The RNA-binding ubiquitin ligase MKRN1 functions in ribosomeassociated quality control of poly(A) translation

Andrea Hildebrandt¹, Mirko Brüggemann², Susan Boerner², Cornelia Rücklé², Jan Bernhard Heidelberger¹, Annabelle Dold¹, Anke Busch¹, Heike Hänel¹, Andrea Voigt¹, Stefanie Ebersberger¹, Ingo Ebersberger^{3,4}, Jean-Yves Roignant¹, Kathi Zarnack^{2§}, Julian König^{1§}, Petra Beli^{1§}

¹ Institute of Molecular Biology (IMB), Ackermannweg 4, 55128 Mainz, Germany

² Buchmann Institute of Molecular Life Sciences (BMLS), Goethe University, Maxvon-Laue-Str. 15, 60438 Frankfurt am Main, Germany

 ³ Department for Applied Bioinformatics, Institute of Cell Biology and Neuroscience, Goethe University, Max-von-Laue-Str. 13, 60438 Frankfurt am Main, Germany
 ⁴ Senckenberg Biodiversity and Climate Research Centre (BiK-F), Georg-Voigt-Straße 14-16, 60325 Frankfurt am Main, Germany

§ Corresponding authors:

KZ (kathi.zarnack@bmls.de), JK (j.koenig@imb-mainz.de) and PB (p.beli@imbmainz.de)

1 Abstract

2 Cells have evolved quality control mechanisms to ensure protein homeostasis by 3 detecting and degrading aberrant mRNAs and proteins. A common source of aberrant 4 mRNAs is premature polyadenylation, which can result in non-functional protein 5 products. Translating ribosomes that encounter poly(A) sequences are terminally 6 stalled, followed by ribosome recycling and decay of the truncated nascent polypeptide 7 via the ribosome-associated guality control (RQC). Here, we demonstrate that the 8 conserved RNA-binding E3 ubiguitin ligase Makorin Ring Finger Protein 1 (MKRN1) 9 promotes ribosome stalling at poly(A) sequences during RQC. We show that MKRN1 10 interacts with the cytoplasmic poly(A)-binding protein (PABP) and is positioned upstream of poly(A) tails in mRNAs. Ubiquitin remnant profiling uncovers PABP and 11 12 ribosomal protein RPS10, as well as additional translational regulators as main 13 ubiguitylation substrates of MKRN1. We propose that MKRN1 serves as a first line of 14 poly(A) recognition at the mRNA level to prevent production of erroneous proteins, thus 15 maintaining proteome integrity.

16 Keywords

- 17 MKRN1, ubiquitylation, RNA binding, ribosome-associated quality control, RQC,
- 18 poly(A), iCLIP, ubiquitin remnant profiling, translation

19 Abbreviations

20 Adenosine (A), Makorin Ring Finger Protein 1 (MKRN1), ribosome-associated guality control (RQC), poly(A)-binding protein (PABP), mass spectrometry (MS), affinity 21 22 purification (AP), stable isotope labelling with amino acids in cell culture (SIALC), false discovery rate (FDR), Gene Ontology (GO), Biological Process (BP), Molecular 23 24 Function (MF), mRNA ribonucleoprotein particle (mRNP), PABP-interacting motif (PAM2), MKRN1 variant with point mutations in the PAM2 motif (GFP-MKRN1^{PAM2mut}), 25 MKRN1 variant with point mutation in RING domain (GFP-MKRN1^{RINGmut}), individual-26 nucleotide resolution UV crosslinking and immunoprecipitation (iCLIP), signal-over-27 background (SOB), nucleotides (nt), 4-thiouridine (4SU), Pearson correlation 28 coefficients (r), A-rich stretches (A-stretches), lysine (K), knock down (KD), RNA 29 recognition motif (RRM), Polyethylenimine (PEI), modified RIPA (mRIPA), N-30 31 ethylmaleimide (NEM), dithiothreitol (DTT), chloroacetamide (CAA), higher-energy 32 collisional dissociation (HCD), glycine-glycine (GlyGly), strong-cation exchange 33 chromatography (SCX)

34 Introduction

During gene expression, quality control pathways monitor each step to detect aberrant mRNAs and proteins. These mechanisms ensure protein homeostasis and are essential to prevent neurodegenerative diseases (Chu et al. 2009). A common source of aberrant mRNAs is premature polyadenylation, often in combination with missplicing, which results in truncated non-functional protein products (Kaida et al. 2010). Therefore, mechanisms are in place that recognise such homopolymeric adenosine (poly(A)) sequences and abrogate their translation (Bengtson and Joazeiro 2010).

42 In eukaryotes, ribosomes that terminally stall for diverse reasons during translation are 43 detected by the ribosome-associated quality control (RQC) (reviewed in Brandman and 44 Hegde 2016; Joazeiro 2017). Upon splitting of the 60S and 40S ribosomal subunits, the RQC complex assembles on the stalled 60S subunit to initiate the release and 45 rapid degradation of the truncated tRNA-bound polypeptide. The E3 ubiquitin ligase 46 Listerin (LTN1) modifies the truncated polypeptide with K48-linked ubiquitin chains to 47 target it for degradation in a p97-dependent manner through the proteasome 48 49 (Bengtson and Joazeiro 2010; Brandman et al. 2012; Verma et al. 2013). Whereas 50 peptide release and ribosome recycling by the RQC complex are relatively well 51 understood, less is known about the mechanisms that promote poly(A) recognition and 52 initial ribosome stalling.

53 Several recent studies demonstrated a role for the RNA-binding E3 ubiquitin ligase 54 ZNF598 in initiating RQC for prematurely polyadenylated mRNAs (Garzia et al. 2017; 55 Juszkiewicz and Hegde 2017; Sundaramoorthy et al. 2017). It was suggested that 56 ZNF598 senses the translation of poly(A) segments through binding the cognate lysine 57 tRNAs (Garzia et al. 2017). In addition, ZNF598 recognises the collided di-ribosome 58 structure that arises when a trailing ribosome encounters a slower leading ribosome

(Juszkiewicz et al. 2018). This is followed by site-specific, regulatory ubiquitylation of
the 40S ribosomal proteins RPS10 and RPS20 by ZNF598. In addition to ZNF598, the
40S ribosomal subunit-associated protein RACK1 was shown to regulate ubiquitylation
of RPS2 and RPS3 upstream of ribosomal rescue (Sundaramoorthy et al. 2017).

63 Makorin Ring Finger Protein 1 (MKRN1) belongs to a family of evolutionary conserved 64 RNA-binding E3 ubiguitin ligases. Up to four paralogs exist in vertebrates (MKRN1-4), which combine a RING domain with one or more CCCH zinc finger domains (Gray et 65 66 al. 2000; Böhne et al. 2010) (Supplemental Fig. 1A). MKRN1 has been implicated in 67 the regulation of telomere length, RNA polymerase II transcription and the turnover of tumour suppressor protein p53 and cell cycle regulator p21 (Kim et al. 2005; 68 69 Omwancha et al. 2006; Lee et al. 2009; Salvatico et al. 2010), but its RNA-related 70 functions remain poorly understood. A study in mouse embryonic stem cells (mESC) 71 reported its interaction with hundreds of mRNAs as well as multiple RNA-binding 72 proteins (RBPs), including the cytoplasmic poly(A)-binding protein (PABP) PABPC1, 73 IGF2BP1 and ELAVL1 (Cassar et al. 2015). The interaction with PABP was further 74 corroborated in human HEK293 cells (Miroci et al. 2012). The same study 75 demonstrated that a shortened isoform of MKRN1 controls local translation via its 76 PABP-interacting motif 2 (PAM2 motif) in rat neurons (Miroci et al. 2012). In line with 77 a role in translation, MKRN1 was found in association with ribosomes, from which it 78 could be released together with PABP and other proteins by RNase digestion (Simsek 79 et al. 2017). Nevertheless, the RNA binding specificity and functional role of MKRN1 80 in human cells remained largely elusive.

Here, we introduce MKRN1 as a novel factor in RQC. We propose that MKRN1 is recruited to A-rich sequences in mRNAs in a PABP-dependent manner, where it acts as a first line of defence against poly(A) translation. MKRN1 depletion abrogates

ribosome stalling in reporter assays, accompanied by reduced ubiquitylation of RQC-

related proteins. We therefore hypothesise that MKRN1 allows recognition of poly(A)

86 sequences prior to their translation.

87 **Results**

88 MKRN1 interacts with PABPC1 and other RBPs

89 In order to learn about potential functions, we first characterised the protein interaction profile of MKRN1 in HEK293T cells. To this end, we used affinity purification (AP) 90 91 coupled to stable isotope labelling with amino acids in cell culture (SILAC)-based quantitative mass spectrometry (MS) using GFP-MKRN1^{wt} or GFP as a bait. We 92 identified 53 proteins that were significantly enriched in GFP-MKRN1^{wt} compared to 93 94 the control APs (false discovery rate [FDR] < 5%, combined ratios of three independent 95 experiments). In line with previous reports (Miroci et al. 2012; Cassar et al. 2015; Hildebrandt et al. 2017), we found the cytoplasmic poly(A)-binding proteins (PABP) 96 97 PABPC1 and PABPC4 among the highly enriched MKRN1 interactors (z-score > 4. corrected *P* values = 7.18e-10 and 6.16e-16, respectively) (Fig. 1A, Supplemental 98 99 Fig. S2A, and Supplemental Table S1). Moreover, we detected 14 ribosomal proteins 100 as well as four proteins that were previously shown to co-purify with ribosomes (Simsek 101 et al. 2017), including IGF2BP1, LARP1, UPF1, and ELAVL1 (Fig. 1A). Consistently, 102 "translation" was among the significantly enriched Gene Ontology (GO) terms for the 103 MKRN1 interaction partners (Biological Process [BP], Supplemental Fig. S2B). 104 Almost all interactors were previously found in association with polyadenylated 105 transcripts (50 out of 53 proteins have been annotated with the GO term "poly(A) RNA binding", Molecular Function [MF], **Supplemental Fig. S2B**). We confirmed the MS 106 107 results in reciprocal AP experiments with GFP-tagged PABPC1, ELAVL1, and

108 IGF2BP1 as baits followed by Western blot for endogenous MKRN1 (Supplemental 109 Fig. S2C). All detected interactions persisted in the presence of RNases (RNase A 110 and T1), demonstrating that MKRN1 interacts with these proteins in an RNA-111 independent manner (Supplemental Fig. S2C). Together, these observations suggest 112 that MKRN1 is part of a larger mRNA ribonucleoprotein particle (mRNP) together with 113 PABP and other RBPs. This is further supported by a parallel study on the Mkrn1 114 ortholog in Drosophila melanogaster, which consistently identified pAbp, Larp, Upf1 115 and Imp (IGF2BP in mammals) as interaction partners (Dold et al, parallel submission; 116 preprint available at bioRxiv, doi: 10.1101/501643).

117 Many proteins interact with PABP via a PABP-interacting motif (PAM2) motif, which 118 specifically binds to the MLLE domain present almost exclusively in PABP (Deo et al. 119 2001; Kozlov et al. 2010). Accordingly, a previous study demonstrated that MKRN1 120 associates with PABP via a PAM2 motif at amino acid positions 161-193 (Miroci et al. 121 2012). In support of a putative functional relevance, a phylogenetic analysis illustrated 122 that the presence and positioning of the PAM2 motif are preserved in MKRN1 orthologs 123 across metazoans (Supplemental Fig. S1A,B). AP of a MKRN1 variant with point 124 mutations in the PAM2 motif (GFP-MKRN1^{PAM2mut}) (Pohlmann et al. 2015) no longer 125 recovered PABPC1 and PABPC4 (Fig. 1B-D and Supplemental Table S1). For comparison, we also tested a previously described point mutation in the RING domain 126 127 that abolishes the E3 ubiguitin ligase function (ligase-dead, GFP-MKRN1^{RINGmut}) (Kim 128 et al. 2005). This mutation did not impair the interaction of MKRN1 with PABPC1, but 129 led to a slight increase, possibly due to stabilisation of MKRN1 and/or PABPC1 (Fig. 130 **1C, Supplemental Fig. S7, and Supplemental Table S1**). Surprisingly, 131 MKRN1^{PAM2mut} lost interaction not only with PABPC1 and PABPC4, but also with several other identified proteins (**Fig. 1D**), suggesting that MKRN1^{PAM2mut} no longer 132

133 resided within the mRNPs. These results confirmed that MKRN1 interacts with PABP

134 proteins, and suggested that this association is required for mRNP formation.

