UPF3A and UPF3B are redundant and modular activators of nonsense-mediated mRNA decay in human cells

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27 Abstract

28 The paralogous human proteins UPF3A and UPF3B are involved in recognizing mRNAs 29 targeted by nonsense-mediated mRNA decay (NMD). While UPF3B has been demonstrated to support NMD, contradicting reports describe UPF3A either as an NMD activator or inhibitor. 30 Here, we present a comprehensive functional analysis of UPF3A and UPF3B in human cells 31 32 using combinatory experimental approaches. Overexpression or knockout of UPF3A as well 33 as knockout of UPF3B did not detectably change global NMD activity. In contrast, the co-34 depletion of UPF3A and UPF3B resulted in a marked NMD inhibition and a transcriptome-wide 35 upregulation of NMD substrates, demonstrating a functional redundancy between both NMD 36 factors. Although current models assume that UPF3 bridges NMD-activating exon-junction 37 complexes (EJC) to the NMD factor UPF2, UPF3B exhibited normal NMD activity in rescue 38 experiments when UPF2 or EJC binding was impaired. Further rescue experiments revealed 39 partially redundant functions of UPF3B domains in supporting NMD, involving both UPF2 and 40 EJC interaction sites and the central region of UPF3. Collectively, UPF3A and UPF3B serve 41 as fault-tolerant NMD activators in human cells.

43 Introduction

44 Precisely regulated expression of correct gene products is indispensable for eukaryotic life. 45 This is underlined by the existence of several quality control mechanisms for gene expression, 46 one of which is the nonsense-mediated mRNA decay (NMD). NMD is primarily known for its 47 ability to eliminate mature mRNAs that contain a premature termination codon (PTC). Thereby, 48 NMD prevents the synthesis and accumulation of C-terminally truncated proteins, which may 49 possess undesirable and potentially disease-causing properties (Frischmeyer & Dietz, 1999). 50 Although the removal of PTC-containing mRNAs was initially considered the most important 51 function of NMD, later studies showed that NMD plays an important role in the post-52 transcriptional regulation of a substantial part of the transcriptome (He et al., 2003; Lelivelt & 53 Culbertson, 1999; Mendell et al., 2004; Rehwinkel et al., 2005). The importance of the factors 54 involved in NMD is underscored by the severe impact that mutations in components of this 55 machinery have on development in metazoans, up to causing embryonic lethality in mammals 56 (Hwang & Maguat, 2011; Li et al., 2015; McIlwain et al., 2010; Medghalchi et al., 2001; 57 Metzstein & Krasnow, 2006; Weischenfeldt et al., 2008; Wittkopp et al., 2009).

58 The final step of gene expression is the cytoplasmic translation of the mRNA by ribosomes. 59 Previous studies suggested that prolonged ribosome stalling at a termination codon indicates 60 improper translation termination and thereby triggers NMD (Amrani et al., 2004; Peixeiro et al., 61 2012). This could be caused by a long 3' untranslated region (UTR) that increases the distance 62 between the stalled ribosome and the poly(A)-binding protein (PABPC1), which normally 63 promotes proper translation termination (Amrani et al., 2004). Alternatively, NMD can also be 64 activated by any PTC located more than 50-55 nt upstream of the 3'-most exon-exon junction. 65 Transcripts with such a PTC may be transcribed from mutant genes with nonsense mutations 66 but could also be generated by defective or alternative splicing (Kervestin & Jacobson, 2012). 67 The aforementioned 50-55 nt boundary between NMD-activating and NMD-resistant PTCs is 68 determined by the RNA-binding exon junction complex (EJC), which is deposited by the 69 spliceosome 20-24 nt upstream of every spliced exon-exon junction (Le Hir et al., 2000). The

EJCs remain attached on the mature mRNA during export into the cytoplasm, where they are removed by translating ribosomes or the disassembly factor PYM1 (Dostie & Dreyfuss, 2002; Le Hir *et al.*, 2000). If translation terminates prematurely due to the presence of a PTC, EJCs bound downstream of the PTC serve as a marker for the NMD machinery and the initial activation of NMD (Kim *et al.*, 2001; Le Hir *et al.*, 2001).

75 Extensive research over many decades has resulted in a model for EJC-dependent NMD. 76 According to this model, the central factor UPF1 is bound non-specifically to all present 77 transcripts in the cell and is removed from the coding sequence by translating ribosomes (Hogg 78 & Goff, 2010; Hurt et al., 2013; Kurosaki & Maguat, 2013; Zund et al., 2013). If translation 79 terminates prematurely, UPF1 interacts with the stalled ribosome and serves as the anchoring 80 point for the other NMD factors. The presence of a downstream EJC is detected by the 81 interaction of UPF1 with UPF2, which in turn binds to UPF3. The latter can bind directly to the 82 EJC, resulting in a bridged connection of UPF1 to the EJC (Chamieh et al., 2008; Kim et al., 83 2001; Le Hir et al., 2001; Weng et al., 1996). This series of interactions marks the termination 84 codon as premature and stimulates the phosphorylation of N- and C-terminal SQ motifcontaining regions of UPF1 by the kinase SMG1 (Yamashita et al., 2001). In its phosphorylated 85 86 state UPF1 recruits the heterodimer SMG5-SMG7 and/or SMG6, which are responsible for 87 both exoribonucleolytic and endoribonucleolytic degradation of the mRNA, respectively (Boehm et al., 2021; Chen & Shyu, 2003; Lejeune et al., 2003). The endonuclease SMG6 88 89 cleaves the mRNA in close proximity to the PTC, resulting in two mRNA fragments (Eberle et 90 al., 2009) of which the 3' fragment is degraded by the 5'-to-3' exoribonuclease XRN1 (Eberle 91 et al., 2009; Huntzinger et al., 2008).

The protein UPF3 plays an important role in the NMD pathway. As detailed above, its main function is believed to bridge the NMD machinery to the EJC providing a physical link between UPF2 and the EJC (Chamieh *et al.*, 2008; Kashima *et al.*, 2006; Lykke-Andersen *et al.*, 2000; Serin *et al.*, 2001). With its conserved RNA recognition motif (RRM) in the N-terminus, UPF3 can interact with the C-terminal MIF4G (middle portion of eIF4G) domain of UPF2 (Kadlec *et* 97 *al.*, 2004). The association with the EJC-binding site, formed by the core components EIF4A3,

98 MAGOHB and RBM8A, is enabled by a C-terminal sequence referred to as EJC binding motif

99 (EBM) (Buchwald *et al.*, 2010; Gehring *et al.*, 2003; Kim *et al.*, 2001).

In vertebrates two genes encode for two UPF3 paralogues: UPF3A and UPF3B, each of which expressing two different isoforms generated by alternative splicing (Lykke-Andersen *et al.*, 2000; Serin *et al.*, 2001). Both human UPF3 proteins contain the same important domains, but differ in details regarding their interactions. According to previous studies, UPF3A and UPF3B are in constant competition for their binding partner UPF2. However, UPF3B binds tighter to UPF2 than UPF3A and therefore UPF3A gets destabilized, and its protein levels are barely detectable under normal conditions (Chan *et al.*, 2009).

107 Recently, UPF3B was reported to interact with the eukaryotic release factor 3 (eRF3) via the
108 so far uncharacterized middle domain (amino acids (aa) 147-256) (Neu-Yilik *et al.*, 2017). Due
109 to this interaction and binding of the terminating ribosome, it can delay translation termination,
110 which is known to define aberrant termination events and trigger NMD (Amrani *et al.*, 2004;
111 Neu-Yilik *et al.*, 2017; Peixeiro *et al.*, 2012).

112 Due to the different molecular characteristics of the UPF3 paralogs, their exact role in NMD is 113 a long-discussed topic. On the one hand, previous studies showed that UPF3A and UPF3B 114 both trigger degradation of a reporter construct when tethered downstream of a termination 115 codon. Notably, the efficiency of UPF3A to elicit NMD was weaker in comparison to UPF3B, 116 which was attributed to a weaker interaction with the EJC (Kunz et al., 2006; Lykke-Andersen 117 et al., 2000). This would suggest that, at least with respect to their NMD activity, UPF3A and 118 UPF3B serve a similar, perhaps even redundant, function. This notion is supported by the 119 observation that in patients with mutated UPF3B the amount of stabilized UPF3A inversely 120 correlated with the severity of the patients' clinical phenotypes (Nguyen et al., 2012). On the 121 other hand, it was recently reported that loss of UPF3A results in increased transcript 122 destabilization, and UFP3A overexpression in NMD inhibition (Shum et al., 2016). This would rather indicate opposing functions of the two UPF3 paralogs with UPF3A being an antagonistof UPF3B and broadly acting as an NMD inhibitor.

