females. Atxn2l exon 5-8 deletion triggered a growth deficit phenotype rather than the weight excess observed in mice with Atxn2 exon 1 deletion. Apparently, ATXN2L has a similar function as ATXN2, which is not only transcriptionally induced upon deprivation from nutrients and especially lipids, but also upon exposure to excess glucose or amino acids. Hence, we believe that elucidating the native role of ATXN2L throughout development and in response to stress provides important insights into the complementary spatio-temporal roles of ATXN2-ATXN2L and will further our understanding of neurodegenerative pathomechanisms.

2. Results

We obtained a null allele of murine Atxn2l via Crispr/Cas9-mediated deletion between two sgRNA sites flanking the exons 5-8 of the gene (outsourced work by TIGM, College Station, TX, USA) (see Figure 1).
Figure 1. Scheme of genetic ablation within Atxn2l, with its splicing and translation effects. Ataxin-2-like (A) at the DNA level with neighbor genes Tufm and Sh2b1, indicating the deletion region in letters and lines of magenta color (also showing alternative translation start codons (ATG), structure from exon 1 to exon 22 with introns, alternative translation STOP codons, start of poly(A) tail); (B) at the mRNA level with alternatively spliced isoforms in separate red boxes; (C) at the protein level with relevant known sequence motifs (pro-rich refers to proline-rich-domain, MPL motif was reported by Meunier et al. 2002), highlighting the phylogenetically conserved Lsm, LsmAD and PAM2 motifs in a red box (deletion and frameshift are limiting translation to an N-terminal fragment until magenta line).

The absence of ATXN2L protein epitopes from residue 150 until residue 1050 was confirmed in Atxn2l−/− mouse embryonal fibroblast (MEF) (Figure 2A). ATXN2L abundance was higher in cerebral cortex than in cerebellum. While MEFs showed two specific ATXN2L bands in immunoblots, in brain tissue the larger band dominated (Figure 2B). The separate quantification of several Atxn2l mRNA exon junctions, namely 7-8 to represent the deleted region, 10-11 further downstream, and 1-2 upstream versus the alternatively spliced 1c-2 suggested the Atxn2l−/− limb cells to sense the loss-of-function and either trigger the Atxn2l promoter or minimize the Atxn2l mRNA decay, thus maximizing the levels of the main Atxn2l mRNA variant. This feedback did not occur in MEF lines that are maintained in nutrient abundance (Figure 2C), suggesting that deprivation of overall energy or specific nutrients modulates Atxn2l transcription.
particularly for females at the prenatal stage (see Tables 1 and 2). There was also reduced viability of heterozygous mice, particularly for females at the prenatal stage (see Tables 1 and 2).

Figure 2. (A) Protein abundance of 3 different epitopes spanning ATXN2L in control Atxn2l+/+ and homozygous Atxn2l−/− MEF (n = 4 vs 4). The protein was completely absent in three assessed epitopes from residue 150 until the most C-terminal region (aa: amino acid). (B) Protein abundance of ATXN2L in heterozygous Atxn2l+/− cerebellum (cb) compared to wildtype littermates, or in heterozygous Atxn2l+/− cortex (ctx), in comparison to one wildtype Atxn2l+/+ MEF line. MEFs exhibit two specific bands of ATXN2L, in contrast to brain tissue where the larger band dominates. The overall ATXN2L levels were decreased to approx. 50% in heterozygous tissues (n = 2). Detection was done for the epitope aa 712-1050. (C) In Atxn2l+/− MEF (upper row, n = 4 vs 4), RT-qPCR showed that mRNA expression of Atxn2l was absent at the boundary between exons 7-8 that was genetically deleted, and significantly reduced at the boundaries of exons 1-2 and 10-11, possibly due to nonsense-mediated RNA decay. In tissue from fore-/hind-limbs [lower row, 3 wildtype (Atxn2l+/+, white bars), 4 heterozygous (Atxn2l+/−, grey bars), 2 Atxn2l-null embryos (Atxn2l−/−, black bars)], the Atxn2l mRNA expression measured at the boundary of exons 7-8 was again absent in Atxn2l−/− and reduced by 50% in Atxn2l+/− mice, whereas 5’-upstream and 3’-downstream from the deletion the increased Atxn2l expression level of null mutants suggested transcript upregulation efforts either via the Atxn2l promoter or via altered mRNA stabilization/degradation to compensate the loss of protein function. The exon 1c-2 assay detects an alternatively spliced isoform. Asterisks represent significance (p < 0.01 **, p < 0.0001 ****).

While the heterozygous mice bred normally, a lethality at the embryonic/prenatal stage was observed for the homozygous mutants. There was also reduced viability of heterozygous mice, particularly for females at the prenatal stage (see Tables 1 and 2).
Table 1. Postnatal genotype distribution for offspring from Atxn2l+/− intercrosses.

| Observed/Expected Number of Live Born Mice with Indicated Genotype | Number of offspring |
|---------------------------------------------------------------|---------------------|
| Live born female                                              | 127/220             |
| +/+                                                           | 50/55               |
| +/−                                                           | 77/110              |
| −/−                                                           | 0/55                |
| Live born male                                                | 143/220             |
| +/+                                                           | 60/55               |
| +/−                                                           | 83/110              |
| −/−                                                           | 0/55                |
| Live born total                                               | 270/440             |
| +/+                                                           | 110/110             |
| +/−                                                           | 160/220             |
| −/−                                                           | 0/110               |

The dissection at different stages of pregnancy showed the death of homozygous mutants to have mostly occurred by embryonic day E14-16. Female Atxn2l−/− embryos were more severely affected, most of them did not reach E11 (Table 2). These observations were in good agreement with reports in C. elegans worms that embryonal patterning is arrested, stem cell proliferation is reduced, and germline abnormally masculinized upon depletion of the ortholog ATX-2 [53,54]. Interestingly, at the 2-4 cell stage of bovine embryos, a 2.4-fold expression difference of Atxn2l was recently observed to correlate with fertility and that the knock-down of Atxn2l enhanced the development of blastocysts [55].