135 MKRN1 binds to poly(A) tails and at internal A-stretches

In order to characterise the RNA-binding behaviour of human MKRN1 in vivo, we 136 137 performed individual-nucleotide resolution UV crosslinking and immunoprecipitation (iCLIP) (König et al. 2010) in combination with 4-thiouridine (4SU) labelling to enhance 138 139 UV crosslinking (Hafner et al. 2010). In three replicate experiments with GFP-tagged 140 MKRN1 (GFP-MKRN1^{wt}) expressed in HEK293T cells, we identified more than 4,6 141 million unique crosslink events, cumulating into 7,331 MKRN1 binding sites (see 142 Materials and methods; Supplemental Table S2). These were further ranked 143 according to the strength of MKRN1 binding, which was estimated from the enrichment 144 of crosslink events within a binding site relative to its local surrounding, which served as a proxy for transcript abundance ("signal-over-background", SOB; see Materials 145 and Methods) (Sutandy et al. 2018). SOB values were highly reproducible between 146 replicates (Pearson correlation coefficients r > 0.72, **Supplemental Fig. S3**). 147

148 Across the transcriptome, MKRN1 almost exclusively bound to protein-coding mRNAs 149 with a strong tendency to locate in 3' UTRs (Fig. 2A,E). Binding sites generally 150 harboured uridine-rich tetramers (Supplemental Fig. S4A), likely reflecting 4SU-151 based UV crosslinking (Hafner et al. 2010). Strikingly, the top 20% MKRN1 binding 152 sites were massively enriched in AAAA tetramers (A, adenosine) within 5-50 nucleotides (nt) downstream of the binding sites (Fig. 2B and Supplemental Fig. 153 154 **S4A**). These were situated within A-rich stretches (A-stretches), which ranged from 8-155 30 nt in length (Supplemental Fig. S4B; see Materials and methods). Within 3' UTRs, 156 30% (1,848 out of 6,165) of MKRN1 binding sites resided immediately upstream of an 157 A-stretch (Fig. 2C,E) and longer A-stretches associated with stronger MKRN1 binding (Supplemental Fig. S4C,D). Intriguingly, we detected a requirement for a continuous run of at least 8 A's to confer strong MKRN1 binding (Fig. 2D), which precisely matched the RNA footprint of PABP (Webster et al. 2018). Since PABP was previously reported to also bind within 3' UTRs (Bag 2001; Lyabin et al. 2011; Kini et al. 2016), these observations indicated that MKRN1 binds together with PABP to mRNAs.

163 Prompted by this notion, we analysed the unusually high fraction of unmapped iCLIP 164 reads in the MKRN1 dataset (Supplemental Table S2). In accordance with binding of MKRN1 immediately upstream of poly(A) tails, more than 13% of the unmapped reads 165 166 displayed an increased A-content (Fig. 3B), compared to only 2% for an unrelated 167 control RBP (Braun et al. 2018). In addition, the mapped GFP-MKRN1^{wt} crosslink 168 events were enriched upstream of annotated polyadenylation sites, as exemplified in 169 the SRSF4 gene (Fig. 2E and Fig. 3A,C). Together, these results support the notion 170 that MKRN1 binds upstream of poly(A) tails, possibly in conjunction with PABP. In order to test whether PABP is required for MKRN1 binding, we performed UV 171 crosslinking experiments with GFP-MKRN1^{PAM2mut}, which no longer interacts with 172 173 PABP (Fig. 1C,D). Strikingly, RNA binding of this mutant was globally reduced 174 compared to GFP-MKRN1^{wt} (Fig. 3D and Supplemental Fig. S5), indicating that 175 PABP might recruit MKRN1 to RNA. In summary, these results strongly imply that 176 MKRN1 binds upstream of poly(A) tails, which could be implemented via its interaction 177 with PABP. In a concordant scenario, it was found that *Drosophila* Mkrn1 bound before 178 an extended A-stretch in the 3' UTR of oskar mRNA, and that this binding was 179 significantly reduced upon depletion of pAbp (Dold et al., parallel submission; preprint 180 available at bioRxiv, doi: 10.1101/501643).

181 MKRN1 promotes ribosome stalling at poly(A) sequences

182 As outlined above, our iCLIP data evidenced that MKRN1 marks the beginning of 183 poly(A) tails. Hence, it is conceivable that MKRN1 will also bind upstream of premature 184 polyadenylation events within open reading frames. Based on MKRN1's binding 185 pattern, its interaction partners and its previously reported association with ribosomes 186 (Simsek et al. 2017), we hypothesised that MKRN1 may be involved in the clearance of such transcripts by ribosome-associated quality control (RQC). In this process, 187 188 ribosomes that translate into a poly(A) sequence, for instance upon stop codon 189 readthrough and premature polyadenylation, are stalled and eventually recycled 190 (Brandman and Hegde 2016; Joazeiro 2017). To test this hypothesis, we employed a 191 recently introduced flow cytometry-based assay that monitors ribosome stalling in a 192 dual fluorescence reporter (Juszkiewicz and Hegde 2017) (Fig. 4A).

193 As reported previously, inserting a K(AAA)₂₀ linker (encoding for 20 lysine residues) 194 into the reporter resulted in predominant ribosome stalling compared to the starting 195 vector (K₀, Fig. 4B and Supplemental Fig. 6A). Importantly, *MKRN1* depletion with 196 two independent siRNA sequences led to a reproducible recovery of RFP expression 197 downstream of K(AAA)₂₀, demonstrating that many ribosomes failed to stall at 198 K(AAA)₂₀ (*MKRN1* KD1 and KD2; Fig. 4C and Supplemental Fig. S6A,B). *MKRN1* KD2 seemed slightly more effective, possibly because this siRNA simultaneously 199 200 decreased the transcript levels of the close paralogue MKRN2 (Supplemental Fig. 201 **S6C**). Notably, *MKRN1* KD2 impaired ribosome stalling to a similar level as KD of 202 ZNF598, the E3 ubiquitin ligase that was recently reported to function in RQC (Garzia 203 et al. 2017; Juszkiewicz and Hegde 2017; Sundaramoorthy et al. 2017). Moreover, 204 simultaneous depletion of *MKRN1* and *ZNF598* was not additive, indicating that both 205 proteins are necessary for function (Fig. 4C and Supplemental Fig. S6A). In addition, 206 we noted a certain level of cross-regulation, such that ZNF598 expression was

decreased in *MKRN1* KD1 (but not in *MKRN1* KD2), whereas *ZNF598* overexpression
reduced *MKRN1* expression (Supplemental Fig. S6D,E). Taken together, we
conclude that MKRN1 contributes to efficient ribosome stalling in RQC.

210 MKRN1 mediates the ubiquitylation of ribosome-associated proteins

211 RQC builds on a series of ubiquitylation events by multiple E3 ubiquitin ligases, 212 including Listerin and ZNF598 (Brandman and Hegde 2016). In order to identify 213 putative ubiquitylation substrates of MKRN1, we first determined the protein interactome of the ligase-deficient mutant GFP-MKRN1^{RINGmut}. In three replicate 214 215 experiments, we quantified 1,097 protein groups present in at least two out of three 216 replicates (Supplemental Table S1), revealing 137 proteins that were significantly 217 enriched compared to GFP-MKRN1^{wt} (Supplemental Fig. S7). Intriguingly, these 218 included RPS10, a ribosomal protein that was previously reported to be modified by 219 ZNF598 during RQC (Garzia et al. 2017; Juszkiewicz and Hegde 2017; 220 Sundaramoorthy et al. 2017).

221 In order to directly test for ubiquitylation of putative substrates of MKRN1, we 222 performed ubiquitin remnant profiling to compare the relative abundance of di-glycine-223 modified lysines in wild type and MKRN1 KD cells. We quantified 2,324 ubiquitylation 224 sites (in 1,264 proteins) that were detected in all four replicate experiments 225 (Supplemental Table S3). Notably, *MKRN1* depletion led to a significantly decreased 226 abundance of 29 ubiguitylation sites on 21 proteins (FDR < 10%, Fig. 5A). The majority 227 of the ubiquitylation targets assembled into a coherent cluster of translational 228 regulators based on previously reported protein-protein interactions and functional 229 annotations (Fig. 5B,C and Supplemental Fig. 8A). Among these proteins, we had 230 already detected PABPC1/4, IGF2BP1, ELAVL1, MOV10, LARP1, and RPS10 as significant interactors of GFP-MKRN1^{wt} and/or GFP-MKRN1^{RINGmut} (Fig. 1A, Fig. 5F 231

232 and **Supplemental Fig. S7**). Importantly, we detected a significant decrease in 233 ubiguitylation at lysine 107 of RPS10 (K107; Fig. 5D). In order to distinguish differential 234 ubiguitylation from protein level changes, we also measured the total protein levels in 235 *MKRN1* KD cells and did not observe changes in RPS10, PABPC1/4, IGF2BP1/2/3, 236 ELAVL1, and MOV10 protein levels (Supplemental Fig. 8B and Supplemental Table 237 **S4**). Taken together, we conclude that MKRN1 mediates ubiguitylation of the ribosomal 238 protein RPS10 and several translational regulators during ribosome-associated quality 239 control.

240 Discussion

Ribosome-associated quality control is essential to recognise and clear terminally stalled ribosomes. Here, we uncover MKRN1 as a novel factor in RQC. Our data indicate that MKRN1 is positioned upstream of poly(A) sequences through direct interaction with PABP, thereby marking the beginning of poly(A) tails. We propose that in case of premature polyadenylation, MKRN1 stalls the translating ribosome and initiates RQC by ubiquitylating ribosomal protein RPS10, PABP and other translational regulators (**Fig. 6**).