125 In this study, we resolved the controversy about the functions of UPF3A and UPF3B in the 126 NMD pathway using different UPF3 overexpression and knockout (KO) HEK293 cell lines. We 127 found that neither overexpression nor genomic KO of UPF3A resulted in substantial changes 128 of NMD activity or global alterations of the transcriptome. In UPF3B KO cells UPF3A protein 129 levels were upregulated, but NMD activity was maintained at almost normal level. In contrast, 130 the co-depletion of both UPF3 paralogs resulted in a marked NMD inhibition and a global 131 upregulation of PTC-containing transcripts. Moreover, rescue experiments revealed that UPF3 132 proteins have additional functions besides bridging the EJC and the NMD machinery. Taken 133 together, our data support a model of human NMD, in which UPF3A and UPF3B can replace 134 each other and therefore perform redundant functions.

135 **RESULTS**

136 UPF3A overexpression or knockout does not affect NMD efficiency

137 Prior work using different mammalian models and various experimental approaches reached 138 different conclusions regarding the function of UPF3A in NMD (Fig 1A). Therefore, we set out 139 to re-examine the role of UPF3A in human cells by specifically manipulating its expression 140 levels. Under regular conditions UPF3A is barely present in cultured cells, presumably due to 141 its lower binding affinity to the stabilizing interaction partner UPF2 compared to UPF3B, 142 resulting in a rapid turnover of "free" UPF3A (Chan et al., 2009). We hypothesized that 143 increasing the abundance of UPF3A should lead to the stabilization of NMD targets if UPF3A 144 is an NMD inhibitor. To test this hypothesis, we generated Flp-In T-REx 293 (HEK 293) cells 145 inducibly overexpressing FLAG-tagged wildtype UPF3A to high protein levels (Fig 1B). Global 146 analysis of the transcriptome using RNA-seq (Fig EV1A and Datasets EV1-EV3) revealed, 147 except for UPF3A itself, no significant differential gene expression (DGE), differential transcript 148 usage (DTU) or alternative splicing (AS) events upon UPF3A overexpression compared to 149 control conditions (Figs 1C and D). Using these RNA-seq data, we analyzed NMD targets that 150 were previously described to be strongly upregulated in UPF3A overexpressing HeLa cells 151 (Shum et al., 2016). The DGE analysis and visualization of the respective read coverage 152 showed no substantial effects in our setup (Figs 1E and EV1B-E). Furthermore, quantification 153 of differential transcript usage via IsoformSwitchAnalyzeR (Vitting-Seerup & Sandelin, 2019) 154 could neither detect any differences in the global isoform fraction distribution, nor an 155 accumulation of PTC-containing transcripts (Fig EV1F). Collectively, these analyses indicated 156 that UPF3A overexpression in HEK 293 cells does not negatively affect gene expression in 157 general or NMD in particular.

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С

A





| cell line | FLAG- | SMG5 | | | | | | Log2FC |
|-----------|-------|---------------|-------|--|-------|---|---------------|--------|
| WT | - | a int | | | i ili | 1 | n ka saninin. | 0 |
| WT | UPF3A | a int | 1.1.1 | | 6.0 | | din talimit | 0.065 |
| SMG5-201 | | · · · · · · · | | | 1 11 | + | | |

Figure 1- UPF3A overexpression does not affect NMD A Schematic representation of the bridge between UPF1 and the EJC during NMD. Binding of UPF3A instead of the stronger bound UPF3B is discussed to either activate or inhibit NMD. B Western blot analyses after induced expression of FLAG-tagged UPF3A in VT HEK 293 cells. Tubulin serves as control. C Fraction of expressed genes (genes with non-zero counts in DESeq2) were calculated which exhibit individual or combinations of differential gene expression (DGE), differential transcript usage (DTU) and/or alternative splicing (AS) events in VT cells overexpressing UPF3A using the respective computational analysis (cutoffs are indicated). AS and DTU events were collapsed on the gene level. For DGE, p-values were calculated by DESeq2 using a two-sided Wald test and corrected for multiple testing using the Benjamini-Hochberg method. For AS, p-values were calculated by LeafCutter using an asymptotic Chi-squared distribution and corrected for multiple testing using the Benjamini-Hochberg method. using the Benjamini-Hochberg method.

D Volcano plot showing the differential gene expression analyses from the RNA-Seq dataset of WT cells overexpressing UPF3A. The log2 fold change is plotted against the -log10 adjusted p-value (adj. p-value). P-values were calculated by DESeq2 using a two-sided Wald test and corrected for multiple testing using the Benjamini-Hochberg method. OE = overexpression. E Read coverage of SMG5 from WT HEK 293 RNA-seq data with or without induced UPF3A overexpression shown as Integrative Genomics Viewer (IGV) snapshot. Differential gene expression (from DESeq2) is indicated as Log2 fold change (Log2FC) on the right. Schematic representation of the protein coding transcript below.

159 Next, we approached the question of UPF3A function in the opposite way by generating 160 UPF3A knockout (KO) HEK 293 cell lines. Using CRISPR-Cas9 genome editing we isolated 161 three clones that lacked the UPF3A-specific band on the Western blot even after 162 downregulation of UPF3B (Fig 2A). Two clones (14 and 20) were characterized in detail.

163 In both cell lines, the UPF3A genomic locus contained insertions and/or deletions causing 164 frame-shifts and eventually PTCs (Figs EV2A-C). To gain a first impression of the NMD activity 165 in the UPF3A KO cells, the transcript levels of three known exemplary endogenous NMD 166 targets, RSRC2, SRSF2 and ZFAS1 were determined by qPCR (Boehm et al., 2021; Lykke-167 Andersen et al., 2014; Sureau et al., 2001). WT HEK 293 cells treated with SMG6 and SMG7 siRNAs were used as a positive control for severe NMD inhibition (Fig 2B)(Boehm et al., 2021). 168 169 While the absence of UPF3A did not result in abundance changes of SRSF2 and RSRC2 170 NMD-sensitive isoforms (mean log2FC between -0.44 and 0.79), ZFAS1 mRNA levels were 171 slightly decreased in the UPF3A KO cells compared to WT cells (mean log2FC -0.42 and -1.01 172 for UPF3A KO clones 14 and 20, respectively; Fig 2B). However, this effect was not rescued by the (over)expression of transgenic UPF3A, indicating that it is not caused by the lack of 173 174 UPF3A but rather represents random variations in gene expression or clonal effects (Figs 175 EV2D-E). To get a complete overview of the effects of the UPF3A KO, we performed RNA-seq 176 of two UPF3A KO cell lines with or without an additional UPF3B knockdown (KD; Fig EV2F 177 and Datasets EV1-EV3). Initially, we focused on the UPF3A KO cell lines without KDs, for 178 which the global transcriptome analysis revealed that about 4-9 % of the expressed genes are 179 altered (Figs 2C-E). The observation that in the absence of UPF3A more genes were 180 downregulated than upregulated (879 vs. 676) could be an indicator for UPF3A NMD inhibiting 181 properties (Fig 2F). However, the majority of genes with altered expression were clone specific 182 and only 110 genes showed downregulation in both UPF3A KO cell lines (Fig 2F). Investigation 183 of selected targets that were significantly up- or downregulated in both clones revealed that 184 the changes were not rescued after UPF3A overexpression, suggesting that they are UPF3A-185 independent (Figs EV2D-E). Another indication that UPF3A depletion does not generally affect 186 NMD efficiency came from the DTU analysis.





Figure 2 - UPF3A KOs show light NMD-independent transcriptome alterations A Western blot analysis of WT and UPF3A KO cells (clones 4, 14 and 20) with the indicated siRNA treatments. UPF3A and UPF3B protein levels were detected, Tubulin serves as control. B Quantitative RT-PCR of the indicated cell lines treated with the indicated siRNAs for 2 or 6 days. For RSRC2 and SRSF2 the ratio of NMD isoform to canonical isoform was calculated. ZFAS1 expression was normalized to C1orf43 reference. Data points and means are plotted as Log2 fold change (n=3 for RSRC2 and SRSF2, n=4 for ZFAS1). C Fraction of expressed genes (genes with non-zero counts in DESeq2) were calculated which exhibit individual or combinations of differential gene expression (DGE), differential transcript usage (DTU) and/or alternative splicing (AS) events in the indicated conditions using the respective computational analysis (cutoffs are indicated). AS and DTU events were collapsed on the gene level. For DGE, p-values were calculated by DESeq2 using a two-sided Wald test and corrected for multiple testing using the Benjamini-Hochberg method. For AS, p-values were calculated by LeafCutter using an asymptotic Chi-squared distribution and corrected for multiple testing using the Benjamini-Hochberg method. D, E Volcano plots showing the differential gene expression analyses from the indicated RNA-Seq datasets (D UPF3AKO clone 14, E UPF3AKO clone 20). The log2 fold change is plotted against the -log10 adjusted

Benjamini-Hochberg method. **D**, E Volcano plots showing the differential gene expression analyses from the indicated RNA-Seq datasets (D UPF3AKO clone 14, E UPF3AKO clone 20). The log2 fold change is plotted against the -log10 adjusted p-value (padj). P-values were calculated by DESeq2 using a two-sided Wald test and corrected for multiple testing using the Benjamini-Hochberg method F nVenn Diagram showing the overlap of up- or downregulated genes in the UPF3AKO cell lines 14 and 20. Log2 fold change <1 (downregulated) or >1 (upregulated) and adjusted p-value (padj) < 0.05. DGE =

Differential Gene Expression

Although clone 14 showed a minor downregulation of PTC-containing transcripts, which could indicate more active NMD, this effect was not reproducible in the second clone (Fig EV2G). In view of and in combination with the results shown in Fig 1, this strongly suggests that neither the overexpression nor the depletion of UPF3A substantially alters (negatively or positively) the efficiency of NMD. In conclusion, our data argue against a role for UPF3A as a general negative NMD regulator in human cell lines.