Table 2. Embryonic day E11-20 genotype distribution for offspring from Atxn2l+/− intercrosses. Most lethality occurred before E11, with females dying before males.

| Observed/Expected Number of Mice with Indicated Genotype | Number of offspring |
|---------------------------------------------------------|---------------------|
| Dissected embryos female                                | 110/156             |
| +/+                                                     | 48/39               |
| +/−                                                     | 57/78               |
| −/−                                                     | 5/39                |
| Dissected embryos male                                  | 117/156             |
| +/+                                                     | 30/39               |
| +/−                                                     | 71/78               |
| −/−                                                     | 16/39               |
| Dissected embryos total                                 | 227/312             |
| +/+                                                     | 78/78               |
| +/−                                                     | 128/156             |
| −/−                                                     | 21/78               |

In the male homozygotes surviving beyond E11, there was a weight loss of 25% on average (Table 3).

Table 3. Weight of embryos in mg directly after dissection at three different embryonal ages. n: Atxn2l+/+ = 1-2, Atxn2l+/− = 4-6, Atxn2l−/− = 1, E = embryonal day.

| Weight of Embryos [mg] | +/+ | +/− | −/− |
|------------------------|-----|-----|-----|
| Day E14–15             | 322 | 306 | 226 |
| Day E15–16             | 447 | 444 | 313 |
| Day E20                | 1472| 1481| 372 |

Only one dead male Atxn2l−/− embryo was identified at E20 (Figure 3A, right side), exhibiting disintegrating tissue surrounded by dense liquid as expected during resorption processes [56]. An Atxn2l−/− embryo at E16 was observed to display retarded growth (Figure 3B, right side). The Atxn2l−/− embryos were not pale, had livers of usual size, and showed normal blood-filling of liver tissue (Figure 3C, right side), so there was no evidence for the cardio-vascular or blood circulation deficits that are frequent causes of early or mid-gestational lethality [57]. Strong developmental differences between Atxn2l−/− versus WT littermates (Figure 3D) sometimes amounted to several Theiler stages [58–60]. No selective organ affection in gross morphology was noticed during embryogenesis (Figure 3A–D), so an evaluation by high-resolution episcopic microscopy [61] was deemed unnecessary. In general, many fetal sacs in utero were found to contain only placenta rests and a fluid-filled cavity.
Figure 3. Comparison of Atxn2l−/− embryos with their littermates, at different gestational stages. Embryonal days reflect approximate values. (A) The only Atxn2l−/− mouse (male) identified at E20 in comparison to a male WT brother. (B) An Atxn2l−/− mouse (male) reaching E16, exhibiting retarded development, and decreased weight, in comparison to the heterozygous Atxn2l+/− female sibling. The petechial bleeds were not seen in other null mutants. (C) An Atxn2l−/− mouse (male) at E15-16 with liver size and blood filling similar to male WT brother. (D) An Atxn2l−/− female embryo at E13 with retarded growth and development and its male WT sibling.

Histological analyses at different embryonic stages from approximately E13 via E14 to E20 (Figure 4) confirmed that the growth deficit is generalized throughout the embryos.

The most frequent causes of early/mid-gestation embryonic lethality before E14 include deficient placentation, followed by blood vessel/heart, skeleton/joint and nervous system malformations [62]. In Atxn2l−/− embryos, no obvious phenotypes apart from developmental delay were detected upon hematoxylin/eosin (H&E) staining of placenta (Supplementary Materials Figure S1), heart, and bones. Interestingly, mice with genetic ablation of the EGF receptor, which was observed in cell lines to influence ATXN2L expression, also exhibit a mid-gestational lethality. However, they display reduced placenta size [63,64] in contrast to the normal Atxn2l−/− placentas. Mid-gestation embryonic lethality was recently also documented for factors of endocytic uptake, such as DENND1A and VPS54 [65,66].

In Atxn2l−/− embryos, preferentially the nervous tissue showed frequent apoptosis, together with reduced cortical layer formation (Figure 5). Similar mid-gestation lethality with abnormal development of the brain cortex was observed, e.g., upon deficient methylation of eukaryotic mRNAs [67]. Neurons more than other cells have a need to modify the 3’-untranslated regions of mRNAs via alternative polyadenylation and microRNA repression [68,69] in a pathway that involves the ATXN2/ATXN2L indirect interactor TDP-43 and stress granules.
Analyzing development of the brain cortex was observed muscle coordination reduced cortical layer formation (Figure 4).

Interestingly, mice with genetic ablation of the EGF receptor, which was observed in cell lines to influence thinning brain (left); in littermate Atxn2l−/− mice (right), less lamina definition and many neurons were observed with nuclear condensation, which is indicative of apoptosis; the diameter of brain cortex was also thinner.

Evident growth/weight phenotype: Atxn2l+/+ embryos with Atxn2l−/− littermates (WT above, homozygote mutants below, E13 on left side, E14 in center, E20 on right side).