248 **PABP recruits MKRN1 upstream of A-stretches and poly(A) tails**

Central to our model is the specific RNA-binding behaviour of MKRN1, which is recruited to mRNA by PABP to mark the beginning of poly(A) tails. This builds on the following observations: (i) We and others show that MKRN1 and PABP interact via the PAM2 motif (Miroci et al. 2012). (ii) MKRN1 binding to RNA is strongly reduced when interaction with PABP is abolished. (iii) The association of strong MKRN1 binding with continuous A-runs of \geq 8 A's mirrors the footprint of one RNA recognition motif (RRM) domain of PABP, indicating that the binding of one RRM to poly(A) is sufficient for

256 MKRN1 recruitment (Webster et al. 2018). On such short A-stretches, MKRN1 might 257 stabilise PABP binding, while on longer A-stretches, PABP might be the major driving 258 force to recruit MKRN1. This interaction might also anchor the first PABP at the 259 beginning of the poly(A) tail. One possible function could be the stabilisation of PABP 260 on short poly(A) tails to promote efficient translation (Lima et al. 2017). In yeast, where 261 a MKRN1 ortholog is missing (see below), this anchoring is thought to be achieved by Pab1p itself via its fourth RRM domain (Webster et al. 2018). Of note, a parallel study 262 263 with the Mkrn1 ortholog from *D. melanogaster* demonstrates binding of a Mkrn1/pAbp 264 complex at an A-stretch in the 3' UTR of oskar mRNA, which is involved in translational 265 control and required for oogenesis (Dold et al., parallel submission; preprint available at bioRxiv, doi: 10.1101/501643). 266

267 MKRN1 ubiquitylates RPS10 and translational regulators to stall ribosomes

268 Our data suggest that ribosomes encountering the MKRN1-PABP complex are stalled, 269 possibly via ubiquitylation of RPS10 and other MKRN1 interactors. Concordantly, 270 ZNF598, a factor that was recently shown to function in RQC, was also found to 271 mediate ubiquitylation of RPS10 (Juszkiewicz et al. 2018). In conjunction with its 272 unique RNA-binding behaviour, we therefore hypothesise that MKRN1 acts as a first 273 line of defence against poly(A) translation. We propose that MKRN1 is recruited by 274 PABP to the beginning of poly(A) tails, including premature polyadenylation events 275 within open reading frames, where it represents a physical "roadblock" to the 276 translating ribosome. Upon contact with the translating ribosome, MKRN1 ubiguitylates 277 K107 on RPS10, thereby stalling the ribosome before it translates the poly(A) tail. 278 Subsequently, the trailing ribosomes collide with the initially stalled ribosome. ZNF598 279 recognises the collision interface and ubiquitylates the collided ribosomes (Simms et 280 al. 2017; Juszkiewicz et al. 2018). In summary, we suggest that a sequence of MKRN1-

281 mediated and ZNF598-mediated ubiquitylation events on ribosomal proteins and 282 possibly other factors, including PABPC1, triggers ribosome-associated quality control.

283 Differences between human and yeast RQC explain the requirement for MKRN1

284 Many known components of the RQC machinery, such as Listerin (Ltn1p in yeast) and 285 ZNF598 (Hel2p in yeast), are identical from yeast to human, however the molecular signals that are recognised differ partially. In yeast, RQC can be triggered by an excess 286 287 of positively charged amino acids (lysine and arginine), which are sensed while they 288 pass through the ribosomal exit tunnel (Lu and Deutsch 2008; Letzring et al. 2013). In 289 contrast, in human, sensing the aberrant mRNAs does not occur via the encoded 290 amino acids but at the level of the mRNA sequence and corresponding tRNAs, such 291 that only poly(A) effectively results in ribosome stalling (Arthur et al. 2015; Garzia et al. 2017; Juszkiewicz and Hegde 2017). We propose that MKRN1 acts as direct reader 292 293 of poly(A) sequences based on its interaction with PABP. Consistent with this 294 conceptual difference, there is no functionally equivalent ortholog of MKRN1 in yeast 295 (Yth1p and Lee1p are similar, but lack RING domain and PAM2 motif; Supplemental 296 Fig. 1C). Why yeast and human employ partially different mechanisms to detect 297 poly(A) translation is currently unclear, but it has been suggested that spurious 298 translation of poly-lysine stretches from long human poly(A) tails might target the 299 aberrant proteins to the nucleus (Juszkiewicz and Hegde 2017). Loss of mRNA 300 surveillance and RQC deficiency can lead to protein aggregation and culminate in 301 proteotoxic stress, which in turn is lined to neurological disorders such as amyotrophic 302 lateral sclerosis (Choe et al. 2016; Jamar et al. 2018). Hence, recognition of poly(A) 303 sequences prior to their translation might be particularly beneficial in humans.

bioRxiv preprint doi: https://doi.org/10.1101/516005; this version posted January 25, 2019. The copyright holder for this preprint (which was not certified by peer review) is the author/funder. All rights reserved. No reuse allowed without permission.

304 Materials and methods

305 Cell culture

HEK293T cells were obtained from DSMZ and cultured in DMEM (Life Technologies) 306 with 10% fetal bovine serum (Life Technologies), 1% penicillin/streptomycin (Life 307 308 Technologies), and 1% L-glutamine (Life Technologies). All cells were maintained at 309 37°C in a humidified incubator containing 5% CO₂ and routinely tested for mycoplasma 310 infection. For SILAC labelling, cells were maintained in media containing either L-311 arginine and L-lysine (light SILAC label), L-arginine (¹³C₆) and L-lysine (²H₄) (medium SILAC label), or L-arginine $({}^{13}C_{6} - {}^{15}N_{4})$ and L-lysine $({}^{13}C_{6} - {}^{15}N_{2})$ (heavy SILAC label) 312 313 (Cambridge Isotope Laboratories).

314 Vectors

315 The following vectors, suitable for Gateway Cloning, were obtained either from the IMB 316 Core Facility ORFeome Collection (Collaboration 2016) or from the Harvard PlasmID 317 Repositorv (https://plasmid.med.harvard.edu/PLASMID/): pENTR221-MKRN1. pENTR221-PABPC1, pENTR223.1-IGF2BP1, pENTR221-ELAVL1, pCMV-SPORT-318 319 ZNF598. Coding sequences from the entry vectors were cloned into the mammalian 320 expression vectors pMX-DEST53-IP-GFP by LR Gateway cloning according to the 321 manufacturer's recommendations (Gateway LR Clonase II Enzyme mix; Life 322 Technologies). Dual fluorescence reporter plasmids (pmGFP-P2A-K₀-P2A-RFP, pmGFP-P2A-(KAAA)12-P2A-RFP, pmGFP-P2A-(KAAA)20-P2A-RFP, and pmGFP-P2A-323 (R^{CGA})₁₀-P2A-RFP) were generously provided by Ramanujan S. Hegde (MRC 324 Laboratory of Molecular Biology, Cambridge, UK) (Juszkiewicz and Hegde 2017). 325

326 Cloning

All MKRN1 mutant plasmids were generated with the Q5 Site-Directed Mutagenesis
 Kit (NEB) according to the manufacturer's recommendations. In order to disrupt

MKRN1's interaction with PABP (MKRN1^{PAM2mut}), three point mutations were introduced into the PAM2 motif (A169S, F172A, P174A; **Fig. 1B**) as previously described (Pohlmann et al. 2015). In MKRN1^{RINGmut}, a previously described mutation in the RING domain (H307E) was introduced to abolish E3 ubiquitin ligase function (Kim et al. 2005). All primers used for introducing mutations into MKRN1 are listed in **Supplemental Table S5**.

335 Transfections

336 Overexpression of vectors was performed using Polyethylenimine MAX 4000 337 (Polysciences, 24885-2) with a DNA:PEI ratio of 1:10. Knockdowns were performed 338 with siRNAs (**Supplemental Table S6**) using Lipofectamine RNAiMAX (Life 339 Technologies) according to the manufacturer's recommendations.

340 Affinity purification (AP) for Western blot analyses

341 GFP-based affinity purifications (APs) were performed as described before (Hildebrandt et al. 2017). In brief, HEK293T cells transiently expressing GFP (empty 342 343 vector) or a GFP-tagged target protein were used. The cells were lysed in modified RIPA (mRIPA) buffer supplemented with protease inhibitors (protease inhibitor 344 345 cocktail, Sigma), 1 mM sodium orthovanadate, 5 mM β-glycerophosphate, 5 mM 346 sodium fluoride, and 10 mM N-ethylmaleimide (NEM) (all from Sigma). Protein 347 concentrations were determined using the Pierce BCA Protein Assay Kit (Thermo 348 Fisher). GFP-trap agarose beads (Chromotek) were incubated with the cleared lysate 349 for 1 h at 4°C. After five washes with mRIPA buffer, the beads were resuspended in 350 LDS sample buffer (Life Technologies) and heated to 70°C for 10 min. For RNase 351 digests, the enriched proteins were incubated with 0.5 U/µI RNase A (Qiagen) and 352 20 U/µl RNase T1 (Thermo Fisher Scientific) for 30 min at 4°C after the first two washes 353 in mRIPA buffer.

354 Sample preparation for the protein interactome analysis

355 GFP-based APs were performed as described before (Hildebrandt et al. 2017). In brief, 356 HEK293T cells transiently expressing GFP (empty vector) were cultured in light SILAC 357 medium, while cells expressing N-terminally GFP-tagged MKRN1 wt or mutants were 358 cultured in medium or heavy SILAC medium. The cells were lysed as described above. 359 After washing in mRIPA buffer, GFP-trap agarose beads were incubated with the 360 cleared lysate for 1 h at 4°C. All AP samples were washed four times with mRIPA buffer, combined and washed again in mRIPA buffer. The beads were heated in LDS 361 362 sample buffer, supplemented with 1 mM dithiothreitol (DTT; Sigma, D5545) for 10 min 363 at 70°C and alkylated using 5.5 mM 2-chloroacetamide (CAA; Sigma, C0267) for 30 364 min at RT in the dark (Nielsen et al. 2008).

365 **Sample preparation for the proteome analysis**

MKRN1 KD using siRNA2 was performed in heavy labelled SILAC cells and control KD was performed in light labelled SILAC cells in two replicates. For the third replicate, a label swop was performed, knocking down *MKRN1* (siRNA2) in light labelled SILAC cells and control in heavy labelled SILAC cells. For proteome analysis, cells were lysed as described above. Subsequently, 25 μg protein from each SILAC condition (50 μg in total) were pooled and processed as described below.

372 Sample preparation for mass spectrometry

The enriched proteins were resolved by SDS-PAGE on a NuPAGE 4-12% Bis-Tris protein gel (Thermo Fisher Scientific) and stained using the Colloidal Blue Staining Kit (Life Technologies). Proteins were in-gel digested using trypsin, before peptides were extracted from the gel. To concentrate, clear and acidify the peptides, they were bound to C18 StageTips as described previously (Rappsilber et al. 2007).

378 Mass spectrometry data acquisition

379 Peptide fractions were analysed on a guadrupole Orbitrap mass spectrometer (Thermo Q Exactive Plus, Thermo Scientific) coupled to an uHPLC system (EASY-nLC 1000, 380 381 Thermo Scientific) (Michalski et al. 2011). Peptide samples were separated on a C18 382 reversed phase column (length: 20 cm, inner diameter: 75 µm, bead size: 1.9 µm) and 383 eluted in a linear gradient from 8 to 40% acetonitrile containing 0.1% formic acid in 105 384 min for the interactome analyses, in 175 min for the proteome analyses, or in 125 min for the ubiguitylome analyses. The mass spectrometer was operated in data-385 386 dependent positive mode, automatically switching between MS and MS² acquisition. 387 The full scan MS spectra (m/z 300–1650) were acquired in the Orbitrap. Sequential 388 isolation and fragmentation of the ten most abundant ions was performed by higher-389 energy collisional dissociation (HCD) (Olsen et al. 2007). Peptides with unassigned 390 charge states, as well as with charge states less than +2 were excluded from 391 fragmentation. The Orbitrap mass analyser was used for acquisition of fragment 392 spectra.