194 NMD is functional in the absence of UPF3B

195 Next, we investigated the RNA-seq data of UPF3B knockdowns in the UPF3A KO cells 196 (Datasets EV1-EV3). We observed that this combination resulted in more transcriptome 197 alterations and an increase of PTC-containing isoforms (Figs 2B-C, EV2H and I). Although the 198 UPF3B KD alone could be in principle responsible for this effect, the results could also be an 199 indicator for redundant functions of the two UPF3 paralogs. To explore this hypothesis, we 200 decided to generate UPF3B KO cell lines. Western blot analysis demonstrated that a UPF3B 201 KD is less efficient in reducing the produced protein than the UPF3B KO in the two clones 202 designated as 90 and 91 (Figs 3A and EV3A-B). In addition, we observed a strong upregulation 203 of UPF3A after depletion (KO) or reduction (KD) of UPF3B, as described before (Chan et al., 204 2009). In the absence of UPF3B no changes in the expression of the respective NMD-sensitive 205 isoforms of the NMD-targets RSRC2 and SRSF2 was observed (Fig 3B). This indicates that 206 either UPF3B is not essential for NMD or that another protein is able to compensate for its 207 loss. The most obvious candidate for this function is its own paralog UPF3A, which was also 208 suggested previously to functionally replace UPF3B in NMD. Indeed, knocking down UPF3A in the UPF3B KO cells resulted in the increase of NMD-sensitive RSRC2 and SRSF2 isoforms 209 210 (Fig 3B). Of note, the combination of the UPF3B KO with UPF3A KD showed stronger effects 211 than the previously analyzed UPF3A KO plus UPF3B KD. This is probably caused by the lower 212 KD efficiency of the UPF3B siRNAs which can be observed by comparing the respective 213 protein levels (Fig 2A vs. Fig 3A). We suspect that the remaining UPF3B levels after siRNAmediated UPF3B KD still support NMD. 214

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Figure 3 - Loss of UPF3B does not affect NMD efficiency, only in combination with KD of UPF3A

A Western blot analysis of WT and UPF3B KO cells (clones 90 and 91) combined with the indicated knockdowns. UPF3A and UPF3B (AK-141) protein levels were detected, Tubulin serves as control. B Quantitative RT-PCR of the indicated cell lines with the indicated knockdowns. For RSRC2 and SRSF2 the ratio of NMD isoform to canonical isoform was calculated. Data points and means are plotted as Log2 fold change (n=3).

C Fraction of expressed genes (genes with non-zero counts in DESeq2) were calculated which exhibit individual or combinations of differential gene expression (DGE), differential transcript usage (DTU) and/or alternative splicing (AS) events in the indicated conditions using the respective computational analysis (cutoffs are indicated). AS and DTU events were collapsed on the gene level. For DGE, p-values were calculated by DESeq2 using a two-sided Wald test and corrected for multiple testing using the Benjamini-Hochberg method. For DTU, p-values were calculated by IsoformSwitchAnalyzeR using a DEXSeq-based test and corrected for multiple testing using the Benjamini-Hochberg method. For AS, p-values were calculated by LeafCutter using an asymptotic Chi-squared distribution and corrected for multiple testing using the Benjamini-Hochberg method.

DE Volcano plots showing the differential transcript usage (via IsoformSwitchAnalyzeR) in various RNA-Seq data. Isoforms containing GENCODE (release 33) annotated PTC (red, TRUE), regular stop codons (blue, FALSE) or having no annotated open reading frame (grey, NA) are indicated. The change in isoform fraction (dIF) is plotted against the -log10 adjusted p-value (adj.p-value). Density plots show the distribution of filtered isoforms in respect to the dIF, cutoffs were |dIF| > 0.1 and adj. p-value < 0.05. P-values were calculated by IsoformSwitchAnalyzeR using a DEXSeq-based test and corrected for multiple testing using the Benjamini-Hochberg method.

216 To gain more transcriptome-wide information, we performed RNA-seq of the UPF3B KO clone 217 90, with and without UPF3A siRNA treatment (Fig EV3C and Datasets EV1-EV3). The global 218 effects detected in the RNA-seg data correlated well with the NMD inhibition seen for single 219 targets (Figs 3C and EV3D-G). Analysis of the differential transcript usage revealed an 220 upregulation of transcripts annotated with a PTC only in the UPF3B KO cells with additionally 221 downregulated UPF3A (Figs 3D and E). This increase of NMD-sensitive transcripts could not 222 be observed in the plain UPF3B KO cells with UPF3A naturally upregulated. All together this 223 data strongly suggested at least partial redundancy of UPF3A and UPF3B, since KO of only 224 one paralog was irrelevant for NMD functionality. Furthermore, we were able to show that also 225 in the absence of UPF3B an overexpression of UPF3A has no negative effects on NMD 226 efficiency, supporting the previous conclusion of UPF3A not being a negative NMD regulator 227 (Figs EV3H and I).

228 Stronger NMD impairment in UPF3A-UPF3B double KO cells

229 Considering that UPF3A and UPF3B seemingly carry out redundant functions, we decided to 230 create a cell line completely lacking both paralogs and aimed to generate UPF3A-UPF3B 231 double KO cells (UPF3 dKO). These cells should show stronger effects than the combination 232 of a KO and a KD, since residual amounts of protein were typically still detected after siRNA 233 treatment. Using the UPF3B-KO clone 90 as parental cell line, two potential UPF3 dKO clones 234 1 and 2 were generated (Fig 4A), which differed in the guide RNAs used to target exon 1 of 235 UPF3A. We confirmed that both cell lines contained frame shift-inducing insertions/deletions 236 at the respective positions in the UPF3A gene (Figs 4B, EV4A and B). We first explored by 237 aPCR how strongly the dKO affected NMD (Fig 4C). For all three tested genes, the expression 238 of the NMD-sensitive isoform was further increased compared to the previously used 239 combination of UPF3B KO with additional UPF3A KD. Of note, the NMD inhibitory effect 240 observed in the dKOs became more pronounced after UPF3B siRNA transfection, suggesting 241 that low levels of residual UPF3B protein were still present in the dKO cells (Fig EV4C).



Figure 4 - KO of both UPF3 paralogs results in strongly impaired NMD A Western blot analysis of WT, UPF38 KO and UPF3A-UPF38 doubleKO cells (clones 1 and 2). UPF3A and UPF3B protein levels were detected, Tubulin serves as control. B Schematic depiction of the insertion/deletion in the UPF3A open reading frame resulting in the additional UPF3AKO in the UPF3B KO clone 90 generating UPF3 dKO clones. C Quantitative RT-PCR of the indicated samples with the indicated KDs. For RSRC2 and SRSF2 the ratio of NMD isoform to canonical isoform was calculated. ZFAS1 expression was normalized to C1orf43 reference. Data points and means are plotted as Log2 fold change (n=3). D Schematic overview of the globin reporter constructs and their functional elements. E Northern blot analysis of globin reporter and xrFrag. Ethidium bromide stained 28S and 18S rRNAs are shown as controls. Quantification results are shown as data points and mean (n=3).

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243 The expression levels of endogenous NMD substrates could be influenced by transcription 244 rates or other indirect effects, which could lead to over- or underestimating NMD inhibition. 245 Therefore, we investigated the NMD efficiency in the dKO cells using the well-established β -246 globin NMD reporter. To this end, we stably integrated β -globin WT or PTC39 constructs in 247 WT and UPF3 dKO cells (Fig 4D). These reporters also contained XRN1-resistant sequences 248 (xrRNAs) in their 3' UTRs, which allowed us to analyze not only the degradation of the full-249 length reporter mRNA but also to quantify decay intermediates (called xrFrag) (Boehm et al., 250 2016; Voigt et al., 2019). The PTC39 mRNA was efficiently degraded and a strong xrFrag 251 observed in WT cells, whereas in dKO cells the PTC39 reporter accumulated to high levels 252 (72% compared to the WT mRNA), which was accompanied by a decrease in the amount of 253 xrFrag. (Fig 4E, lane 2 vs. lane 4). In line with the previous observations, this result indicated 254 a strong decrease of NMD activity upon the KO of both UPF3 paralogs using a robust NMD 255 reporter pair expressed independently of endogenous NMD substrates, supporting the 256 functional redundancy of UPF3A and UPF3B in NMD.