Figure 4. Evident growth/weight phenotype: Atxn2l+/+ embryos with Atxn2l−/− littermates (WT above, homozygote mutants below, E13 on left side, E14 in center, E20 on right side).

Figure 5. H&E histology at E14 showed good separation of neuron layers in the cortex of Atxn2l+/+ brain (left); in littermate Atxn2l−/− mice (right), less lamina definition and many neurons were observed with nuclear condensation, which is indicative of apoptosis; the diameter of brain cortex was also thinner.
In view of the postnatal survival of Atxn2l<sup>−/−</sup> mice, a cohort of 12 WT animals and 12 heterozygous littermates was aged until 12 months and assessed every 3 months regarding survival, growth, and locomotor behavior. This preliminary survey found no increased mortality or weight loss, and detected no prominent deficits in spontaneous movements in the open-field paradigm or in muscle coordination under stress on an accelerating rotarod (Supplementary Materials Figure S2).

Analyzing cell growth in four ATXN2L-null MEF lines versus four sex-matched WT littermates that were derived from dissected embryos, we noticed the presence many multinucleated giant cells. This cell type occurs in primary cell cultures when nuclear division is not followed by cytokinesis [70]. In ATXN2L-null MEF lines, their frequency was consistently increased, with a >2-fold effect size across all lines (Figure 6A,B). Cholesterol is necessary for cytokinesis, its deficiency induces polyploid cell formation, so it is noteworthy that cholesterol anomalies are well documented in patients and mice with ATXN2 mutations [31,71–73]. In addition, the furrow progression during cytokinesis depends on receptor tyrosine kinase activation of SRC and on vesicle endocytosis, where ATXN2 is known to play a modulating role [9,10,74–76].

![Figure 6](image_url)

Figure 6. (A) Cell culture images of Atxn2l<sup>+/+</sup> and Atxn2l<sup>−/−</sup> MEF illustrating the increased number of giant multinucleated cells in the absence of Atxn2l. (B) Quantification of giant cells for 4 different age-matched littermate lines with 3 technical replicates each. The diagram on the left side shows the increase with consistency for the individual MEF lines 1-4, while the diagram on the right side represents the overall increase across all lines. Statistical testing was done by two-way ANOVA and t-test with Welch’s correction, respectively, significance levels were illustrated by ** for p < 0.01, by T for 0.05 < p < 0.1.

Then, we assessed whether the deprivation of nutrients, such as glucose and lipids, indeed alter the expression of Atxn2l, as was previously observed for its homolog Atxn2 [31]. The human neuronal cell line SH-SY5Y, first cultured in high-glucose DMEM medium with fetal calf serum (FCS) supplement, was switched to the starvation medium HBSS without FCS for a time course of 48
h. The sudden exposure to conditions of low glucose, no amino acids, no FCS (containing growth factors/apolipoproteins with lipids/transferrin with iron, etcetera) triggered a phasic induction that peaked at 8–24 h with a 3-fold $\text{Atxn2l}$ mRNA increase, similar to the induction of $\text{Atxn2}$ (Figure 7A,B). Preliminary testing in MEF cells suggest that RNA levels of $\text{Atxn2}$ levels decreased slightly upon stress by a toxic RNA-analogon, Poly(I:C), while $\text{Atxn2l}$ failed to respond (Supplementary Materials Figure S3E–H). These MEF analyses provided no evidence that the depletion of $\text{ATXN2}$ triggers compensatory dysregulation of its homolog $\text{Ataxin-2-like}$ at mRNA or protein level, or that conversely the depletion of $\text{ATXN2L}$ triggers expression or abundance changes of $\text{Ataxin-2}$ (Supplementary Materials Figure S3E–J).

Further analysis of individual nutrients demonstrated that the transcriptional induction of $\text{ATXN2L}$ in HBSS starvation medium could be diminished by the supplementation with FCS (which contains components such as insulin, transferrin, and apolipoproteins), and by low level cholesterol administration, two effects that were similarly observed for $\text{ATXN2}$. This was in contrast to supplementation with excess glucose, amino acids, or excess cholesterol that were unable to downregulate $\text{ATXN2L}$ or further enhanced its levels (Figure 7C,E). The increased transcription of $\text{ATXN2}$ upon glucose and amino acid addition was significant, as well as the induction by cholesterol excess (Figure 7D,F).

Further analyses about components of FCS that could be responsible for the rescue effect in the induction of $\text{ATXN2L}$ are shown in Supplementary Materials Figure S3. Different cell culture media were used as basal media (DMEM having the highest abundance of nutrients, MEM containing fewer nutrients such as vitamins or amino acids, and HBSS containing no nutrients apart from low glucose amounts). The supplements included SPITE (a mixture of sodium selenite, sodium pyruvate, bovine insulin, human transferrin, and ethanolamine), ITS (a mixture of bovine insulin, human transferrin, and sodium selenite), and SITE+3 (containing analogous components as SPITE, but 5 mg/mL BSA, linoleic acid and oleic acid as a lipid energy source instead of pyruvate). However, none of these supplements was sufficient for a significant rescue of the starvation-triggered $\text{ATXN2L}$ induction, to a degree that approached FCS. The comparison of SPITE versus SITE+3 supplementation suggested that fatty acid administration is more helpful than pyruvate for the reduction of $\text{ATXN2L}$ and $\text{ATXN2}$ mRNA levels (Supplementary Materials Figure S3C,D). Overall, these in vitro observations confirm the notion derived from limb tissues in vivo and from MEF cells, that either the promoter activity of the Ataxin-2-like gene is upregulated or the $\text{Atxn2l}/\text{Atxn2}$ mRNA decay is downregulated during periods of growth deficits and then this regulation can be reversed upon availability of specific nutrients. In this context, it is noteworthy that the RegRNA2.0 webtool predicts both transcripts to contain AU-rich elements in their 3′-untranslated region, which are known to modulate the stability of selected mRNA according to growth needs. It is also relevant to mention that the promoters/enhancers of human Ataxin-2/Ataxin-2-like share 476 transcription factor binding sites, according to predictions found at the GeneCards website, including stress-response regulator families that are strongly implicated in neurodegeneration, such as ATF1/2/3/4/7, E2F1/5/6/8/ELF1/3/4/ETS1/ETV1/4/5/6, HNRNP-H1/K/L/L1, or IRF1/2/3/4.