393 Peptide identification and quantification

394 Raw data files were analysed and peptides were identified using the MaxQuant 395 software (version 1.5.28) (Cox et al. 2009). Parent ion and MS² spectra were compared 396 to a database containing 92,578 human protein sequences obtained from UniProtKB 397 (release June 2018), coupled to the Andromeda search engine (Cox et al. 2011). Cysteine carbamidomethylation was set as a fixed modification. N-terminal acetylation, 398 399 oxidation, and N-ethylmaleimide (NEM) were set as variable modifications. For 400 ubiquitylome data analysis, glycine-glycine (GlyGly) modification of lysine was 401 additionally set as a variable modification. The mass tolerance for the spectra search was set to be lower than 6 ppm in MS and 20 ppm in HCD MS² mode. Spectra were 402

403 searched with strict trypsin specificity and allowing for up to three mis-cleavages. Site 404 localisation probabilities were determined by MaxQuant using the PTM scoring 405 algorithm as described previously (Elias and Gygi 2007; Cox and Mann 2008). Filtering 406 of the dataset was based on the posterior error probability to arrive at a false discovery 407 rate (FDR) < 1% estimated using a target-decoy approach. Proteins that were 408 categorised as "only identified by site", potential contaminants and reverse hits were 409 removed. Only proteins identified with at least two peptides (including at least one 410 unique peptide) and a SILAC ratio count of at least two were used for analysis. For AP 411 experiments, proteins that were quantified in at least two out of three experiments were 412 kept for further analysis. In total, we quantified 1,106 and 1,097 protein groups in the AP experiments with GFP-MKRN1^{wt} (Fig. 1A), GFP-MKRN1^{PAM2mut} (Fig. 1D) and 413 GFP-MKRN1^{RINGmut} (Supplemental Fig. S7), respectively (Supplemental Table S1). 414 415 The SILAC ratios were log₂ transformed and converted into an asymmetric z-score 416 based on the mean and interguartile range of the distribution as described previously 417 (Cox and Mann 2008). For statistical analysis, a moderated t-test from the limma 418 algorithm was used (Ritchie et al. 2015). Enriched proteins with an FDR < 5% were 419 determined to be significantly enriched interactors (for GFP-MKRN1^{wt}). For proteins enriched in GFP-MKRN1^{RINGmut} over GFP-MKRN1^{wt}, proteins with an FDR < 5% and 420 421 a GFP-MKRN1^{wt}/GFP z-score > 1 were selected. In the proteome experiment, we 422 quantified 6,439 protein groups, present in all three replicates. Ratio-ratio and ratio-423 intensity plots were created in R (version 3.4.3) using RStudio 424 (http://www.rstudio.com/).

425 Functional annotation of MKRN1 interactors and MKRN1-ubiquitylation targets.

In order to assess the functions of MKRN1-interacting proteins and proteins with
 MKRN1-dependent ubiquitylation sites, we performed gene ontology (GO) enrichment

analyses using the Database for Annotation, Visualization and Integrated Discovery
(DAVID 6.7) for three GO domains (Jiao et al. 2012). Enriched GO terms (modified
Fisher exact test, adjusted *P* value < 0.05, Benjamini-Hochberg correction;
Supplemental Fig. S2B, S8A) were visualised using REVIGO (Reduce & Visualize
Gene Ontology) allowing medium GO term similarity (Supek et al. 2011).

433 Western blot

434 Denatured proteins were separated by SDS-PAGE on a NuPAGE 4-12% Bis-Tris 435 protein gel (Life Technologies) and transferred to a 0.45 µm nitrocellulose membrane 436 (VWR). For detection, either fluorophore-coupled secondary antibodies or HRP-437 conjugated secondary antibodies and WesternBright Chemiluminescent Substrate 438 (Biozym Scientific) or SuperSignal West Pico Chemiluminescent Substrate (Life 439 Technologies) were used. Western blots were quantified by determining the 440 background-subtracted densities of the protein of interest using ImageJ (Schindelin et al. 2015). The signal from the AP (against GFP-tagged protein of interest) was 441 442 normalised to the respective control samples expressing the empty vector or to the 443 input.

444 Antibodies

The following antibodies were used: anti-GFP (B-2 clone; Santa Cruz; sc-9996), antiMKRN1 (Bethyl Laboratories, A300-990A), anti-PABPC1/3 (Cell Signaling, 4992), antiZnf598 (N1N3; GeneTex; GTX119245), anti-αTubulin (Sigma Aldrich, T-5168), antiRabbit IgG (Cell Signaling; 7074), anti-Mouse IgG (Cell Signaling; 7076), IRDye®
680RD Goat anti-Mouse IgG (P/N 925-68070), and IRDye® 800CW Goat anti-Rabbit
IgG (P/N 925-32211) (both LI-COR Biosciences GmbH).

451 **RNA isolation, cDNA synthesis and qPCR**

452 Cells were washed twice in ice-cold PBS and harvested. RNA was isolated using the 453 RNeasy Plus Mini Kit (Qiagen) according to the manufacturer's recommendations. 454 500 ng total RNA was transcribed into cDNA using random hexamer primers (Thermo 455 Scientific) and the RevertAid Reverse Transcriptase (Thermo Scientific) according to 456 the manufacturer's recommendations. qPCR was performed using the Luminaris 457 HiGreen gPCR Master Mix, low ROX (Thermo Scientific) according to the manufacturer's recommendations with 10 µM forward and reverse primers 458 459 (Supplemental Table S5).

460 iCLIP experiments and data processing

461 iCLIP libraries were prepared as described previously (Huppertz et al. 2014; Sutandy 462 et al. 2016). HEK293T cells ectopically expressing either GFP alone (empty vector) or N-terminally GFP-tagged MKRN1 wild type (GFP-MKRN1^{wt}), GFP-MKRN1^{PAM2mut}, or 463 464 GFP-MKRN1^{RINGmut} were used. For crosslinking, confluent cells were irradiated once with 150 mJ/cm² at 254 nm in a Stratalinker 2400 or treated with 4-thiouridine (100 µM 465 for 16 h) and irradiated with 3x 300 mJ/cm² in a Stratalinker 2400 with 365 nm bulbs. 466 For IP, 10.5 µg anti-GFP antibody (goat, Protein Unit, MPI-CBG, Dresden) were used 467 per sample. The libraries were sequenced as 50-nt single-end reads on an Illumina 468 469 MiSeq platform (Supplemental Table S2).

Basic sequencing quality checks were applied to all reads using FastQC (version
0.11.5) (https://www.bioinformatics.babraham.ac.uk/projects/fastqc/). Afterwards,
reads were filtered based on sequencing qualities (Phred score) of the barcode region.
Only reads with at most one position with a sequencing quality < 20 in the experimental
barcode (positions 4 to 7) and without any position with a sequencing quality < 17 in
the random barcode (positions 1-3 and 8-9) were kept for further analysis. Remaining

476 reads were de-multiplexed based on the experimental barcode on positions 4 to 7
477 using Flexbar (version 3.0.0) (Dodt et al. 2012) without allowing mismatches.

All following steps of the analysis were performed on all individual samples after demultiplexing. Remaining adapter sequences were trimmed from the right end of the reads using Flexbar (version 3.0.0) allowing up to one mismatch in 10 nt, requiring a minimal overlap of 1 nt of read and adapter. After trimming off the adapter, the barcode is trimmed off of the left end of the reads (first 9 nt) and added to the header of the read, such that the information is kept available for downstream analysis. Reads shorter than 15 nt were removed from further analysis.

Trimmed and filtered reads were mapped to the human genome (assembly version GRCh38) and its annotation based on GENCODE release 25 (Harrow et al. 2012) using STAR (version 2.5.4b) (Dobin et al. 2013). When running STAR, up to two mismatches were allowed, soft-clipping was prohibited and only uniquely mapping reads were kept for further analysis.

Following mapping, duplicate reads were marked using the dedup function of bamUtil (version 1.0.13), which defines duplicates as reads whose 5' ends map to the same position in the genome (https://github.com/statgen/bamUtil). Subsequently, marked duplicates with identical random barcodes were removed since they are considered technical duplicates, while biological duplicates showing unequal random barcodes were kept.

Resulting bam files were sorted and indexed using SAMtools (version 1.5) (Li et al.
2009). Based on the bam files, bedgraph files were created using bamToBed of the
BEDTools suite (version 2.25.0) (Quinlan and Hall 2010), considering only the position
upstream of the 5' mapping position of the read, since this nucleotide is considered as

500 the crosslinked nucleotide. bedgraph files were then transformed to bigWig file format

⁵⁰¹ using bedGraphToBigWig of the UCSC tool suite (Kent et al. 2010).

502 Identification and characterisation of MKRN1 binding sites

503 Peak calling was performed on merged iCLIP coverage tracks (crosslink events per 504 nucleotide) from the three replicates based on GENCODE annotation (release 27, 505 GRCh38) using ASPeak (version 2.0; default setting plus -nornaseq to estimate 506 parameters p and r for the negative binomial distributions in a 500-nt window around 507 each peak) (Kucukural et al. 2013). The initially predicted peaks were resized to 508 uniform 9-nt windows around their weighted centred as defined by ASPeak. To avoid 509 artefacts, we removed sparsely covered peaks that harbour crosslink events on less 510 than three nucleotides within the 9-nt region window. We iteratively merged all 511 remaining windows if overlapping by at least 1 nt, by defining the position with the 512 cumulative half maximum count of crosslink events as new window centre. We further 513 kept only reproducible windows with at least three crosslink events from any two 514 replicates. Finally, we excluded all windows overlapping with none or multiple protein-515 coding genes (GENCODE annotations support level \geq 2 and transcript support level \geq 516 3), and assign each binding site to a distinct genomic region (3' UTR, 5' UTR, CDS, 517 intron). Consistent with the mostly cytoplasmic localisation of MKRN1 (Miroci et al. 518 2012; Cassar et al. 2015; Hildebrandt et al. 2017), less than 6% of the binding sites 519 were predicted within introns, which were excluded from further analysis. This 520 procedure yielded a total of 7,331 MKRN1 binding sites in 2,163 genes.

In order to estimate binding site strength and to facilitate comparisons between binding sites (**Fig. 2B,D** and **Supplemental Fig. 3D-F, 4A,C,D**), we corrected for transcript abundance by representing the crosslink events within a binding site as a 'signal-overbackground' ratio (SOB). The respective background was calculated as the sum of

525 crosslink events outside of binding sites (plus 5 nt to either side) by the merged length 526 of all exons. 3' UTR lengths were restricted to 10 nt past the last MKRN1 binding site 527 or 500 nt if no binding site was present. SOB calculations were performed separately 528 for each replicate and then averaged. No SOB value was assigned for genes with a 529 background of < 10 crosslink events, resulting in SOB values for 97% of all binding 530 sites.

In order to assess the local RNA sequence context of MKRN1 binding sites (**Fig. 2B** and **Supplemental Fig. S4A**), enriched 4-mers were counted inside the 9-nt binding sites as well as within 40-nt before and after. To estimate an empirical background distribution, 1,000 9-nt windows were randomly picked in 3' UTRs and 4-mer frequencies were counted in the same windows. This process was repeated 100 times, and the resulting mean and standard deviation were used to calculate the z-score for each 4-mer.

538 In order to define the A-rich regions downstream of MKRN1 binding sites in 3' UTRs 539 (A-rich stretches), we used a maximisation approach in a 55-nt search space starting 540 from the binding site centre. Within this space, we calculated the percentage of A 541 nucleotides (A-content) for windows of increasing size (8-30 nt) and selected the 542 stretch with highest value for each window size. In case of ties, the window closer to 543 the binding site was preferred, resulting in a set of 23 candidate A-stretches with the 544 maximal A-content for each length. Next, we computed the longest continuous A run 545 (LCA) and a weighted A-content (multiplying the A-content with the number of A 546 nucleotides) for each candidate A-stretch. Candidate A-stretches with an A-content < 547 70%, a weighted A-content < 11 and an LCA < 4 were excluded. The final A-stretch 548 for each binding site was then selected in a hierarchical manner, preferring LCA over 549 weighted A-content. Lastly, overlapping A-stretches of neighbouring binding sites were

550 merged by selecting the highest scoring A-stretch, based on LCA and weighted A-551 content. In total, this procedure identified 1,412 non-overlapping A-stretches, 552 associated with 1,848 binding sites.

553 In order to estimate the extent of MKRN1 binding to poly(A) tails (Fig. 3B), we 554 evaluated the percentage of adenosine within the iCLIP reads that could not be 555 mapped to the human genome without soft-clipping (see above). iCLIP data for 556 heterogeneous nuclear ribonucleoprotein H (HNRNPH) served as control (Braun et 557 al. 2018). Annotated transcript 3' ends (i.e. polyadenylation sites) were taken from 558 GENCODE (all annotated protein-coding transcripts with support level ≤ 2 and 559 transcript support level ≤ 3; release 28, GRCh38.p12; <u>www.gencodegenes.org</u>). For 560 **Fig. 3C**, all crosslink events within a 2-kb window around the polyadenylation sites 561 for 3' UTR longer than 1 kb were counted.