257 To establish transcriptome-wide insights into UPF3A and UPF3B function, we carried out RNA-258 seq for both dKO clones, which was combined with and without UPF3B KD treatment to 259 eliminate as many of the potentially present remaining UPF3B proteins (Fig EV5A and 260 Datasets EV1-EV3). Differential gene expression analysis showed that nearly three times as 261 many genes were upregulated than downregulated in both dKO cells (Figs 5A and EV5B). This 262 is consistent with the redundant role of UPF3B and UPF3A as supporting NMD factors. The 263 considerable overlap between both clones also suggests that we identified high-confidence 264 UPF3 targets. Furthermore, 890 of these gene were also significantly upregulated in previously 265 generated SMG7 KO plus SMG6 KD data (Fig EV5C, ref data: (Boehm et al., 2021)) indicating 266 that these are universal NMD-targets and not specific to a certain branch of the NMD pathway. 267 In addition to DGE, subsets of the expressed genes showed changes in alternative splicing or/and differential transcript usage (Fig 5B). In total, 14-16 % of the global transcriptome 268 269 showed single or combined changes (DGE, DTU and/or AS) and up to 20% when the cells 270 were treated with an additional UPF3B KD.



exon inclusion

3' UTR intron

Figure 5 - RNA-seq reveals strong global upregulation of NMD-sensitive targets upon UPF3 dKO in HEK293 cells A nVenn Diagram showing the overlap of up- or downregulated genes in the UPF3 dKO cell lines 1 and 2. Log2 fold change <-1 (downregulated) or >1 (upregulated) and adjusted p-value (padj) < 0.05. DGE =

Bifferential Gene Expression. B Fraction of expressed genes (genes with non-zero counts in DESeq2) were calculated which exhibit individual or combinations of differential gene expression (DGE), differential transcript usage (DTU) and/or alternative splicing (AS) events in the indicated conditions using the respective computational analysis (cutoffs are indicated). AS and DTU events were collapsed on the gene level. For DGE, p-values were calculated by DESeq2 using a two-sided Wald test and corrected for multiple testing using the Benjamini-Hochberg method. For DTU, p-values were calculated by IsoformSwitchAnalyzeR using a DEXSeq-based test and corrected for multiple testing using the Benjamini-Hochberg method. For AS, p-values were calculated by LeafCutter using an asymptotic Chi-squared distribution and corrected for multiple testing using the

corrected for multiple testing using the Benjamini-Hochberg method. For AS, p-values were calculated by LeatCutter using an asymptotic Chi-squared distribution and corrected for multiple testing using the Benjamini-Hochberg method. C Volcano plots showing the differential transcript usage (via IsoformSwitchAnalyzeR) in various RNA-Seq data. Isoforms containing GENCODE (release 33) annotated PTC (red, TRUE), regular stop codons (blue, FALSE) or having no annotated open reading frame (grey, NA) are indicated. The change in isoform fraction (dIF) is plotted against the -log10 adjusted p-value (adj.p-value). Density plots show the distribution of filtered isoforms in respect to the dIF, cutoffs were |dIF| > 0.1 and adj.p-value < 0.05. P-values were calculated by IsoformSwitchAnalyzeR using a DEXSeq-based test and corrected for multiple testing using the Benjamini-Hochberg method. D Read coverage of SRSF2 from the indicated RNA-seq sample data with or without UPF3A siRNA treatment shown as Integrative Genomics Viewer (IGV) snapshot. The canonical and NMD-sensitive isoforms are schematically indicated below. Quantification of isoforms by IsoformSwitchAnalyzeR (right).

С

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In agreement with NMD inhibition in the dKOs, we saw that many transcripts containing a PTC were up-regulated, while the corresponding transcripts without a PTC were down-regulated. (Figs 5C and EV5D). Under these conditions, the IGV snapshot of the NMD-target SRSF2 showed NMD-inducing exon inclusion and 3' UTR splicing events, which were not visible in combined UPF3B KO/UPF3A KD cells (Fig 5D). Collectively, the RNA-seq data support the previously observed strong NMD inhibition in response to the complete absence of both UPF3 paralogs and hence their proposed redundancy.

279 UPF3A supports NMD independent of a bridge function

280 Next, we aimed to analyze whether the severe effects in the dKOs are at least partly due to 281 the loss of a protein-protein interaction bridge between UPF2 and the EJC, while the presence 282 of UPF3A in the UPF3B KOs preserved this function ensuring NMD functionality. Therefore, 283 we expressed FLAG-tagged UPF2 in WT, UPF3B KO and UPF3A-UPF3B dKO cells and 284 analyzed the UPF2 interactome using mass spectrometry (Dataset EV4). Consistent with the 285 previously described interaction partners, we found many NMD factors as well as EJC proteins 286 in the UPF2 interactome in WT cells (Fig 6A). Contrary to our expectation, the three EJC core 287 components (EIF4A3, RBM8A, MAGOHB) barely co-precipitated with UPF2 in the absence of 288 UPF3B (compared to control: log2 FC = 0.58, 0.63 and 0.74, respectively; Fig 6B) and were 289 therefore strongly decreased in comparison to the WT cells (Fig EV6A). Hence, the UPF2-290 bound UPF3A was unable to establish a stable interaction with the EJC. Surprisingly, in the 291 UPF3B KO cells the EJC-associated CASC3 still showed relatively high levels of co-292 precipitation (log2 FC = 4.42), which therefore appears to be independent of the interaction 293 with the other EJC components. In the dKOs all interactions with EJC proteins including 294 CASC3 were completely lost (Figs 6C and EV6B). The latter was also observed in a 295 comparable approach employing stable isotope labeling with amino acids in cell culture 296 (SILAC) to analyze the UPF2 interactome in the WT and dKO cells (Dataset EV5). All EJC core components that were highly co-precipitated in WT cells were lost in the absence of both 297 298 UPF3 paralogs (Figs EV6C-E).



Figure 6 - Interaction of UPF3A and UPF3B with the EJC is dispensable to elicit NMD A-C Volcano plots of label free mass spectrometry-based analysis of the interaction partners of UPF2 in WT cells treated with control siRNAs and the UPF3B KO clone 90 and dKO clone 1 both treated with siRNAs targeting UPF3B (n = 4 biologically independent samples). (A) FLAG-UPF2 in WT against FLAG-GST control in WT cells, (B) UPF2 in 3B KO cells against FLAG control in WT cells, (C) UPF2 in dKO cells against FLAG control in WT cells. Points labeled in purple indicate NMD factors; points labeled in turquoise indicate EJC components. Cut offs: Log2 fold change ≥ 0 D Schematic representation of the UPF3B protein domains and the respective functions. Below are the mutated rescue constructs and their respective abstract placeholders. E Western blot analysis of WT and UPF3B KO clone 90 with Luciferase and UPF3AKDs respectively. Monitored expression of the FLAG-tagged UPF3B rescue construct shown in (D). Rescue construct protein levels were detected with anti-UPF3B (KAL-141) antibodies. Tubulin serves as control. F Quantitative RT-PCR of the samples from (E). For RSRC2 and SRSF2 the ratio of NMD isoform to canonical isoform was calculated. ZFAS1 expression was normalized to C1orf43 reference. Data points and means are plotted as log2 fold change (n=3).

300 Partially redundant functions of UPF3B domains are required for NMD

With regard to the surprising observation that UPF3A apparently elicits NMD without interacting with the EJC, we aimed to investigate the molecular features required by UPF3 to support NMD via rescue experiments. In principle, the UPF3 dKO cells represent an ideal system for this approach. However, apart from the residual amounts of UPF3B that were still expressed, we noticed that the UFP3 dKOs were able to upregulate the expression of a shortened UPF3B variant after long-term cultivation.

307 Since this phenomenon did not occur in the single UPF3B KOs, we generated stable UPF3B 308 KO cell lines expressing various inducible UPF3B constructs with individual or combined 309 binding site mutations (Figs 6D and E and EV6F). Transfection of these cells with UPF3A 310 siRNAs resulted in the robust depletion of UPF3 for the analysis of the rescue activities of 311 individual UPF3B variants. Considering the established role of UPF3B as a bridge between 312 UPF2 and the EJC, which we validated in the mass spec analysis, it was surprising to see that 313 disruption of either of these interactions did not affect UPF3B's rescue capacity (Fig 6F, lane 314 3 vs. lanes 4 and 5). This is partially consistent with the observed functional NMD in UPF3B 315 KOs, despite the apparent inability of UPF3A to form a bridge between UPF2 and the EJC 316 (Figs 3C and 6B). However, mutating both UPF3B binding sites (disrupting UPF2 and EJC 317 binding) largely inactivated the NMD-related function of UPF3B.