A dysregulation of $\text{ATXN2L}$ levels upon depletion of its ortholog $\text{ATXN2}$ was not observed in MEF cells, nor a converse dysregulation of $\text{ATXN2}$ upon depletion of $\text{ATXN2L}$, neither at protein nor at mRNA level (Supplementary Materials Figure S3E–J). As mentioned in the Introduction however, unpublished global proteome mass spectrometry data in our lab documented a significant increase of $\text{ATXN2L}$ in the brain of $\text{Atxn2-CAG100}$-knockin mice. This finding may therefore be specific to the CAG-repeat expansion mutation and be caused by the polyQ aggregation process in stress granules, where $\text{ATXN2}$ and $\text{ATXN2L}$ have common protein interactors such as DDX6 and may co-precipitate.
The mRNA expression of ATXN2L in human SH-SY5Y neuroblastoma cells was induced by nutrient deprivation in HBSS medium over 48 h with low glucose/no amino acids, in the absence of lipids normally supplied via FCS, similar to the expression of ATXN2 mRNA. For the induction peak at 24 h, the expression upregulation of ATXN2L was partially prevented upon supplementation of 10% FCS, while glucose or amino acids alone had no rescue effect; FCS addition also diminished ATXN2 expression, while glucose and amino acid administration enhanced the expression further; The HBSS induction of ATXN2L was reduced at least as much by low cholesterol concentrations as by FCS, while higher dosage of cholesterol elicited even stronger upregulations. This effect was similarly observed for ATXN2. A-D: n = 3; E/F: n = 8. Statistical testing was done by one-way ANOVA, significance levels were illustrated by asterisks: * p < 0.05; ** p < 0.01; *** p < 0.001; **** p < 0.0001. Comparison was always in respect to nutrient abundant control condition, if not stated otherwise.
3. Discussion

ATXN2L was described as direct RNA-binding protein with sequence homology to nuclear spliceosome factors [37,77]. Its alternatively spliced isoforms may interact with tyrosine kinase receptor signaling at the plasma membrane [36]. The constitutive isoform of ATXN2L was mainly characterized for its role as surveillance factor during transient periods of cell damage, which promotes the formation of stress granules in the cytosol [34]. Now, our ATXN2L-deficient mouse data demonstrate that homozygous loss of its RNA-interaction domains is incompatible with embryonic development from early stages. Their absence delays the cytokinesis stage of cell division and impairs the layer formation and survival of brain neurons.

Human data agree with selective roles of ATXN2L for the growth process, the nervous system, and immunity. Genome-wide association studies (GWAS) of single nucleotide polymorphism (SNP) alleles versus phenotypes in healthy populations (summarized at the NHGRI-EBI Catalog of human genome-wide association studies, online available at: https://www.ebi.ac.uk/gwas/) demonstrated significant effects in several independent studies of the ATXN2L gene locus. Its impact on growth was reflected by associations with body mass index (in diverse reports for different polymorphisms \( p = 4e^{-29} \); \( 2e^{-15} \); \( 9e^{-15} \); \( 2e^{-13} \); \( 3e^{-8} \); \( 2e^{-7} \) [79–81]), body weight \( (p = 2e^{-8}) [81] \), and waist circumference \( (p = 4e^{-8}) [81] \). Its impact on the nervous system was reflected by associations with cognitive function \( (p = 8e^{-35}) [82] \), grip strength \( (p = 1e^{-25}) [83] \), intelligence \( (p = 1e^{-19} \); \( 2e^{-9} \); \( 1e^{-8} \); \( 2e^{-9} \); \( 4e^{-5} \) [84–87]), and educational attainment \( (p = 4e^{-14} \); \( 7e^{-11} \); \( 1e^{-7} \); \( 1e^{-6} \) [85,88,89]). Its impact on immunity was reflected by associations with albumin/globulin ratio \( (p = 6e^{-10}) [90] \), inflammatory bowel disease \( (p = 2e^{-9}) [91] \), and pediatric autoimmune diseases \( (p = 6e^{-7}) [92] \).