562 Evolutionary characterisation of Makorin protein family

563 Four different ortholog searches were performed using HaMStR-OneSeg (Ebersberger 564 et al. 2014) against the Quest for Orthologs Consortium protein set, containing 78 species (release 2017 04) (Sonnhammer et al. 2014). For each run, a different seed 565 protein was chosen: human MKRN1-3 (UniProt identifiers Q9UHC7, Q9H000 and 566 567 Q13064) and MKRN4 from zebrafish (A9C4A6). In order to identify proteins with a 568 similar domain architecture, we calculated a unidirectional feature architecture 569 similarity (FAS) score which compares the domain architecture of the seed protein and 570 the predicted ortholog (Koestler et al. 2010). Predicted orthologues with FAS < 0.7 571 were removed after initial assessment. Finally, all vertebrate species and selected 572 invertebrate species were used for reconstruction of a maximum likelihood (ML) tree. 573 For this, protein sequences were aligned using MAFFT v7.294b L-INS-i (Katoh and 574 Standley 2013), and ML trees with 100 bootstrap replicates were calculated using

575 RAxML version 8.1.9 (Stamatakis 2014). Settings for a rapid bootstrap analysis and 576 searching for the best scoring ML tree in one program run (-f a) and an automatic 577 selection of the best fitting amino acid substitution model (-m PROTGAMMAAUTO) 578 were chosen. Reconstructed trees were visualised using FigTree v1.4.2 579 (http://tree.bio.ed.ac.uk/ software/figtree/).

The phylogenetic tree and FASTA sequences from the ortholog dataset were loaded into DoMosaics (<u>http://www.domosaics.net</u>) and Pfam domains were annotated with HMMER (<u>http://hmmer.org/</u>, default parameters). Since the PAM2 motif in all Makorin proteins differs from the described consensus motif (Albrecht and Lengauer 2004), a custom Hidden Markov Model was trained on PAM2 motifs from selected Makorin orthologs and used for a HMMER scan of the orthologs (no E-value cutoff). The same procedure was repeated for the PAM2-like motif (PAM2L) (Pohlmann et al. 2015).

587 Dual fluorescence translation stall assay via flow cytometry

588 Knockdowns were performed for 24 h, before the dual fluorescence reporter plasmids 589 were ectopically expressed for 48 h. Cells were washed in PBS and trypsinised. After 590 sedimentation, cells were resuspended in DPBS supplemented with 2 mM EDTA. 591 Cellular GFP and RFP fluorescence was measured using flow cytometry on a 592 LSRFortessa SORP (BD Biosciences). Data analysis was done using FlowJo (v10) 593 (FlowJo, LLC). For statistical testing, paired two-tailed Student's t-tests with Benjamini-594 Hochberg correction were performed on $n \ge 6$ replicates.

595 Ubiquitin remnant profiling

596 Di-glycine remnant profiling was performed as described before (Wagner et al. 2011; 597 Heidelberger et al. 2018). In four different experiments, isotope labels were assigned 598 as follows: experiment 1, *MKRN1* KD1 (siRNA1), *MKRN1* KD2 (siRNA2) and control 599 siRNA with light, medium and heavy SILAC labels, respectively; experiment 2, *MKRN1* 600 KD2 (siRNA2) and control siRNA with heavy and light SILAC labels, respectively; experiment 3, MKRN1 KD2 (siRNA2) and control siRNA with heavy and light SILAC 601 602 labels, respectively; experiment 3, MKRN1 KD2 (siRNA2) and control siRNA with light 603 and heavy SILAC labels, respectively. Cells were treated with the proteasome 604 inhibitors bortezomib (1 µM, 8h, replicate 1; Santa Cruz Biotechnology) or MG132 (10 605 µM, 2 h, replicates 2, 3, 4; Sigma). Proteins were precipitated in acetone. Proteins were digested with endoproteinase Lys-C (Wako Chemicals) and sequencing-grade 606 607 modified trypsin (Sigma). To purify the peptides, reversed-phase Sep-Pak C18 608 cartridges (Waters) were used. Modified peptides were enriched using di-glycine-609 lysine antibody resin (Cell Signaling Technology). The enriched peptides were eluted 610 with 0.15% trifluoroacetic acid in water, then fractionated using micro-column-based 611 strong-cation exchange chromatography (SCX) (Weinert et al. 2013) before being 612 desalted on reversed-phase C18 StageTips (Rappsilber et al. 2007). Samples were 613 analysed by quantitative mass spectrometry and MaxQuant as described above. To 614 identify significantly regulated ubiquitylation sites, the limma algorithm was applied 615 (Ritchie et al. 2015). A P value < 0.1 after multiple testing correction was used as a 616 cut-off to determine up- and downregulated ubiguitylation sites. Volcano and dot plots were created in R (version 3.4.3). 617

618 Functional interaction network of MKRN1 ubiquitylation target proteins

The functional protein interaction network analysis was performed by integrating interaction data from the STRING database (score > 0.4), the BioGrid database and our own findings (Franceschini et al. 2013; Chatr-Aryamontri et al. 2017). Cytoscape (version 3.6.1) was used to visualise the protein interaction network (Saito et al. 2012).

623

624

625 Availability of data and materials

The mass spectrometry proteomics data have been deposited to the
ProteomeXchange Consortium (http://proteomecentral.proteomexchange.org) via the
PRIDE partner repository with the dataset identifier PXD011772.

Raw and processed iCLIP data are available at GEO under the accession numberGSE122869.

631 Acknowledgements

We would like to thank all members of the Zarnack, König and Beli labs, as well as 632 René Ketting, Nadine Wittkopp, and Miguel Almeida for fruitful discussions. The 633 634 authors gratefully thank the Ramanujan S. Hegde for providing the dual fluorescence reporter plasmids. We thank Anja Freiwald for assistance with mass spectrometry 635 analysis and Dr. Stefan Simm for assistance with evaluating PAM2 mutations. The 636 637 support of the IMB Core Facilities Bioinformatics, Flow Cytometry, Genomics, and the use of its Illumina MiSeq, as well as the DFG-funded Mass Spectrometer Q Exactive 638 639 Plus (INST 247/766-1 FUGG) are gratefully acknowledged. K.Z. was supported by the 640 LOEWE program Ubiquitin Networks (Ub-Net) of the State of Hesse (Germany) and 641 the SFB 902 of the German Research Foundation. P.B. is supported by the Emmy 642 Noether Program (BE 5342/1-1), the SFB 1177 of the German Research Foundation 643 and the Marie Curie Career Integration Grant from the European Commission (grant 644 agreement number: 630763). The project was funded by the German Research 645 Foundation (DFG) as part of SPP1925 to J.-Y.R. (RO 4681/4-1) and J.K. (KO 4566/3-1). Animal shapes in Supplemental Fig. S1A were obtained from PhyloPic and are 646 647 used under the Creative Common Attribution-NonCommercial-ShareAlike 3.0 648 Unported license.

649 Author contributions

650 A.H. performed iCLIP experiments, flow cytometry measurements of dual fluorescence reporters and most proteomics experiments. M.B. performed most bioinformatics 651 652 analyses of MKRN1 iCLIP data. C.R. analysed MKRN1 binding at polyadenylation sites and poly(A) tails. A.B. and S.B. performed initial iCLIP data processing and 653 654 analysis. A.H. and A.B. analysed the proteomics data. J.B.H. and A.V. contributed to 655 replicate ubiquitin remnant profiling experiments and AP-Western blot experiments, respectively. H.H. performed replicate iCLIP and replicate AP-Western blot 656 657 experiments. C.R. and I.E. contributed evolutionary characterisation of Makorin proteins. A.D. and J.-Y.R. performed complementary studies in *D. melanogaster*. S.E. 658 and K.Z. supervised the bioinformatics analyses. J.K. and P.B. conceived the project 659 660 with K.Z. and supervised the experimental work. A.H., J.K., K.Z. and P.B. wrote the 661 manuscript with help and comments from all co-authors.

662 References

669

- Albrecht M, Lengauer T. 2004. Survey on the PABC recognition motif PAM2. *Biochem Biophys Res Commun* **316**: 129-138.
- Arthur L, Pavlovic-Djuranovic S, Smith-Koutmou K, Green R, Szczesny P, Djuranovic
 S. 2015. Translational control by lysine-encoding A-rich sequences. *Sci Adv* 1.
- Bag J. 2001. Feedback inhibition of poly(A)-binding protein mRNA translation. A
 possible mechanism of translation arrest by stalled 40 S ribosomal subunits. *J Biol*
- 670 Bengtson MH, Joazeiro CA. 2010. Role of a ribosome-associated E3 ubiquitin ligase 671 in protein guality control. *Nature* **467**: 470-473.

Chem 276: 47352-47360.

- Böhne A, Darras A, D'Cotta H, Baroiller JF, Galiana-Arnoux D, Volff JN. 2010. The
 vertebrate makorin ubiquitin ligase gene family has been shaped by large-scale
 duplication and retroposition from an ancestral gonad-specific, maternal-effect
 gene. *BMC Genomics* 11: 721.
- Brandman O, Hegde RS. 2016. Ribosome-associated protein quality control. *Nat Struct Mol Biol* 23: 7-15.
- Brandman O, Stewart-Ornstein J, Wong D, Larson A, Williams CC, Li GW, Zhou S,
 King D, Shen PS, Weibezahn J et al. 2012. A ribosome-bound quality control
 complex triggers degradation of nascent peptides and signals translation stress. *Cell* **151**: 1042-1054.
- Braun S, Enculescu M, Setty ST, Cortés-López M, de Almeida BP, Sutandy FXR,
 Schulz L, Busch A, Seiler M, Ebersberger S et al. 2018. Decoding a cancerrelevant splicing decision in the *RON* proto-oncogene using high-throughput
 mutagenesis. *Nat Commun* **9**: 3315.
- Cassar PA, Carpenedo RL, Samavarchi-Tehrani P, Olsen JB, Park CJ, Chang WY,
 Chen Z, Choey C, Delaney S, Guo H et al. 2015. Integrative genomics positions
 MKRN1 as a novel ribonucleoprotein within the embryonic stem cell gene
 regulatory network. *EMBO Rep* 16: 1334-1357.
- Chatr-Aryamontri A, Oughtred R, Boucher L, Rust J, Chang C, Kolas NK, O'Donnell L,
 Oster S, Theesfeld C, Sellam A et al. 2017. The BioGRID interaction database:
 2017 update. *Nucleic Acids Res* 45: D369-D379.