318 It was recently reported that EJC-bound or free UPF3B can interact with the eukaryotic release 319 factor 3 (eRF3) via the so far uncharacterized middle domain (aa 147-256) (Neu-Yilik et al., 320 2017). With this interaction and binding of the terminating ribosome, UPF3B can delay 321 translation termination, which defines aberrant termination events and triggers NMD (Amrani 322 et al., 2004; Neu-Yilik et al., 2017; Peixeiro et al., 2012). To investigate the impact of this 323 interaction on NMD, we created UPF3B variants lacking that specific middle domain or 324 combined the deletion with the previously used interaction mutations (Fig 6D and E). We 325 observed a similar pattern as the UPF3B mutants examined in the previous experiment: when 326 only the middle domain was deleted, UPF3B was able to rescue NMD comparable to the WT

327 protein (Fig 6F lane 7). In combination with a mutation in the UPF2- or the EJC binding site its 328 function in NMD was severely impaired (lanes 8 and 9). This suggests that if the classic bridge 329 formation is inhibited by removing either of the interaction sites, UPF3B relied on the function 330 carried out by the uncharacterized middle domain.

As further support for the NMD promoting effect of UPF3A on NMD, we performed a rescue experiment in the above used UPF3B KO UPF3A KD cells (Figs 7A and B). Expression of the siRNA insensitive UPF3A construct did not only restore NMD functionality, it was even more efficient than the UPF3B construct (Fig 7C lane 3 vs. lane 5), underlining our previous statement: UPF3A supports and elicits NMD comparably to its paralog UPF3B.

Also comparable to UPF3B, UPF3A has a second naturally occurring isoform but instead of skipping exon 8 (like UPF3B) it excludes its fourth exon. This isoform is transcribed in approximately one third of the cases (Fig EV6G) in WT HEK 293 cells but cannot be detected on protein levels. We were interested, whether this isoform was as potent to elicit NMD as the full-length construct. Expression of the exon 4 UPF3A deletion construct in the UPF3 depleted cells showed no rescue (Fig 7C lane 4). Hence, exon 4 must encode for an essential region required for the bridge-independent function of UPF3A.

343 In view of these observations and the close proximity of exon 4 (124-157) to the middle domain 344 (147-256), we decided to investigate which effect the deletion of the homologous exon 4 in the 345 paralog UPF3B has. The UPF3B Ae4 construct behaved like the corresponding UPF3A 346 construct and showed no NMD rescue activity (lane 6). However, the expression of the N-347 terminus of UPF3B was able to restore NMD comparably to the WT protein (lane 7). This is 348 consistent with all our previous findings, since the first 279 amino acids contain the UPF2 349 binding site as well as the middle domain, which was shown to be sufficient to elicit NMD (Fig 350 6F lane 5). Due to the fact that the C-terminus contains only one interaction site and lacks exon 351 4, its incapability to rescue NMD is in line with our previous experiments. Overall, our results 352 identify exon 4 of UPF3 as a previously unnoticed region that is essential for its function in 353 NMD.





Figure 7 - WT UPF3A can rescue NMD in full extent. Deletion on exon 4 disrupts functionality in both paralogs. A Schematic representation of the UPF3A and UPF3B protein domains. Below are the respective mutated rescue constructs. B Western blot analysis of WT and UPF3B KO clone 90 with Luciferase and UPF3AKDs respectively. Monitored expression of the FLAG-tagged UPF3A and UPF3B rescue construct shown in (A). Rescue construct protein levels were detected with anti-FLAG, anti-UPF3A and anti-UPF3B (AK-141) antibodies. Tubulin serves as control. C Quantitative RT-PCR of the samples from (B). For RSRC2 and SRSF2 the ratio of NMD isoform to canonical isoform was calculated. Data points and means are plotted as log2 fold change (n=3).

355 Discussion

356 Methodological advances - be it improved analytics or novel experimental approaches - can 357 help to find new answers to old biological problems. Equipped with powerful new molecular 358 biology methods, we set out to answer the question which functions the two UPF3 paralogs 359 carry out in human cells. Since the initial description of mammalian UPF3A and UPF3B (Lykke-360 Andersen et al., 2000; Serin et al., 2001), many researchers have been engaged in determining 361 the function and work distribution of these two proteins in NMD. It became clear relatively early 362 that UPF3A and UPF3B can both interact with UPF2 as well as the EJC and activate NMD 363 (Gehring et al., 2003; Kim et al., 2001; Lykke-Andersen et al., 2000; Serin et al., 2001). But 364 significant differences in these interactions and the amounts of UPF3A and UPF3B proteins 365 were also found, leading to the hypothesis that UPF3B is the central player and UPF3A more 366 its backup (Chan et al., 2009; Kunz et al., 2006). The investigations were further intensified 367 upon the discovery that mutations in the human UPF3B gene lead to various forms of mental 368 disorder (Nguyen et al., 2014). Disease severity seems to be dictated by the amount of UPF3A 369 present, suggesting again a compensatory mechanism and redundant functions of UPF3B and 370 UPF3A (Nguyen et al., 2012).

371 Later, it was reported that in mouse cells UPF3B is an NMD activator and UPF3A is an NMD 372 inhibitor (Shum et al., 2016). Accordingly, removal of UPF3A resulted in enhanced NMD, 373 whereas removal of UPF3B inhibited NMD. Recently, another function of UPF3B was 374 discovered, namely that it is involved in different phases of translation termination (Neu-Yilik 375 et al., 2017). UPF3B not only interacts with release factors, but also slows down translation 376 termination and promotes dissociation of post-termination ribosomal complexes. Whether 377 these functions of UPF3B in translation termination are related to its function in NMD has not 378 been clarified to date.

Although our results cannot fully answer this last question, we have more or less definite answers for the functions of the two proteins UPF3A and UPF3B in NMD. All our data support the notion that the presence of UPF3B or UPF3A is sufficient to maintain NMD activity in

382 human cells. First, we see at most a weak inhibition of NMD in our HEK 293 UPF3B KO cells. 383 However, this does not necessarily mean that all cell types display full NMD activity after a KO 384 of UPF3B. Instead, it is conceivable that HEK 293 cells are just particularly robust against the 385 UPF3B depletion or particularly efficient at the compensatory upregulation of UPF3A. Likewise, 386 we see no NMD inhibition in cells overexpressing UPF3A or NMD "boosting" in UPF3A KO 387 cells. Only when we deplete in UPF3A or UPF3B KO cell lines the respective other protein by 388 RNAi or genomic KO, NMD efficiency substantially decreases. It should be noted that the 389 effects of a KO on NMD activity are typically stronger than those of a KD, which is consistent 390 with our earlier observations (Boehm et al., 2021; Gerbracht et al., 2020). Although our data 391 clearly argue against an NMD-inhibitory function of UPF3A, experimental differences exist 392 between our work and the work describing the NMD inhibition by UPF3A. While we have used 393 human HEK293 cells, the results of Shum et al. were obtained in P19 mouse cells and mouse 394 embryonic fibroblasts (MEFs). Both, the different organisms and the different types of cells 395 could have influenced the results. Interestingly, in a parallel manuscript, Yi et al. find that 396 mouse UPF3A can rescue NMD in human HCT116 UPF3A-UPF3B dKO cells (Yi et al., 2021). 397 Although these results were obtained in a heterologous context, mouse UPF3A does not 398 appear to be a general NMD inhibitor.

399 The different KO cells that we have generated in the course of this project enabled us to 400 conduct experiments that went beyond investigating UPF3-dependent NMD substrates in 401 human cells. Specifically, we were able to study the composition of NMD complexes without 402 UPF3B or both UPF3 proteins and to carry out rescue experiments with different UPF3A and 403 UPF3B protein variants. Not entirely unexpected, we observed that in the absence of UPF3A 404 and UPF3B, the interaction between UPF2 and the EJC is lost. This bridging by UPF3 between 405 UPF2-containing NMD complexes and the EJC was previously considered to be essential for 406 NMD. However, two observations argue against UPF3 being mainly a bridging protein. First, 407 UPF3B mutants that cannot interact with either the EJC or UPF2 fully rescue NMD. Only when 408 both interaction sites were mutated, UPF3 lost its NMD function. This indicates that the 409 interaction with one of the two interaction partners is sufficient to maintain NMD. Second, we

410 observed that not only in UPF3 double-KO cells, but also in UPF3B KO cells, the bridge 411 between UPF2 and the EJC was lost. Although quite surprising at first glance, this is in good 412 agreement with previous results showing that the interaction between the EJC and UPF3A is 413 substantially weaker than that between the EJC and UPF3B (Kunz et al., 2006). Indeed, earlier 414 structural data also argue against a bridging function of UPF3. In the cryo-EM structure of an 415 EJC-UPF3-UPF2-UPF1 complex, UPF1 did not face towards a possible terminating ribosome 416 in the 5' direction, but instead in 3' direction (Melero et al., 2012). Therefore, one could 417 conclude that the interactions between all these proteins do not take place at a single time 418 point during NMD.