Our studies of the expression regulation of \( Atxn2l \) and \( Atxn2 \) by diverse nutrients indicate a strong similarity in their role, with a transcriptional induction being triggered by nutrient deprivation as well as by an excess of cholesterol and glucose (both of which have a role for plasma membrane flexibility/stickiness), while the transcription was reduced upon the supplementation of FCS. Given that the depletion of the ATXN2L ortholog in flies is neuroprotective, there may be therapeutic value in the transcriptional regulation by diverse nutrients indicate a strong similarity in their role, with a transcriptional induction being triggered by nutrient deprivation as well as by an excess of cholesterol and glucose (both of which have a role for plasma membrane flexibility/stickiness), while the transcription was reduced upon the supplementation of FCS. Given that the depletion of the ATXN2L ortholog in flies is neuroprotective, there may be therapeutic value in the expression of other disease models of ALS, where hypercaloric feeding showed to be beneficial, and evaluations on the relative contribution from carbohydrates versus lipids are ongoing [93,94]. An experimental artifact was observed upon nutrient deprivation and upon Poly(IC) stress in vitro: Although the induction strength of \( Atxn2 \) and \( Atxn2l \) showed limited variance during repeated experiments by a student with a SH-SY5Y aliquot and a specific FCS batch, it could however show quite different fold-changes months later in different hands using a separate SY-SY5Y stock with another FCS batch. It remained unclear to what degree passage number, clonal mutation drift or handling variance played a role here (compare Figure 7C vs. E, and Figure S2 F vs. H).

Publicly available expression data on \( Atxn2l \) and \( Atxn2 \) by RNA sequencing and further processing at the Broad Institute within the Genotype-Tissue-Expression project database, online available at: https://gtexportal.org/home/ [95] with normalization to transcripts per million reads (TPM) confirm that \( Atxn2 \) and \( Atxn2l \) are both preferentially expressed in cerebellar hemispheres and total cerebellum (median TPM value = 30/33 vs. 184/229, respectively), in comparison with other regions of the nervous system. Interestingly, our observations on sex differences in the phenotype of null mutants are also reflected by these public expression data: they document that \( Atxn2l \) expression is stronger in most female organs (36/31/30/29/28/24 TPM in ovary/uterus/endocervix/FallopianTube/ectocervix/vagina, respectively) than in cerebellum. In contrast, \( Atxn2l \) shows the highest expression in testis tissue (TPM 296), as opposed to the average expression in female organs (e.g., ovary TPM 124). Overall, \( Atxn2l \) in cerebellum shows a 6-fold higher expression than \( Atxn2 \), and the preferential expression of \( Atxn2l \) in male testis contrasts with the pronounced expression of \( Atxn2 \) in several female sexual organs. Given that the biosynthesis of all sex hormones starts from cholesterol, the effects of ATXN2L and ATXN2 on the cholesterol homeostasis may account for the sex difference of \( Atxn2l \)-null phenotypes, the increased formation of giant multinucleated MEF cells, and the expression regulation by cholesterol.
It is important to note that *Atxn2*-mutations were already observed to affect cholesterol homeostasis and trigger gender-dependent or hormonal effects [31,73,96,97]. Giant multinucleated cells can be due to increased mTOR activity [98–100]. Given that ATXN2 was reported to inhibit mTOR in yeast, nematodes and mice; also the absence of ATXN2L might lead to enhanced mTOR signaling and, in this way, increase the appearance of multinucleated giant cells in MEF cultures. These phenotypes may be downstream effects of the reported protein interaction between ATXN2L and NHERF2 as a known modulator of cholesterol metabolism and Src-dependent phosphorylation signals for growth. While a gene duplication event in mammals led to the coexistence of two homologous genes, in plants a coexistence of four homologous genes from the ataxin-2 family raises the possibility that mTOR-dependent growth can be regulated by each of them in response to different stressors or to different nutrients, via an RNA-binding mechanism.

Clearly, the regulation of *Atxn2l* and *Atxn2* mRNA levels by nutrients is almost parallel in our cell culture analyses, so it seems paradoxical that our null mutant of ATXN2L has a growth deficit phenotype, while the null mutant of ATXN2 was found by two independent teams to show a converse nutrient excess phenotype [31,32]. An important difference in the recombinant design of these mutants should be noted here. The KO of *Atxn2* was targeting exon 1 and removed all of the protein domains, while the ablation in *Atxn2l* targeted exons 5-8, with a resulting shift of reading frame downstream. In the mutant ATXN2L, the N-terminal Pro-rich domain across residues 4-61 and the MPL interaction domain across residues 96-119 would still be synthesized as a fragment, the available commercial antibodies cannot detect this N-terminal region. In view of the increase of *Atxn2l* non-targeted exons in the limb tissue of homozygous mutants (Figure 2B), this hypothetical fragment could exist in overdose. The excess amounts of such an N-terminal fragment would interact with plasma membrane receptors and with the endocytosis machinery. In analogy to the published observations for ATXN2 overexpression, it is expected to impair trophic uptake mechanisms [9]. At the same time, the absent middle and C-terminal part of ATXN2L would fail to interact with RNAs in the appropriate manner. Overall, our mouse mutant might represent not only a loss of the C-terminal functions of ATXN2L; it is conceivable that it also involves a toxic gain of N-terminal functions of ATXN2L. Perhaps this speculation provides a clue why our mouse mutant has this unexpected phenotype, which is converse to *Atxn2*-KO mice.