- 693 Choe YJ, Park SH, Hassemer T, Korner R, Vincenz-Donnelly L, Hayer-Hartl M, Hartl
- FU. 2016. Failure of RQC machinery causes protein aggregation and proteotoxic
 stress. *Nature* 531: 191-195.
- 696 Chu J, Hong NA, Masuda CA, Jenkins BV, Nelms KA, Goodnow CC, Glynne RJ, Wu

697 H, Masliah E, Joazeiro CA et al. 2009. A mouse forward genetics screen identifies

698 LISTERIN as an E3 ubiquitin ligase involved in neurodegeneration. *Proc Natl Acad*

- 699 Sci U S A **106**: 2097-2103.
- Collaboration O. 2016. The ORFeome Collaboration: a genome-scale human ORFclone resource. *Nat Methods* 13: 191-192.
- Cox J, Mann M. 2008. MaxQuant enables high peptide identification rates,
 individualized p.p.b.-range mass accuracies and proteome-wide protein
 quantification. *Nat Biotechnol* 26: 1367-1372.
- Cox J, Matic I, Hilger M, Nagaraj N, Selbach M, Olsen JV, Mann M. 2009. A practical
 guide to the MaxQuant computational platform for SILAC-based quantitative
 proteomics. *Nat Protoc* 4: 698-705.
- Cox J, Neuhauser N, Michalski A, Scheltema RA, Olsen JV, Mann M. 2011.
 Andromeda: a peptide search engine integrated into the MaxQuant environment.
 J Proteome Res 10: 1794-1805.
- 711 Deo RC, Sonenberg N, Burley SK. 2001. X-ray structure of the human hyperplastic
- discs protein: an ortholog of the C-terminal domain of poly(A)-binding protein. *Proc Natl Acad Sci U S A* **98**: 4414-4419.
- Dobin A, Davis CA, Schlesinger F, Drenkow J, Zaleski C, Jha S, Batut P, Chaisson M,
 Gingeras TR. 2013. STAR: ultrafast universal RNA-seq aligner. *Bioinformatics* 29:
 15-21.
- Dodt M, Roehr JT, Ahmed R, Dieterich C. 2012. FLEXBAR-Flexible Barcode and
 Adapter Processing for Next-Generation Sequencing Platforms. *Biology (Basel)* 1:
 895-905.
- Ebersberger I, Simm S, Leisegang MS, Schmitzberger P, Mirus O, von Haeseler A,
 Bohnsack MT, Schleiff E. 2014. The evolution of the ribosome biogenesis pathway
 from a yeast perspective. *Nucleic Acids Res* 42: 1509-1523.

Elias JE, Gygi SP. 2007. Target-decoy search strategy for increased confidence in
 large-scale protein identifications by mass spectrometry. *Nat Methods* 4: 207-214.

- Franceschini A, Szklarczyk D, Frankild S, Kuhn M, Simonovic M, Roth A, Lin J,
 Minguez P, Bork P, von Mering C et al. 2013. STRING v9.1: protein-protein
 interaction networks, with increased coverage and integration. *Nucleic Acids Res*41: D808-815.
- Garzia A, Jafarnejad SM, Meyer C, Chapat C, Gogakos T, Morozov P, Amiri M, Shapiro
 M, Molina H, Tuschl T et al. 2017. The E3 ubiquitin ligase and RNA-binding protein
 ZNF598 orchestrates ribosome quality control of premature polyadenylated
 mRNAs. *Nat Commun* 8: 16056.
- Gray TA, Hernandez L, Carey AH, Schaldach MA, Smithwick MJ, Rus K, Marshall
 Graves JA, Stewart CL, Nicholls RD. 2000. The ancient source of a distinct gene
 family encoding proteins featuring RING and C(3)H zinc-finger motifs with
 abundant expression in developing brain and nervous system. *Genomics* 66: 7686.
- Hafner M, Landthaler M, Burger L, Khorshid M, Hausser J, Berninger P, Rothballer A,
 Ascano M, Jr., Jungkamp AC, Munschauer M et al. 2010. Transcriptome-wide
 identification of RNA-binding protein and microRNA target sites by PAR-CLIP. *Cell*141: 129-141.
- Harrow J, Frankish A, Gonzalez JM, Tapanari E, Diekhans M, Kokocinski F, Aken BL,
 Barrell D, Zadissa A, Searle S et al. 2012. GENCODE: the reference human
 genome annotation for The ENCODE Project. *Genome Res* 22: 1760-1774.
- Heidelberger JB, Voigt A, Borisova ME, Petrosino G, Ruf S, Wagner SA, Beli P. 2018.
 Proteomic profiling of VCP substrates links VCP to K6-linked ubiquitylation and cMyc function. *EMBO Rep* 19.
- Hildebrandt A, Alanis-Lobato G, Voigt A, Zarnack K, Andrade-Navarro MA, Beli P,
 König J. 2017. Interaction profiling of RNA-binding ubiquitin ligases reveals a link
 between posttranscriptional regulation and the ubiquitin system. *Sci Rep* 7: 16582.
- Huppertz I, Attig J, D'Ambrogio A, Easton LE, Sibley CR, Sugimoto Y, Tajnik M, König
 J, Ule J. 2014. iCLIP: protein-RNA interactions at nucleotide resolution. *Methods*65: 274-287.

Jamar NH, Kritsiligkou P, Grant CM. 2018. Loss of mRNA surveillance pathways
 results in widespread protein aggregation. *Sci Rep* 8: 3894.

Jiao X, Sherman BT, Huang da W, Stephens R, Baseler MW, Lane HC, Lempicki RA.

2012. DAVID-WS: a stateful web service to facilitate gene/protein list analysis. *Bioinformatics* 28: 1805-1806.

- Joazeiro CAP. 2017. Ribosomal Stalling During Translation: Providing Substrates for
 Ribosome-Associated Protein Quality Control. *Annu Rev Cell Dev Biol* 33: 343368.
- Juszkiewicz S, Chandrasekaran V, Lin Z, Kraatz S, Ramakrishnan V, Hegde RS. 2018.
- 763 ZNF598 Is a Quality Control Sensor of Collided Ribosomes. *Mol Cell* 72: 469-481
 764 e467.
- Juszkiewicz S, Hegde RS. 2017. Initiation of Quality Control during Poly(A) Translation
 Requires Site-Specific Ribosome Ubiquitination. *Mol Cell* 65: 743-750 e744.
- Kaida D, Berg MG, Younis I, Kasim M, Singh LN, Wan L, Dreyfuss G. 2010. U1 snRNP
 protects pre-mRNAs from premature cleavage and polyadenylation. *Nature* 468:
 664-668.
- Katoh K, Standley DM. 2013. MAFFT multiple sequence alignment software version 7:
 improvements in performance and usability. *Mol Biol Evol* **30**: 772-780.
- Kent WJ, Zweig AS, Barber G, Hinrichs AS, Karolchik D. 2010. BigWig and BigBed:
 enabling browsing of large distributed datasets. *Bioinformatics* 26: 2204-2207.
- Kim JH, Park KW, Lee EW, Jang WS, Seo J, Shin S, Hwang KA, Song J. 2014.
 Suppression of PPARgamma through MKRN1-mediated ubiquitination and
 degradation prevents adipocyte differentiation. *Cell Death Differ* 21: 594-603.
- Kim JH, Park SM, Kang MR, Oh SY, Lee TH, Muller MT, Chung IK. 2005. Ubiquitin
 ligase MKRN1 modulates telomere length homeostasis through a proteolysis of
 hTERT. *Genes Dev* 19: 776-781.
- Kini HK, Silverman IM, Ji X, Gregory BD, Liebhaber SA. 2016. Cytoplasmic poly(A)
 binding protein-1 binds to genomically encoded sequences within mammalian
 mRNAs. *RNA* 22: 61-74.

- Ko A, Shin JY, Seo J, Lee KD, Lee EW, Lee MS, Lee HW, Choi IJ, Jeong JS, Chun
 KH et al. 2012. Acceleration of gastric tumorigenesis through MKRN1-mediated
 posttranslational regulation of p14ARF. *J Natl Cancer Inst* 104: 1660-1672.
- Koestler T, von Haeseler A, Ebersberger I. 2010. FACT: functional annotation transfer
 between proteins with similar feature architectures. *BMC Bioinformatics* 11: 417.
- König J, Zarnack K, Rot G, Curk T, Kayikci M, Zupan B, Turner DJ, Luscombe NM,
 Ule J. 2010. iCLIP reveals the function of hnRNP particles in splicing at individual
 nucleotide resolution. *Nat Struct Mol Biol* **17**: 909-915.
- 791 Kozlov G, De Crescenzo G, Lim NS, Siddiqui N, Fantus D, Kahvejian A, Trempe JF,

Elias D, Ekiel I, Sonenberg N et al. 2004. Structural basis of ligand recognition by

PABC, a highly specific peptide-binding domain found in poly(A)-binding protein

and a HECT ubiquitin ligase. *EMBO J* 23: 272-281.

- Kozlov G, Menade M, Rosenauer A, Nguyen L, Gehring K. 2010. Molecular
 determinants of PAM2 recognition by the MLLE domain of poly(A)-binding protein. *J Mol Biol* **397**: 397-407.
- Kozlov G, Trempe JF, Khaleghpour K, Kahvejian A, Ekiel I, Gehring K. 2001. Structure
 and function of the C-terminal PABC domain of human poly(A)-binding protein. *Proc Natl Acad Sci U S A* 98: 4409-4413.
- Kucukural A, Ozadam H, Singh G, Moore MJ, Cenik C. 2013. ASPeak: an abundance
 sensitive peak detection algorithm for RIP-Seq. *Bioinformatics* 29: 2485-2486.
- Lee EW, Lee MS, Camus S, Ghim J, Yang MR, Oh W, Ha NC, Lane DP, Song J. 2009.
 Differential regulation of p53 and p21 by MKRN1 E3 ligase controls cell cycle arrest
 and apoptosis. *EMBO J* 28: 2100-2113.
- Letzring DP, Wolf AS, Brule CE, Grayhack EJ. 2013. Translation of CGA codon
 repeats in yeast involves quality control components and ribosomal protein L1.
 RNA 19: 1208-1217.
- Li H, Handsaker B, Wysoker A, Fennell T, Ruan J, Homer N, Marth G, Abecasis G,
 Durbin R, Genome Project Data Processing S. 2009. The Sequence
 Alignment/Map format and SAMtools. *Bioinformatics* 25: 2078-2079.

- Lima SA, Chipman LB, Nicholson AL, Chen YH, Yee BA, Yeo GW, Coller J, Pasquinelli
- AE. 2017. Short poly(A) tails are a conserved feature of highly expressed genes.
 Nat Struct Mol Biol 24: 1057-1063.
- Lu J, Deutsch C. 2008. Electrostatics in the ribosomal tunnel modulate chain elongation rates. *J Mol Biol* **384**: 73-86.
- Lyabin DN, Eliseeva IA, Skabkina OV, Ovchinnikov LP. 2011. Interplay between Ybox-binding protein 1 (YB-1) and poly(A) binding protein (PABP) in specific regulation of YB-1 mRNA translation. *RNA Biol* **8**: 883-892.
- 820 Michalski A, Damoc E, Hauschild JP, Lange O, Wieghaus A, Makarov A, Nagaraj N,
- 821 Cox J, Mann M, Horning S. 2011. Mass spectrometry-based proteomics using Q
- 822 Exactive, a high-performance benchtop quadrupole Orbitrap mass spectrometer.
- 823 *Mol Cell Proteomics* **10**: M111 011015.
- Miroci H, Schob C, Kindler S, Olschlager-Schutt J, Fehr S, Jungenitz T, Schwarzacher
 SW, Bagni C, Mohr E. 2012. Makorin ring zinc finger protein 1 (MKRN1), a novel
 poly(A)-binding protein-interacting protein, stimulates translation in nerve cells. J
- 827 Biol Chem **287**: 1322-1334.
- Nielsen ML, Vermeulen M, Bonaldi T, Cox J, Moroder L, Mann M. 2008.
 Iodoacetamide-induced artifact mimics ubiquitination in mass spectrometry. *Nat Methods* 5: 459-460.
- Olsen JV, Macek B, Lange O, Makarov A, Horning S, Mann M. 2007. Higher-energy
 C-trap dissociation for peptide modification analysis. *Nat Methods* 4: 709-712.
- Omwancha J, Zhou XF, Chen SY, Baslan T, Fisher CJ, Zheng Z, Cai C, Shemshedini
 L. 2006. Makorin RING finger protein 1 (MKRN1) has negative and positive effects
 on RNA polymerase II-dependent transcription. *Endocrine* 29: 363-373.
- Pohlmann T, Baumann S, Haag C, Albrecht M, Feldbrügge M. 2015. A FYVE zinc
 finger domain protein specifically links mRNA transport to endosome trafficking. *Elife* 4.
- Quinlan AR, Hall IM. 2010. BEDTools: a flexible suite of utilities for comparing genomic
 features. *Bioinformatics* 26: 841-842.