419 This raises the question of what UPF3 function is essential for NMD, if it is not its bridging 420 function? Our rescue experiments showed that the middle domain of UPF3B cooperates with 421 the UPF2- and the EJC interaction sites, i.e., its deletion in combination with one other mutation 422 inactivates UPF3B. The middle domain has been described to mediate the interaction of 423 UPF3B with release factor 3 (RF3), but this interaction has not been demonstrated for UPF3A 424 (Neu-Yilik et al., 2017). So, if UPF3A cannot interact with RF3 and also binds weaker to the 425 EJC as noted above, it should be functionally inactive. However, we see no obvious difference 426 between UPF3B and UPF3A in the rescue experiments. These and other observations can, in 427 our view, only be explained with more complex models, which must also consider non-linear 428 relationships and potential auxiliary functions of certain regions of UPF3 (Fig 8).

429 Since the interactions of UPF3 are essential only in combination with each other, we propose that UPF3A and UPF3B exert multiple functions at different time points of NMD and in 430 431 association with different complexes. We only consider here the previously described 432 interactions of UPF3 with UPF2, the EJC and the release factor 3 (Fig 8B). While it was 433 previously described that UPF3B interacts better than UPF3A with UPF2 (Chan et al., 2009), 434 we find both proteins in the FLAG-UPF2 IP. The amount of UPF3A does not seem to increase 435 when UPF3B is depleted, which could be due to the overexpression of UPF2 in our 436 experimental system. The interaction of UPF3 and UPF2 is also conserved in yeast and is thus



Figure 8 - Model for potentially independent functions of UPF3 in NMD. A Both UPF3 paralogs UPF3A and UPF3B can elicit NMD and trigger mRNA degradation. We propose a new model where UPF3B exerts multiple functions at different timepoints during NMD. B Schematic overview of UPF3B domain structure and the postulated functions. Conservation between UPF3A (UniProt ID: Q9H1J1-1) and UPF3B (UniProt ID: Q9BZI7-2) proteins as calculated via Clustal Omega

and Jalview is shown. C Via the RRM-like domain in the N-termination delay. The interaction with UPE2C, which might be important for the assembly of an NMD-inducing complex. The middle domain was ascribed to be responsible for the interaction with eRF3 and potential translation termination delay. The interaction with the EBC via the EBM could stabilize the complex by preventing its interaction with the disassembly factor PYM1. Only if two out of the three functions are ensured, NMD can pursue and degrade the targeted mRNA.

438 likely to have functional significance. However, a UPF3B mutant in which binding to UPF2 was439 inactivated rescues NMD better than the UPF3B WT.

The interaction of the middle domain of UPF3B with RF3 has only recently been described (Neu-Yilik *et al.*, 2017). Again, we find that removing only the middle domain does not substantially inhibit NMD. In combination with an inactivation of the UPF2 binding, the deletion of the middle domain leads to a complete inhibition of the NMD in the rescue assay. This could also happen when exon 4 of UPF3B or UPF3A is removed, which is located at the junction between the UPF2 binding domain and the middle domain. Therefore, UPF3A Δexon4 would be a naturally occurring, NMD-inactive variant of the UPF3 proteins.

447 How might the different regions and domains communicate with each other and regulate the 448 function of UPF3 in NMD? The function of the middle domain in relation to NMD has not yet 449 been investigated. It is conceivable that UPF3B plays a minor role in translation termination 450 and that the events that trigger NMD can also occur without the middle domain - potentially 451 with a delay (Figure 8c). Therefore, the deletion of the middle domain could be tolerated in 452 isolation but would become fatal in combination with other mutations that impair additional 453 functions. With regard to the EBM, we propose that its binding stabilizes the EJC, for example 454 by preventing the interaction of the EJC with PYM1 (Fig 8C). PYM1 is a known EJC 455 disassembly factor and binds to the EJC at a surface area that overlaps with the EBM binding 456 site (Bono et al., 2004; Buchwald et al., 2010; Gehring et al., 2009). We suggest that in cells 457 rescued with the Δ EBM mutant EJCs are more readily dissociated from the mRNA in the 458 cytoplasm. The concomitant loss of UPF3B's termination function (Δ middle domain, Δ EBM) 459 would have a dramatic effect on NMD efficiency, because NMD would be initiated too slowly. 460 Likewise, the interaction of UPF3 with UPF2 might be important for the assembly of an NMD-461 inducing complex (Fig 8C) that needs to be timed with translation termination and that leads 462 to NMD activation only in the presence of the EJC. Although these suggestions may not 463 accurately reflect the molecular events during NMD, they illustrate possible functions of the 464 domains of UPF3, particularly in relation to their NMD-inactive combinations (Fig 8). Overall,

465 our observations fit well with a "synthetic lethal" model in which inactivation of any two domains
466 together disrupts UPF3 activity.

467 One factor whose function needs to be examined in more detail in the context of UPF3 is 468 CASC3. Our mass spectrometry analysis shows that CASC3 immunoprecipitates very well 469 with UPF2 in wildtype cells. CASC3 still partially precipitates with UPF2 in UPF3B KO cells, 470 although the interaction of the other EJC factors is reduced to background levels. In previous 471 work, we observed that UPF3B interacts less well with the EJC when CASC3 is knocked out 472 (Gerbracht et al., 2020). This indicates that an interaction between CASC3 and the UPF3 473 proteins exists that is not well understood so far. What kind of interaction this is and what 474 function it has will be interesting to address in future experiments.

475 Our own work and the work of Yi et al. have re-examined the functions of the human UPF3 476 paralogues UPF3A and UPF3B (Yi et al., 2021). Together, the studies confirmed some 477 previous findings and disconfirmed others, thereby successfully (re-)defining the role of UPF3 478 proteins in human cells. A few questions remain unanswered and need to be addressed in the 479 future, for example how exactly the middle domain supports NMD. As described above, our 480 results have implications for the understanding of the NMD mechanism, as they are 481 incompatible with, and thus exclude, certain models of NMD. In addition, they may also help 482 to better understand the link between UPF3B and intellectual disability and which domains of 483 UPF3A modulate the severity of the disease and may therefore be potential targets for therapy.

484

486 Materials and Methods

487 Cell Culture

Flp-In-T-REx-293 (human, female, embryonic kidney, epithelial; Thermo Fisher Scientific, RRID:CVCL_U427) cells were cultured in high-glucose, GlutaMAX DMEM (Gibco) supplemented with 9% fetal bovine serum (Gibco) and 1x Penicillin Streptomycin (Gibco). The cells were cultivated at 37°C and 5% CO2 in a humidified incubator. The generation of knockout and stable cell lines is described below and all cell lines are summarized in Dataset EV6.

494 siRNA-mediated knockdowns

For reverse transfection the cells were seeded at a density of 2.5x10⁵ cells per well. The transfection solution contained 2.5 µl Lipofectamine RNAiMAX and 60 pmol of the respective siRNAs. For the UPF3A and UPF3B knockdowns 30 pmol of both belonging siRNAs were used. In preparation for mass spectrometry, 2.5x10⁶ were reverse transfected in 10 cm plates using 6.25 µl Lipofectamine RNAiMAX and 150-200 pmol siRNA (or half of it for each of the two siRNAs for UPF3A or UPF3B). All siRNAs used in this study are listed in Dataset EV6.

501 Plasmid transfection

502 For each stable transfection 2.5-3.0x10⁵ cells were seeded one day prior transfection in 6-503 wells. To express the N-terminally FLAG-tagged protein constructs and reporter mRNAs for 504 northern blotting, they were stably integrated using the PiggyBac (PB) Transposon system with 505 the cumate-inducible PB-CuO-MCS-BGH-EF1-CymR-Puro vector. This vector was modified 506 from the original vector (PB-CuO-MCS-IRES-GFP-EF1α-CymR-Puro (System Biosciences)) 507 by replacing the IRES-GFP cassette with a BGH polyA signal. Per well 1.0 µg of the respective 508 PB vector and 0.8 µg PB Transposase were transfected using a calcium phosphate-based 509 system with BES buffered saline (BBS). Additionally, 0.5 µg of pCI-maxGFP was transfected 510 as a visual feedback for transfection efficiency. 48 h later, the cells were pooled in 10 cm plates 511 and selected for positive cells by incubation in media containing 2 µg/ml puromycin for a week. 512 To induce expression of the constructs, 30 µg/ml cumate was added and the cells were 513 harvested after 72h for continuing experiments.

514 The mRNA reporter constructs β -globin WT and β -globin PTC are available on Addgene (IDs

515 108375-108376). All vectors used in this study are listed in Dataset EV6.