In the future, antibodies against the N-terminal portion of ATXN2L should be generated to obtain evidence regarding this doubt, and a new ATXN2L mutant that targets exons 1-4 would also help to clarify this issue. It will be crucial to generate conditional mutants where the postnatal role of ATXN2L for tissues such as the aging brain can be studied selectively. Moreover, it is interesting to note that commercial antibodies such as HPA041506/PA5-59601 in U2OS cells showed only the band of predicted size, and our immunoblots in brain tissue also detected this putative full-length band, while we observed two ATXN2L bands with three different antibodies in MEF cells, and also the A301-369A and 24822-1-AP antibodies detected a smaller band in HeLa/HEK293T and Jurkat cells, according to manufacturer datasheets. Thus, further studies will be necessary to understand the usage of the two alternative start codons and of the alternative splice variants in various tissues/cell populations and after different stressors. Clearly, our data indicated that the depletion of ATXN2 or ATXN2L in MEF cells fail to trigger a compensatory upregulation of the other homolog, so the unpublished accumulation of ATXN2L in *Atxn2*-CAG100-KIN brain appears to be an expansion-driven effect and ATXN2L immunoreactivity should be assessed within the Q100-ATXN2 aggregates. Since TDP-43 is essential for embryonic development in mouse and its aggregation drives the decisive toxicity for motor neuron degeneration, it is conceivable that ATXN2L—a factor that is also essential for embryonic development—could co-aggregate with ATXN2 and drive toxicity in cerebellar Purkinje neurons where it has selective abundance.
4. Materials and Methods

4.1. Generation, Breeding, Maintenance, and Dissection of Atxn2l−/− Mice

To specifically target sites flanking exons 5-8 of the Atxn2l gene, this pair of sgRNAs was used: upstreamGRNA085CCT (atgtggcagggtcagtatcag), downstreamGRNA02CCC (gttacctggataccaagct). To distinguish the long wildtype from the short mutant allele, primers Atxn2l-F (gagagtgtgtgttgggtgac), Atxn2l-F2 (gaagcacagtgctgttcagc), and Atxn2l-R (gtaatttgcagcaacagtagagc) were employed. Heterozygous breeders were shipped, crossed at the ZFE animal facility of Goethe University Medical Faculty in Frankfurt as described [101] and health-monitored with sentinel animals at quarterly intervals. Congenic heterozygous mice were then crossed among them. The studies were ethically approved by Regierungspräsidium Darmstadt, code V54-19c20/15-FK/1083, on 11 March 2019.

4.2. Genotyping

DNA was extracted from adult ear punches or embryo tail biopsies for genotyping by incubating at 95 °C in 100 µL of 25 mM NaOH/0.2 mM EDTA for 30 min before neutralizing with 100 µL 40 mM Tris–HCl, pH 5.0. For the subsequent genotyping reactions, 1 µL of lysed tissue sample was used per reaction. The Atxn2l alleles were amplified by PCR with the primers Atxn2l-F (GAGAGTGTGTGTGTGGGTGAC), Atxn2l-F2 (GAAGCACAGTGCTGTTCAGC) and Atxn2l-R (GCTCTACTGTGTGTGCTGCAAAATTAC), the PCR products 217 bp and 309 bp corresponding to Atxn2l wt and Atxn2l mutant respectively were separated in 1.5% agarose gel.

For the determination of the sex, primers Rmb31F (CACCTTAAGAAGCCAATACA) and Rbm31R (GGCTTGTCCTGAAAACATTTGG) were used to yield a 269 bp product from X chromosome and a 353 bp product from the Y chromosome [102].

4.3. Behavioral Tests

Behavioral tests were done as described in [103]. Twelve Atxn2l+/+ and twelve Atxn2l+/− male mice were aged and assessed at ages between 3 months and 12 months.

4.4. Generation and Cell Culture of MEF Cells

MEF were derived from Atxn2l−/− embryos at day 14-16 and their littermate Atxn2l+/+ controls. As previously described [104], the skin of the embryos was dissected, homogenized, and trypsinized for 10 min at 37 °C. The cells (passage 2-3) were cultured in DMEM, containing 4.5 g/L D-glucose, 1% L-glutamine, 1% Penicillin/Streptomycin and 10% FCS. Atxn2l−/− MEF cells were described before [9].

4.5. Poly(I:C) Treatment

Atxn2+/+ and Atxn2−/− MEF (3 vs. 3 cell lines), as well as Atxn2+/+ and Atxn2−/− MEF (4 vs. 4 cell lines) were treated with 1 µg/mL Poly(I:C) (Invivogen) for 16 h with subsequent RNA extraction and RT-qPCR [105].

4.6. RNA Extraction and Expression Analysis

RNA extraction from limbs and MEF was performed with TRIzol Reagent (Sigma Aldrich) according to the user manual. RNA from cell culture experiments with SH-SY5Y cells was performed with Extractme Total RNA Kit (7Bioscience EM09.1-250). Synthesis of cDNA from 1 µg of total RNA template was performed with the SuperScript IV VILO kit (ThermoFisher) according to the manufacturer’s instructions. cDNA from 25 ng total RNA was used for each RT-PCR reaction with 0.5 µL TaqMan® Assay, 5 µL FastStart Universal Probe Master 2x (Rox) Mix and ddH2O up to 10 µL of total volume. The mouse-specific TaqMan® Assays utilized for this study: Atxn2 (Mm00805548_m1, Mm00805539_m1, Mm01165459_m1, Mm01276208_m1), Atxn2 (Mm01199894_m1), and Tbp (Mm00446973_m1).
4.7. Immunoblotting

Protein was isolated with RIPA buffer as described before [106]. Samples with 20 µg protein were separated on 8% SDS gels and blotted on nitrocellulose membranes. Membranes were incubated overnight at 4 °C with the following antibodies: ATXN2L (Bethyl A301-369), ATXN2L (Invitrogen PA5-59601), ATXN2L (Proteintech 24822-1-AP), ATXN2 (Proteintech 21776-1-AP) or beta-Actin (Sigma A5441), and for 1 h with the respective secondary antibodies (Li-Cor). Fluorescence was detected with the Li-Cor Odyssey Classic Instrument and bands were analyzed with Image Studio Lite, Version 5.2.