- 841 Rappsilber J, Mann M, Ishihama Y. 2007. Protocol for micro-purification, enrichment,
- pre-fractionation and storage of peptides for proteomics using StageTips. *Nat Protoc* 2: 1896-1906.
- Ritchie ME, Phipson B, Wu D, Hu Y, Law CW, Shi W, Smyth GK. 2015. limma powers
 differential expression analyses for RNA-sequencing and microarray studies. *Nucleic Acids Res* 43: e47.
- Saito R, Smoot ME, Ono K, Ruscheinski J, Wang PL, Lotia S, Pico AR, Bader GD,
 Ideker T. 2012. A travel guide to Cytoscape plugins. *Nat Methods* 9: 1069-1076.
- Salvatico J, Kim JH, Chung IK, Muller MT. 2010. Differentiation linked regulation of
 telomerase activity by Makorin-1. *Mol Cell Biochem* **342**: 241-250.
- Schindelin J, Rueden CT, Hiner MC, Eliceiri KW. 2015. The ImageJ ecosystem: An
 open platform for biomedical image analysis. *Mol Reprod Dev* 82: 518-529.
- 853 Simms CL, Yan LL, Zaher HS. 2017. Ribosome Collision Is Critical for Quality Control
 854 during No-Go Decay. *Mol Cell* 68: 361-373 e365.
- Simsek D, Tiu GC, Flynn RA, Byeon GW, Leppek K, Xu AF, Chang HY, Barna M. 2017.
 The Mammalian Ribo-interactome Reveals Ribosome Functional Diversity and
 Heterogeneity. *Cell* 169: 1051-1065 e1018.
- Sonnhammer EL, Gabaldon T, Sousa da Silva AW, Martin M, Robinson-Rechavi M,
 Boeckmann B, Thomas PD, Dessimoz C, Quest for Orthologs c. 2014. Big data
 and other challenges in the quest for orthologs. *Bioinformatics* **30**: 2993-2998.
- 861 Stamatakis A. 2014. RAxML version 8: a tool for phylogenetic analysis and post-862 analysis of large phylogenies. *Bioinformatics* **30**: 1312-1313.
- Sundaramoorthy E, Leonard M, Mak R, Liao J, Fulzele A, Bennett EJ. 2017. ZNF598
 and RACK1 Regulate Mammalian Ribosome-Associated Quality Control Function
 by Mediating Regulatory 40S Ribosomal Ubiguitylation. *Mol Cell* 65: 751-760 e754.
- Supek F, Bosnjak M, Skunca N, Smuc T. 2011. REVIGO summarizes and visualizes
 long lists of gene ontology terms. *PLoS One* 6: e21800.
- Sutandy FXR, Ebersberger S, Huang L, Busch A, Bach M, Kang HS, Fallmann J,
 Maticzka D, Backofen R, Stadler PF et al. 2018. In vitro iCLIP-based modeling
 uncovers how the splicing factor U2AF2 relies on regulation by cofactors. *Genome Res* 28: 699-713.

- Sutandy FXR, Hildebrandt A, König J. 2016. Profiling the Binding Sites of RNA-Binding
 Proteins with Nucleotide Resolution Using iCLIP. *Methods Mol Biol* 1358: 175-195.
- Verma R, Oania RS, Kolawa NJ, Deshaies RJ. 2013. Cdc48/p97 promotes
 degradation of aberrant nascent polypeptides bound to the ribosome. *Elife* 2:
 e00308.
- Wagner SA, Beli P, Weinert BT, Nielsen ML, Cox J, Mann M, Choudhary C. 2011. A
 proteome-wide, quantitative survey of in vivo ubiquitylation sites reveals
 widespread regulatory roles. *Mol Cell Proteomics* 10: M111 013284.
- 880 Webster MW, Chen YH, Stowell JAW, Alhusaini N, Sweet T, Graveley BR, Coller J,
- 881 Passmore LA. 2018. mRNA Deadenylation Is Coupled to Translation Rates by the
- Differential Activities of Ccr4-Not Nucleases. *Mol Cell* **70**: 1089-1100 e1088.
- 883 Weinert BT, Scholz C, Wagner SA, Iesmantavicius V, Su D, Daniel JA, Choudhary C.
- 884 2013. Lysine succinylation is a frequently occurring modification in prokaryotes and
- eukaryotes and extensively overlaps with acetylation. *Cell Rep* **4**: 842-851.

Supplemental Tables and Legends

Supplemental Tables S1,3,4

Provided as Excel files

Supplemental Tables S2: Summary of MKRN1 iCLIP experiments. iCLIP experiments with GFP-MKRN1 were performed in three independent replicates.

	Sequenced	Uniquely	Crosslink events	Binding sites
	reads	mapped reads		
Replicate 1	3,418,021	1,561,445	957,097	-
Replicate 2	6,527,256	3,149,583	1,972,821	-
Replicate 3	4,660,274	2,515,161	2,293,633	-
Total	14,605,551	7,226,189	5,223,551	7,331

Supplemental Table S5: Primers used in this study.

Name	Sequence 5' – 3'	Comment
PAMmut	GCCGTTGCCGGGCAACCCTACTGTGGC	MKRN1 ^{PAM2mut}
	CTCAATAGAATTCACCCAGTCCTCTGAACC	mutant
H307E	CAACTGCAACGAAACCTACTGTCTCAAG	MKRN1 ^{RINGmut}
	GAGAGGATCCCGAAGCGG	mutant
MKRN1 qPCR	CGATACGGGGAGAACTGTGT	MKRN1 qPCR
	CCTTCTCATGGGCCTCAAT	[–] primer
ZNF598 qPCR	AACCTCGACAAATGGTCCTG	<i>ZNF598</i> qPCR
	GTCTTCGTCCTTGAGCTTCG	[¯] primer
ßActin qPCR	TCCTCCCTGGAGAAGAGCTAC	ß-actin qPCR
	TGGAGTTGAAGGTAGTTCGTG	primer

Supplemental Table S6: siRNAs used in this study.

Name	Sequence 5' – 3'	Comment
MKRN1 siRNA1	CAGGCGAAGCUGAGUCAAGAA[dT][dT]	Ko et al. 2012
MKRN1 siRNA2	CGGGAUCCUCUCCAACUGCAA[dT][dT]	Kim et al. 2014
ZNF598 siRNA	CCCUCUAAAGUUGGGAAGA[dT][dT]	Sigma, Rosetta
		predictions
Control siRNA	UGGUUUACAUGUCGACUAA[dT][dT]	Heidelberger et al.
		2018

886 Figure legends

887 Figure 1. MKRN1 interacts with PABP and other regulators of translation and RNA 888 stability. (A) Protein interactome of GFP-MKRN1^{wt} in HEK293T cells analysed by 889 quantitative MS-based proteomics. Combined SILAC ratios (n = 3 replicates) after zscore normalisation are plotted against log₁₀-transformed intensities. 1,100 protein 890 891 groups were quantified in at least two out of three replicate experiments. MKRN1 and 892 significant interactors are highlighted (FDR < 5%). (B) A PAM2 motif similar to the previously reported consensus (shown on top; Supplemental Fig. S1B) (Albrecht and 893 Lengauer 2004) is present in MKRN1 (first amino acid position indicated on the left). 894 Introduced mutations in MKRN1^{PAM2mut} are indicated in red below. Relevant positions 895 896 are highlighted (Supplemental Fig. S1B). (C) Endogenous PABP interacts with MKRN1^{wt} and MKRN1^{RINGmut}, but not with MKRN1^{PAM2mut}. Western blots for 897 898 endogenous PABPC1 and GFP (two exposure times, exp.) after AP of GFP-MKRN1 899 (wt and mutants). Ratios of PABP signal (normalised to input) in GFP-MKRN APs over 900 control (GFP empty vector, EV) are shown on the right. Replicates 2, 3, and uncropped 901 gel images are shown in Supplemental Fig. S9A-C. (D) Quantitative comparison of the interactomes of GFP-MKRN1^{wt} and GFP-MKRN1^{PAM2mut} shows that PABP and 902 903 several other interactors are lost upon PAM2 mutation. Combined ratios of three replicates are shown in a scatter plot. Only proteins detected in at least two out of three 904 905 replicates are shown. MKRN1^{wt} significant interactors (from A) are highlighted as in (A) 906 (FDR < 5% in MKRN1^{wt}).

907 Figure 2. MKRN1 binds upstream of A-stretches in 3' UTRs. (A) MKRN1 908 predominantly binds in the 3' UTR of protein-coding genes. Piecharts summarising the 909 distribution of MKRN1 binding sites to different RNA biotypes (7,331 binding sites, left) 910 and different regions within protein-coding transcripts (6.913 binding sites, right). (B) 911 MKRN1 binding sites display a downstream enrichment of AAAA homopolymers. 912 Frequency per nucleotide (nt) for four homopolymeric 4-mers in a 101-nt window 913 around the midpoints of the top 20% MKRN1 binding sites (according to signal-over-914 background; see Material and methods). (C) MKRN1 crosslink events accumulate 915 upstream of A-stretches. Metaprofile (top) shows the mean crosslink events per nt in 916 a 201-nt window around the start position of 1,412 MKRN1-associated A-stretches in 917 3' UTRs. Heatmap visualisation (bottom) displays crosslink events per nt (see colour 918 scale) in a 101-nt window around the MKRN1-associated A-stretches. (D) MKRN1 919 binding site strength (signal-over-background, SOB) increases with length of longest 920 continuous run of A's (LCA) within the A-stretch. Mean and standard deviation of 921 MKRN1 binding sites associated with A-stretches harbouring LCAs of increasing 922 length (x-axis). MKRN1 binding sites without associated A-stretches are shown for 923 comparison on the left. Number of binding sites in each category indicated as barchart 924 above. (E) MKRN1 binds upstream of A-stretches in the 3' UTR of the LARP1 gene. 925 Genome browser view of GFP-MKRN1 iCLIP data showing crosslink events per nt 926 (merged replicates, turquois) together with binding sites (lilac) and associated A-927 stretches (dark green).

928 Figure 3. MKRN1 binds at poly(A) tails. (A) MKRN1 binds near the polyadenylation 929 site of the SRSF4 gene. Genome browser view as in Fig. 2E. (B) Unmapped MKRN1 930 iCLIP reads display increased A-content (more than half of all nucleotides in the read), 931 evidencing poly(A) tail binding. Cumulative fraction of iCLIP reads (y-axis, merged 932 replicates) that could not be mapped to the human genome (see Materials and 933 methods) and show at least a given A-content (x-axis). iCLIP data for the unrelated 934 RBP HNRNPH (Braun et al. 2018) are shown for comparison. (C) MKRN1 crosslink 935 events increase towards 3'UTR ends. Metaprofile shows the sum of crosslink events 936 per nt in a 2001-nt window around annotated polyadenylation sites of transcripts with 937 >1 kb 3' UTRs (n = 11,257). (D) Overall RNA binding of MKRN1 is strongly reduced 938 when abrogating PABP interaction. Audioradiograph (left) of UV crosslinking 939 experiments (replicate 1, with 4SU and UV crosslinking at 365 nm; replicates 2 and 3 940 in **Supplemental Fig. S5**) comparing GFP-MKRN1^{PAM2mut} with GFP-MKRN1^{wt} at 941 different dilution steps for calibration. Quantification of radioactive signal of protein-942 RNA complexes and corresponding Western blots shown on the right. Uncropped gel 943 images are shown in Supplemental Fig. S10.