516 Generation of knockout cells using CRISPR-Cas9

517 The knockouts were performed using the Alt-R CRISPR-Cas9 system (Integrated DNA 518 Technologies) and reverse transfection of a Cas9:guideRNA ribonucleoprotein complex using 519 Lipofectamine RNAiMAX (Thermo Fisher Scientific) according to the manufacturer's protocol. 520 (Integrated DNA Technologies) to target UPF3B The crRNA sequence was 521 /AltR1/rArGrArUrArArGrCrArGrGrArUrCrGrCrArArCrArGrUrUrUrUrArGrArGrCrUrArUrGrCrU/ 522 AltR2/. For UPF3A the crRNA sequences were /AltR1/rCrCrGrCrArArCrCrGrGrArGrGrArC 523 rGrArArGrUrGrUrUrUrUrArGrArGrCrUrArUrGrCrU/AltR2/ for clone 1 and /AltR1/rGrCrGrGr 524 UrGrGrArArCrUrGrCrArCrUrUrCrUrArGrUrUrUrUrArGrArGrCrUrArUrGrCrU/AltR2/ for clone 525 2. Reverse transfection was performed on 1.5×10⁵ cells per crRNA in 12-well plates. 48 h after 526 transfection the cells were trypsinized, counted and seeded at a mean density of a single cell 527 per well in 96-well plates. Cell colonies originating from a single clone were then screened via 528 Western blot and genome editing of UPF3A and UPF3B was analyzed on the genomic level 529 via DNA extraction and Sanger sequencing. Alterations on the transcript level were analyzed via RNA extraction followed by reverse transcription and Sanger sequencing. 530

531 DNA and RNA extraction

532 Genomic DNA extraction using QuickExtract DNA Extraction Solution (Lucigen) was 533 performed according to manufacturer's instruction. For RNA extraction cells were harvested 534 with 1 ml RNAsolv reagent (Omega Bio-Tek) per 6 well and RNA was isolated according to 535 manufacturer's instruction, with the following changes: instead of 200 µl chloroform, 150 µl 1-536 Bromo-3-chloropropane (Sigma-Aldrich) was added to the RNAsolv. Additionally, in the last 537 step the RNA pellet was dissolved in 20 µl RNase-free water by incubating for 10 min on a 538 shaking 65 °C heat block.

539 Western blotting

540 SDS-polyacrylamide gel electrophoresis and immunoblot analysis were performed using 541 protein samples harvested with RIPA buffer (50 mM Tris/HCl pH 8.0, 0.1% SDS, 150 mM NaCl, 542 1% IGEPAL, 0.5% deoxycholate) or samples eluted from Anti-FLAG M2 magnetic beads. For 543 protein guantification, the Pierce Detergent Compatible Bradford Assay Reagent (Thermo 544 Fisher Scientific) was used. All antibodies used in this study are listed in Dataset EV6. 545 Detection was performed with Western Lightning Plus-ECL (PerkinElmer) or ECL Select 546 Western Blotting Detection Reagent (Amersham) and the Vilber Fusion FX6 Edge imaging 547 system (Vilber Lourmat).

548 Semi-quantitative and quantitative reverse transcriptase (RT)-PCR

549 Reverse transcription was performed with 1-4 µg of total RNA in a 20 µl reaction volume with 550 10 µM VNN-(dT)₂₀ primer and the GoScript Reverse Transcriptase (Promega). For the semi-551 quantitative end-point PCRs the MyTag Red Mix (Bioline) was used. Quantitative RT-PCRs 552 were performed with the GoTag qPCR Master Mix (Promega), 2% of cDNA per reaction, and 553 the CFX96 Touch Real-Time PCR Detection System (Bio-Rad). Each biological replicate was 554 repeated in technical triplicates and the average Ct (threshold cycle) value was measured. 555 When isoform switches were measured, values for NMD sensitive isoforms were normalized 556 to the canonical isoforms to calculate ΔCt . For differentially expressed genes, the 557 housekeeping gene C1orf43 values were subtracted from the target value to receive the ΔCt . 558 To calculate the mean log2 fold changes three biologically independent experiments were 559 used. The log2 fold changes are visualized as single data points and mean. All primers used 560 in this study are listed in Dataset EV6.

561 RNA-sequencing and computational analyses

562 Four different RNA-seq experiments were performed: 1) unaltered Flp-In T-REx 293 wild type 563 (WT) cells and WT cells overexpressing the UPF3A WT construct via the PB-Transposase 564 system. 2) Flp-In T-REx 293 wild type (WT) cells transfected with Luciferase siRNA and the 565 UPF3A KO clones 14 and 20 treated with either Luciferase or UPF3B siRNAs. 3) The control 566 Flp-In T-REx 293 wild type (WT) cells with a Luciferase KD and the UPF3B KO clone 90 567 transfected with Luciferase or UPF3A siRNAs. 4) WT cells transfected with Luciferase siRNA 568 and the two UPF3 double KO cell lines 1 and 2 transfected with either Luciferase or UPF3B 569 siRNAs. RNA was purified using peqGOLD TriFast (VWR Peqlab; for UPF3B KO samples) or 570 the Direct-zol RNA MiniPrep kit including the recommended DNase I treatment (Zymo 571 Research; all other samples) according to manufacturer's instructions. Three biological 572 replicates were analyzed for each sample.

573 The Lexogen SIRV Set3 Spike-In Control Mix (SKU: 051.0x; for UPF3B KO samples) or ERCC 574 RNA Spike-In Mix (for all other samples) that provides a set of external RNA controls was 575 added to the total RNA to enable performance assessment. The Spike-Ins were used for 576 quality control purposes, but not used for the final analysis of DGE, DTU or AS.

577 Using the Illumina TruSeg Stranded Total RNA kit library preparation was accomplished. This 578 includes removing ribosomal RNA via biotinylated target-specific oligos combined with Ribo-579 Zero gold rRNA removal beads from 1 µg total RNA input. Cytoplasmic and mitochondrial 580 rRNA gets depleted by the Ribo-Zero Human/Mouse/Rat kit. After a purification step, the RNA 581 gets cleaved and fragmented. These fragments are then reverse transcribed into first strand 582 cDNA using reverse transcriptase and random primers. In the next step, using DNA 583 Polymerase I and RNase H second strand cDNA synthesis is performed. The resulting cDNA 584 fragments then have the extension of a single'A' base and adapter ligation. To create the final 585 cDNA library the products are purified and enriched with PCR. Next library validation and 586 quantification (Agilent tape station) are performed, followed by pooling of equimolar amounts of library. The pool itself was then quantified using the Peqlab KAPA Library Quantification Kit 587 588 and the Applied Biosystems 7900HT Sequence Detection System and sequenced on an 589 Illumina HiSeq4000 sequencing instrument with an PE75 protocol (UPF3B KO samples) or 590 Illumina NovaSeq6000 sequencing instrument with an PE100 protocol (all other samples).

Reads were aligned against the human genome (version 38, GENCODE release 33 transcript
annotations (Frankish *et al.*, 2019) supplemented with SIRVomeERCCome annotations from

593 Lexogen; obtained from https://www.lexogen.com/sirvs/download/) using the STAR read 594 aligner (version 2.7.3a) (Dobin *et al.*, 2013). Transcript abundance estimates were computed 595 with Salmon (version 1.3.0) (Patro et al., 2017) with a decoy-aware transcriptome. After the 596 import of transcript abundances, differential gene expression analysis was performed with the 597 DESeq2(Love et al., 2014) R package (version 1.28.1) with the significance thresholds 598 |log2FoldChange|> 1 and adjusted p-value (padj) < 0.05. Differential splicing was detected with 599 LeafCutter (version 0.2.9) (Li et al., 2018) with the significance thresholds |deltapsi| > 0.1 and 600 adjusted p-value (p.adjust) < 0.05. Differential transcript usage was computed with 601 IsoformSwitchAnalyzeR (version 1.10.0) and the DEXSeg method (Anders et al., 2012; Ritchie 602 et al., 2015; Robinson & Oshlack, 2010; Soneson et al., 2015; Vitting-Seerup & Sandelin, 2017, 603 2019). Significance thresholds were |d|F| > 0.1 and adjusted p-value (isoform switch q value) 604 < 0.05.

PTC status of transcript isoforms with annotated open reading frame was determined by IsoformSwitchAnalyzeR using the 50 nucleotide (nt) rule of NMD (Huber *et al.*, 2015; Vitting-Seerup *et al.*, 2014; Vitting-Seerup & Sandelin, 2017; Weischenfeldt *et al.*, 2012). Isoforms with no annotated open reading frame in GENCODE were designated "NA" in the PTC analysis.

All scripts and parameters for the RNA-Seq analysis are available at GitHub [https://github.com/boehmv/UPF3]. Overlaps of data sets were represented via nVenn(Perez-Silva *et al.*, 2018) or the ComplexHeatmap package (version 2.6.2)(Gu *et al.*, 2016). Integrative Genomics Viewer (IGV) (version 2.8.12)(Robinson *et al.*, 2011) snapshots were generated from mapped reads (BAM files) converted to binary tiled data (tdf), using Alfred(Rausch *et al.*, 2019) with resolution set to 1 and IGVtools.

616 SILAC and mass spectrometry

HEK293 WT cells and the UPF3 dKO clone 2 expressing either FLAG-tagged GST or UPF2
were labeled by culturing them for at least 5 passages in DMEM for SILAC medium (Thermo
Fisher Scientific) supplemented with 9% FBS (Silantes), 1% Penicillin-Streptomycin and the

labeled amino acids Lysin and Arginine at final concentrations of 0.798 mmol/L and 0.393
mmol/L, respectively. The three conditions were "light" (unlabeled Lys/ Arg), "medium" (Lys4/
Arg6) and "heavy" (Lys8/ Arg10). Unlabeled proline was added in all conditions to prevent
enzymatic Arginine-to-Proline conversion.