4.8. Sections and Staining

All tissue specimens were fixed in 4% paraformaldehyde (PFA) for at least 48 h. All specimens were embedded in paraffin and aligned as displayed on the respective images (see Figures 4 and 5 and Figure S1). The 4-µm thick sections were made using a microtome. Slides were stained with hematoxylin and eosin according to standard protocols. Images were taken using an Olympus BX50 microscope and an Olympus DP72 camera.

4.9. Microscopy and Cell Counting of MEF

175,000 MEF cells were seeded in 6-well plates and grown overnight (n = 4 Atxn2l+/+ vs. 4 Atxn2l−/− lines in 3 technical replicates each). For each well, 5 random areas were microscopically visualized (Leica) and all giant cells were counted. Means for each cell line was calculated and statistically evaluated. Images were taken with a Nikon microscope after changing the cell culture medium to PBS.

4.10. Cell Culture Experiments

SH-SY5Y cells were purchased from ECACC, and cultured as described [6] in Dulbecco’s Modified Eagle Medium (DMEM) containing 4.5 g/L D-glucose, 2 mM L-glutamine with 10% FCS. The cells were starved of trophic factors by incubating them in Hanks’ balanced salt solution (HBSS) for a range of different periods (time course) to determine when the cells show the strongest deprivation effect. In different experiments, HBSS was supplied with 10% FCS, glucose 4.5%, MEM non-essential amino acids solution 1x, or cholesterol (SIGMA L4666) at 10 µM, 25 µM, cholesterol 125 µM (Figure 7). For the experiments shown in Supplementary Figure S3, cells were incubated with the indicated basal media (DMEM, MEF, or HBSS), supplemented with SPITE medium supplement 1x (Sigma S5666), ITS liquid media supplement 1x (Sigma I3146), or SITE+3 liquid media supplement 1x (Sigma S5295). After 24 h, cells were collected and RNA was extracted with EXTRACTME (Blirt, Gdańsk, Poland). Synthesis of cDNA and RT-qPCR was performed as previously described with human-specific TaqMan® Assay ATXN2L (Hs00944485_g1), ATXN2 (Hs00268077_m1) and HPRT1 (Hs99999909_m1). The numbers of cell lines used is stated in respective figure legends.

4.11. Statistical Analysis

All statistical tests were performed as unpaired Student’s t-test with Welch’s correction or one-way or two-way ANOVA using GraphPad Prism software version 7 after establishing that each population was normally distributed (one-sided Kolmogorov–Smirnov test). Figures display mean values and standard error of the mean (s.e.m.). Values of p < 0.05 were considered significant and marked with asterisks p < 0.05 *, p < 0.01 **, p < 0.001 ***, p < 0.0001 ****.

5. Conclusions

Overall, this first analysis of ATXN2L-depletion in mouse demonstrates its essential role for embryonic development, due to growth delays with prominent vulnerability of brain lamination, especially in females. A preferential affection of female embryos and an increased occurrence of multinucleated giant cells in MEF culture were observed, which may reflect a role of ATXN2L in
cholesterol and hormone homeostasis. This mouse mutant will provide an important tool to dissect the role of ATXN2L domains, such as the impact of LSM and PAM2 motifs on RNA surveillance, which were ablated here, versus the effects of PRD and MPL interaction motifs on trophic endocytosis, during stress periods and aging. Since the data show ATXN2L to be more important than ATXN2 for cell growth, it will be highly interesting to study its role in neurodegenerative disorders due to RNA surveillance problems, such as ALS, where the depletion of mammalian ATXN2 and of the yeast/fly ortholog also of ATXN2L was already shown to be neuroprotective.

Supplementary Materials: Supplementary materials can be found at http://www.mdpi.com/1422-0067/21/14/5124/s1.