624 Experimental setup for SILAC with FLAG-tagged UPF2

625 Expression of FLAG-GST and FLAG-UPF2 was induced for 72 h with 1x cumate. The cells 626 lysed in 250 – 400 µl Buffer E with 1 µg/ml RNase and sonicated using the Bandelin Sonopuls 627 mini20 with 15x 1s pulses at 50% amplitude with a 2.5 mm tip. Protein concentrations were 628 measured using the Bradford assay and protein samples containing 1.6-1.7 mg/ml total protein 629 were diluted. 600 µl of these samples were incubated with 30 µl Anti-FLAG M2 magnetic beads 630 (Sigma) for 2 h on an overhead shaker at 4 °C. The beads were then washed three times for 631 5 min with mild EJC-Buffer before eluting twice with 22 µl of a 200 µg/ml dilution of FLAG-632 peptides (Sigma) in 1x TBS for 10 min at RT and 200 rpm each elution step. Another elution 633 with 1x SDS loading buffer was performed to analyze pull down efficiency via Western blot. 634 The FLAG-peptide eluates were then mixed as followed: 7 µl of both light conditions, 14 µl 635 medium and 14 µl heavy. 1 volume of SP3 (10% SDS in PBS) was added and the samples 636 were reduced with 5 mM DTT and alkylated with 40 mM CAA.

637 Tryptic protein digestion was achieved by following a modified version of the single pot solid 638 phase-enhanced sample preparation (SP3) (Hughes et al., 2014). In brief, paramagnetic Sera-639 Mag speed beads (Thermo Fisher Scientific) were added to the reduced and alkylated protein 640 samples and then mixed 1:1 with 100% acetonitrile (ACN). Protein-beads-complexes form 641 during the 8 min incubation step, followed by capture using an in-house build magnetic rack. 642 After two washing steps with 70% EtOH, the samples were washed once with 100% ACN. 643 Then they were air-dried, resuspended in 5 µl 50 mM Triethylamonium bicarbonate 644 supplemented with trypsin and LysC in an enzyme:substrate ratio of 1:50 and incubated for 16 645 h at 37°C. The next day the beads were again resuspended in 200 µl ACN and after 8 min 646 incubation placed on the magnetic rack. Tryptic peptides were washed with 100% ACN and

647 air-dried before dissolved in 4% DMSO and transfer into 96-well PCR tubes. The last step was 648 the acidification with 1 μ l of 10% formic acid, then the samples were ready for mass spec 649 analysis.

650 Proteomics analysis was performed by the proteomics core facility at CECAD via data-651 dependent acquisition using an Easy nLC1200 ultra high-performance liquid chromatography 652 (UHPLC) system connected via nano electrospray ionization to a Q Exactive Plus instrument 653 (all Thermo Scientific) running in DDA Top10 mode. Based on their hydrophobicity the tryptic 654 peptides were separated using a chromatographic gradient of 60 min with a binary system of 655 buffer A (0.1% formic acid) and buffer B (80% ACN, 0.1% formic acid) with a total flow of 250 656 nl/min. For the separation in-house made analytical columns (length: 50 cm, inner diameter: 657 75 µm) containing 2.7 µm C18 Poroshell EC120 beads (Agilent) that were heated to 50 °C in 658 a column oven (Sonation) were used. Over a time period of 41 min Buffer B was linearly 659 increased from 3% to 27% and then more rapidly up to 50% in 8 min. Finally, buffer B was 660 increased to 95% within 1 min followed by 10 min at 95% to wash the analytical column. Full 661 MS spectra (300-1,750 m/z) were accomplished with a resolution of 70,000, a maximum 662 injection time of 20 ms and an AGC target of 3e6. In each full MS spectrum, the top 10 most 663 abundant ions were selected for HCD fragmentation (NCE:27) with a guadrupole isolation 664 width of 1.8 m/z and 10 s dynamic exclusion. The MS/MS spectra were then measured with a 665 35,000 resolution, an injection time of maximum 110 ms and an AGC target of 5e5.

666 The MS RAW files were then analyzed with MaxQuant suite (version 1.5.3.8) on standard 667 settings with the before mentioned SILAC labels (Cox & Mann, 2008). By matching against the 668 human UniProt database the peptides were then identified using the Andromeda scoring 669 algorithm (Cox et al., 2011). Carbamidomethylation of cysteine was defined as a fixed 670 modification, while methionine oxidation and N-terminal acetylation were variable 671 modifications. The digestion protein was Trypsin/P. A false discovery rate (FDR) < 0.01 was 672 used to identify peptide-spectrum matches and to quantify the proteins. Data processing and 673 statistical analysis was performed in the Perseus software (version 1.6.1.1) (Tyanova et al.,

674 2016). Using the One-sample t-test the significantly changed proteins were identified (H0 = 0,

fudge factor S0 = 0.2). Visualization was performed with RStudio (version 1.2.5033).

676 Label-free quantitative mass spectrometry

677 Twenty-four hours before expression of the FLAG-tagged constructs, the HEK 293 WT cells 678 were treated with Luciferase siRNA and the UPF3B KO clone 90 and UPF3 dKO clone 1 cells 679 were treated with siRNAs targeting residual UPF3B. The expression of either FLAG-GST or 680 FLAG-UPF2 in WT cells and FLAG-UPF2 in the clones 90 and 1 was induced for 48 h with 1x. 681 Lysis and sample preparation were performed as described above. MS analysis was 682 performed as described above with a slightly adjusted gradient as followed: 3 - 30% B in 41 683 min, 30 – 50% B in 8 min, 50-95% B in 1 min, followed by 10 min washing at 95%. LFQ values 684 were calculated using the MaxLFQ algorithm (Cox et al., 2014) in MaxQuant. Significantly 685 changed proteins were identified by two-sample *t*-testing (fudge factor S0 = 0.2).

686 Northern Blotting

687 The cells were harvested in RNAsolv reagent and total RNA extraction was performed as 688 described above. 3.0 µg total RNA were resolved on a 1% agarose/0.4 M formaldehyde gel 689 using the tricine/triethanolamine buffer system (Mansour & Pestov, 2013). Next a transfer on 690 a nylon membrane (Roth) in 10x SSC followed. The blot was incubated overnight at 65°C in 691 Church buffer containing [α -32P]-GTP body-labeled RNA-probes for mRNA reporter detection 692 (Voigt et al., 2019). Ethidium bromide stained 28S and 18S rRNA served as loading controls. 693 RNA signal detected with the Typhoon FLA 7000 (GE Healthcare) was guantified in a semi-694 automated manner using the ImageQuant TL 1D software with a rolling-ball background 695 correction. EtBr-stained rRNA bands were quantified with the Image Lab 6.0.1 software (Bio-696 Rad). Signal intensities were normalized to the internal control (rRNA) before calculation of 697 mean values. The control condition was set to unity (TPI WT for reporter assays), quantification 698 results are shown as data points and mean.

699 **Protein conservation**

- 700 UPF3A (UniProt ID: Q9H1J1-1) and UPF3B (UniProt ID: Q9BZI7-2) protein sequences were
- aligned using Clustal Omega (<u>https://www.ebi.ac.uk/Tools/msa/clustalo/</u>) (Goujon et al., 2010;
- Sievers *et al.*, 2011), viewed using Jalview (Waterhouse *et al.*, 2009), the conservation score
- 703 extracted and used for visualization.

704 Data Presentation

Quantifications and calculations for other experiments were performed - if not indicated otherwise - with Microsoft Excel (version 1808) or R (version 4.0.4) and all plots were generated using IGV (version 2.8.12), GraphPad Prism 5, ggplot2 (version 3.3.3) or ComplexHeatmap (version 2.6.2) (Gu *et al.*, 2016).

709 Data Availability

- The datasets and computer code produced in this study are available in the followingdatabases:
- RNA-Seq data for UPF3B KO samples: ArrayExpress E-MTAB-10711
- 713 (https://www.ebi.ac.uk/arrayexpress/experiments/E-MTAB-10711)
- RNA-Seq data for UPF3 dKO samples: ArrayExpress E-MTAB-10716
- 715 (https://www.ebi.ac.uk/arrayexpress/experiments/E-MTAB-10716)
- RNA-Seq data for UPF3A KO/OE samples: ArrayExpress E-MTAB-10718
- 717 (<u>https://www.ebi.ac.uk/arrayexpress/experiments/E-MTAB-10718</u>)
- Mass spectrometry proteomics data: PRIDE PXD027120
- 719 (https://www.ebi.ac.uk/pride/archive/projects/PXD027120)
- Codes used in this study: GitHub (<u>https://github.com/boehmv/UPF3</u>)
- All relevant data supporting the key findings of this study are available within the article and its
- Expanded View files or from the corresponding author upon reasonable request.

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748 Conflict of interest

749 None.

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