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Abbreviations

°C Degrees Celsius (temperature)
aa amino acid
ACTB beta-actin
Adnp Activity-Dependent Neuroprotector Homeobox Protein
ALS Amyotrophic Lateral Sclerosis
ANOVA Analysis of variance
ARHGef11 Rho Guanine Nucleotide Exchange Factor (GEF) 11
ATF1 Activating Transcription Factor 1
ATG Start codon Adenine-Thymidine/Uracil-Guanine
ATX-2 the single nematode ortholog of mammalian Ataxin-2 and Ataxin-2-like
ATXN2 Ataxin-2
ATXN2L Ataxin-2-like, aka: Ataxin-2-Related Protein, A2LP, A2RP or A2D
BDNF Brain-Derived Neurotrophic Factor
bp Base pair
BSA Bovine serum albumin
Carm1 Coactivator Associated Arginine Methyltransferase 1
Cart1 Cartilage paired class homeoprotein 1 (aka Abx1)
Cb Cerebellum
Cbp cAMP-response element binding protein (Creb) binding protein (aka Crebbp)
cDNA complementary Deoxyribo-Nucleic Acid
CID3 CTC-interacting domain 3 protein in Arabidopsis thaliana
CID4 CTC-interacting domain 4 protein in Arabidopsis thaliana
Cited2 Cbp/P300 Interacting Transactivator With Glu/Asp Rich Carboxy-Terminal Domain 2
Crispr/Cas9 Clustered Regularly Interspaced Short Palindromic Repeats/Crispr-associated protein 9
C-term C-terminal end of the protein
Ctx Cortex
DAPK1 Death-Associated Protein Kinase 1
DDX6 DEAD/H (Asp-Glu-Ala-Asp/His) Box Polypeptide 6 (RNA Helicase, 54kD)
DENND1A  |  DENN/MADD Domain Containing 1A (RAB35-activating guanine nucleotide exchange factor)
DMEM     |  Dulbecco’s modified essential medium
DNA      |  Desoxyribo-Nucleic-Acid chain
E        |  Embryonic development day
E2F1     |  E2F Transcription Factor 1
e.g.,    |  Exemplo gratia (for example, in Latin language)
EGF      |  Epidermal growth factor
ELF1     |  E74 Like ETS Transcription Factor 1
Ep300    |  E1A Binding Protein, Histone Acetyltransferase P300
EST      |  Expressed sequence tag
Ex1      |  Exon 1
FCS      |  Fetal calf serum
FTLD     |  Fronto-temporal lobar dementia
Foxj2    |  Forkhead Box Protein J2, a transcriptional activator
g        |  Gram
G3BP2    |  Ras-GTPase Activating Protein SH3 Domain-Binding Protein 2
GLAST    |  Sodium-Dependent Glutamate/Aspartate Transporter 1 (in astrocytes)
GWAS     |  Genome wide association studies
h        |  hour
H&E      |  Hematoxylin and Eosin staining
HBSS     |  Hank’s buffered salt solution
Het      |  Heterozygote
HNRNP    |  Heterogeneous Nuclear Ribonucleoprotein
Kb       |  kilo-base
IRF1     |  Interferon Regulatory Factor 1
ITS      |  Insulin-Transferrin-Selenium
KD       |  Knock-Down
KO       |  Knock-Out
L        |  liter
lncRNA   |  long non-coding RNA
Lsm      |  Like “Smith antigen” protein domain
LSM12    |  Homolog of yeast cytosolic LSM12 protein
LsmAD    |  Lsm-associated domain
mg       |  milliGram
mm       |  milliMeter
mM       |  milliMolar
µg       |  microGram
µL       |  microLiter
µM       |  microMolar
MEF      |  Mouse embryonal fibroblasts
MEM      |  Minimal Essential Medium
MGI      |  Mouse Genome Informatics
mGluR5   |  metabotropic Glutamate Receptor 5
MPL      |  Interactor domain with Myelo-Proliferative Leukemia gene, encodes Thrombopoietin receptor
mRNA     |  Messenger ribonucleic acid
mTORC1   |  Mechanistic “target of rapamycin” complex 1
Neat1    |  Nuclear Paraspeckle Assembly Transcript 1
NFATC2IP |  Nuclear Factor of Activated T-Cells, Cytoplasmic, Calcineurin-Dependent 2 Interacting Protein
ng       |  nanoGram
NHGRI-EBI|  American National Human Genetics Research Institute & European Bioinformatics Institute
N-term   |  N-terminal end of the protein
NUFIP2   |  Nuclear Fragile X Mental Retardation Protein 1 Interacting Protein 2
ORF      |  open reading frame
p        |  probability value for the occurrence of a given observation by chance
PABPC1   |  Poly(A)-binding-protein cytosolic 1
PAM2     |  Poly(A)-binding mediator domain 2
PBS Phosphate buffered saline
PFA Parafomaldehyde
poly(A) tail adenosine-repeat at the 3’ end of mRNA
Poly(I:C) Poly-Iminosinic: Poly-Cytidyl acid chain
Pro Proline
PRD Proline-rich domain
PSD-95 Postsynaptic Density Protein 95 (aka DLK4, Disks Large Homolog 4)
PSP progressive supranuclear palsy (aka Parkinson plus)
Q, 22Q glutamine, repeat of 22 consecutive glutamines
RA Retinoic acid
RT-qPCR Reverse-Transcriptase quantitative Polymerase Chain Reaction
S6R Small ribosomal subunit protein 6
SCA1 Spinocerebellar Ataxia type 1
SCA2 Spinocerebellar Ataxia type 2
SDS Sodium-Dodecyl-Sulfate detergent
s.e.m. Standard error of the mean
Setd5 Histone-Lysine N-Methyltransferase, SET domain containing 5
SG Stress granule
sgRNA Single guide Ribo-Nucleic-Acid chain
Sh2b1 Pro-Rich, PH And SH2 Domain-Containing Signaling Mediator gene
SH3 Src-homology 3 domain
SH-SY5Y Cell line subcloned from SK-N-SH line from 4-year-old female neuroblastoma patient
SITE+3 Selenite, Insulin, Transferrin, Ethanolamine +3
SLC9A3R2 Solute Carrier Family 9 (Sodium/Hydrogen Exchange) Member 3 Regulator 2 (aka NHERF2)
SNP Single nucleotide polymorphism
SPITE Selenite, Pyruvate, Insulin, Transferrin, Ethanolamine +3
SR-B1 Scavenger Receptor Class B Member 1 (aka CD36, Thrombospondin Receptor-Like 1)
Src V-Src Avian Sarcoma (Schmidt-Ruppin A-2) Viral Oncogene
Tbp TATA-binding factor of transcription
TDP-43 TAR (transcription active response element) DNA-Binding Protein 43 (encoded by TARDBP)
Tfap2a Transcription Factor Activator-Protein-2 alpha
TIGM, TX Texas Institute of Genomic Medicine, Texas, United States of America
TPM Transcripts per million (expression unit as normalization method in RNA-sequencing)
Tufm Tu Translation Elongation Factor, Mitochondrial gene
ULK1 Unc-51 Like Autophagy Activating Kinase 1
VPS54 Vacuolar Protein Sorting-Associated Protein 54 (endosomal factor for retrograde transport)
WT Wild-Type

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