944 **Figure 4.** MKRN1 stalls ribosomes at poly(A) sequences. (A) The dual fluorescence 945 reporter harbours an N-terminal GFP, followed by a FLAG-SR-X linker and a C-946 terminal RFP, which are separated by P2A sites to ensure translation into three 947 separate proteins (Juszkiewicz and Hegde 2017). The resulting GFP:RFP ratio was 948 determined using flow cytometry. The inserted fragment K(AAA)₂₀ encodes 20 lysines 949 by repeating the codon AAA. The starting vector without insert (K₀) served as control. 950 Schematic ribosomes illustrate translation of the respective reporter segments. (B) 951 Ribosomes are efficiently stalled at K(AAA)₂₀ in HEK293T cells. Median RFP:GFP 952 ratios, normalised to K₀, are shown. Error bars represent standard deviation of the 953 mean (s.d.m., n = 6 replicates). P value indicated above (paired two-tailed t-test). (C) 954 Ribosomes fail to stall in the absence of MKRN1. HEK293T cells were transfected with 955 control siRNA or siRNAs targeting MKRN1 (KD1 and KD2) or ZNF598 for 24 h, 956 followed by transfection of the reporter plasmids for 48 h. Western blots for KDs are 957 shown in Supplemental Fig. S6B. RFP and GFP signals were analysed by flow 958 cytometry. Median RFP:GFP ratios, normalised to K₀ in control, are shown. Error bars 959 represent s.d.m.; P values indicated above (paired two-tailed t-test, Benjamini-960 Hochberg correction, $n \ge 6$ replicates; ns, not significant).

961 Figure 5. MKRN1 ubiquitylates ribosomal protein RPS10 and translational regulators. (A) Ubiquitin remnant profiling to compare the relative abundance of ubiquitylation sites 962 963 in *MKRN1* KD2 and control HEK293T cells. Ubiquitin remnant peptides were enriched 964 and analysed by quantitative mass spectrometry, quantifying a total of 15,528 965 ubiguitylation sites on 4,790 proteins. 29 putative MKRN1 target sites with significantly 966 decreased ubiquitylation upon MKRN1 KD2 (FDR < 10%, n = 4 replicates) are 967 highlighted and labelled with the respective protein name. Note that many proteins contain several differentially regulated ubiquitylation sites. (B) Protein interaction 968 969 network of 21 proteins with putative MKRN1 ubiquitylation target sites (significantly 970 reduced, shown in (A)). The functional interactions were obtained from the STRING 971 and BioGrid databases and our study. Visualisation by Cytoscape. (C) Ubiquitin 972 remnant profiling results for significantly regulated ubiguitylation sites (FDR < 10%) in 973 proteins from network in (B). Mean and standard deviation of the mean (s.d.m., error 974 bars) are given together with all data points. (D) Ubiquitin remnant profiling results for 975 seven quantified ubiquitylation sites in RPS10 and RPS20. Significant changes are 976 shown in black (FDR < 10%) and non-significant changes in grey. Representation as 977 in (C). (E) Comparison of ubiquitylation sites in selected target proteins that are 978 modified by ZNF598 and MKRN1 during RQC. (F) Comparison of enriched proteins 979 from the interactomes for GFP-MKRN1^{wt} (over GFP, see Fig. 1A) and GFP-MKRN1^{RINGmut} (over GFP-MKRN1^{wt}, see **Supplemental Fig. S7B**) with the proteins 980 981 containing MKRN1 ubiquitylation targets sites (UB, see (A)). Protein names of 982 overlapping targets are given.

983

984 Figure 6. MKRN1 is a sensor for poly(A) sequences that stalls ribosomes to initiate 985 ribosome-associated quality control. Proposed model of MKRN1 function: MKRN1 is positioned upstream of (premature) poly(A) tails via interaction with PABP. Ribosomes 986 987 translating the open reading frame run into MKRN1 that acts as a roadblock to prohibit 988 poly(A) translation. Upon contact with the translating ribosome, MKRN1 ubiguitylates 989 the 40S ribosomal protein RPS10. This stalls the ribosome, causing the trailing 990 ribosomes to collide. ZNF598 recognises the collided ribosomes and ubiquitylates 991 ribosomal proteins to promote RQC.

992

993

994 Supplemental figure legends

995 Supplemental Figure S1. Maximum likelihood tree of Makorin orthologs with their 996 protein domain architecture. (A) Maximum likelihood tree with 100 bootstrap replicates 997 of selected vertebrate and invertebrate orthologs and C. elegans as an outgroup. 998 Bootstrap values at each node indicate the number of replicates (out of 100) that 999 support the local tree structure and thereby serve as confidence estimates. Protein 1000 schematics (drawn to scale) on the right depict protein domains corresponding to the 1001 following PFAM domains: RING-type zinc finger, PF13445; MKRN1 C-terminus, 1002 PF15815; CCCH zinc finger, PF15663, PF14608 and PF00642. PAM2 motifs, predicted to interact with the MLLE domain of PABP proteins (Kozlov et al. 2001) as 1003 1004 well as the recently reported derivative PAM2L (Pohlmann et al. 2015), were added 1005 separately (see Material and methods). Abbreviated and full species names with 1006 corresponding UniProt identifiers in order of appearance: ANOGA, Anopheles 1007 gambiae, Q7QF83; BOVIN, Bos taurus, F1MF12, F6QQR5; BRAFL, Branchiostoma floridae, C3Y7M0; CAEEL, Caenorhabditis elegans, Q9N373; CANLF, Canis lupus, 1008 J9P921, E2RRA5, E2REH2, J9P9K3; DANRE, Danio rerio, Q4VBT5, Q9DFG8, 1009

1010 A9C4A6; DROME, Drosophila melanogaster, Q9VP20; CHICKEN, Gallus gallus, Q9PTI4, F1NI93; GORGO, Gorilla gorilla, G3S6Y3, G3QDU4, G3RZ99; HUMAN, 1011 1012 Homo sapiens, Q9UHC7, Q9H000, Q13064; IXOSC, Ixodes scapularis, B7QIJ9, 1013 B7Q4B2; LEPOC, Lepisosteus oculatus, W5NGW8, W5N9B2, W5LWJ1; MONDO, 1014 Monodelphis domestica, F6QPR3, F7F0I3; MOUSE, Mus musculus, Q9QXP6, 1015 Q9ERV1, Q60764; ORYLA, Oryzias latipes, H2MBR3, H2M1P4, H2LQG1; PANTR, 1016 Pan troglodytes, H2QVH8, H2QM29, H2Q915; RAT, Rattus norvegicus, 1017 A0A0G2QC40, Q5XI23, D3ZY41; XENTR, Xenopus tropicalis, Q6GLD9, B4F720. (B) 1018 The PAM2 motif in Makorin proteins from vertebrates (bottom, species abbreviations 1019 as in (A)) shows similarities to PAM2 in known PABP-interacting proteins from human (top, protein names given; first amino acid position for all PAM2 motifs indicated on the 1020 left in grey). The PAM2 consensus (Kozlov et al. 2001) is given above. Positions 9, 10 1021 1022 and 12 within the aligned regions that are highly consistent between all aligned proteins 1023 and important for PAM2 function (Kozlov et al. 2004) are highlighted in brown. 1024 Mutations that were introduced to abrogate the function of the PAM2 motif in human 1025 MKRN1 (MKRN1^{PAM2mut}) are shown below. The corresponding UniProt identifiers are 1026 Q8IYD1, Q8NDV7, Q99700, Q9H074, Q9BPZ3 (known PABP-interacting proteins 1027 from human), Q9UHC7, H2QVH8, G3S6Y3, J9P921, E2RRA5, F1MF12, Q5XI23, 1028 Q9QXP6, Q9PTI4, F6QPR3, W5NGW8, Q4VBT5, H2MBR3 (Makorin orthologs from 1029 vertebrates). (C) The closest Makorin orthologs in Saccharomyces cerevisiae lack RING domain and PAM2 motif. Domain architecture of Yth1p and Lee1p, which were 1030 1031 detected as closest orthologs by HaMStR-OneSeq (Ebersberger et al. 2014), but were 1032 not considered as orthologs due to low FAS scores (0,59 and 0,60, respectively). The 1033 annotated PFAM domains are CCCH zinc finger, PF15663, PF00642, PF16131.

1034

1035 Supplemental Figure S2. MKRN1 interacts with translational regulators and other RBPs. (A) Overlap of the 53 significant interaction partners of GFP-MKRN1^{wt} in human 1036 HEK293T cells with previously published interactors of MKRN1 in mouse embryonic 1037 1038 stem cells (mESC) (Cassar et al. 2015). (B) GO terms enriched for the 53 MKRN1 1039 interactors. P values (modified Fisher exact test, Benjamini-Hochberg correction) are depicted for all significant GO terms (corrected *P* value < 0.05) for Biological Process, 1040 1041 Molecular Function and Cellular Compartment, together with the number of interactors 1042 associated with the respective term. (C) Reciprocal APs show that MKRN1 interacts 1043 with PABPC1, ELAVL1 and IGF2BP1 independently of RNA. AP with GFP-PABPC1, 1044 GFP-ELAVL1 and GFP-IGF2BP1 as baits were performed from HEK293T cells in the presence or absence of RNase A and T1. Bait proteins and endogenous MKRN1 were 1045 detected by Western blots (replicate 1). Different exposure times (exp.) for MKRN1 are 1046 1047 shown for GFP-ELAVL1 and GFP-IGF2BP1 APs. Quantifications (fold changes of the 1048 MKRN1 signal over empty vector (EV)) of three replicates are shown on the right. 1049 Replicates 2 and 3, and uncropped gel images are shown in Supplemental Fig. S9D-1050 F.

1051

1052 Supplemental Figure S3. Signal-over-background transformation allows to estimate 1053 MKRN1 binding strength. (A-C) Raw iCLIP signal before signal-over-background 1054 transformation. (A) Scatter plots show pairwise comparisons of crosslink events per 1055 binding site in three replicate MKRN1 iCLIP experiments. Pearson correlation 1056 coefficients (r) and associated P values are given. (B) Density plots depict the 1057 distribution of crosslink events per binding site in the three replicate experiments. 1058 Shades of blue indicate 20% quantiles; top 20% of binding sites with highest counts 1059 are denoted by a dashed line. (C) Raw iCLIP counts are strongly influenced by the

1060 expression level of the underlying gene. MKRN1-bound genes were stratified into 50 1061 bins with increasing expression (using the total number of MKRN1 crosslink events within the 3' UTR as a proxy of a gene's expression level). Shown is the average 1062 1063 number of crosslink events per binding site for all binding sites in each bin. Dashed 1064 line denotes median across all bins. (D-F) Signal-over-background (SOB) values allow 1065 to correct for expression-level differences. (D) Pairwise comparison of SOB values for 1066 the three MKRN1 iCLIP replicate experiments. Scatter plots as in (A). (E) Distribution 1067 of SOB values in the three replicates. Density plots as in (B). Shades of blue indicate 1068 20% quantiles. Dashed lines denote the top 20% MKRN1 binding sites with strongest 1069 binding that were used for the analyses in Fig. 2B and Supplemental Fig. S4A. (F) 1070 SOB values are independent of the expression level of the underlying gene. Average 1071 SOB values for all binding sites in each expression bin are shown as in (C).

1072

1073 Supplemental Figure S4. MKRN1 binds upstream of long A-stretches. (A) Binding sites with associated A-stretches show stronger MKRN1 binding. Boxplot compares 1074 1075 the SOB values of MKRN1 binding sites in 3' UTRs with and without associated A-1076 stretches. Number of binding sites indicated inside box. (B) Heatmap representation 1077 of 1,412 non-overlapping A-stretches at MKRN1 binding sites, sorted by increasing 1078 length (8-30 nt). Only A's are coloured. (C) MKRN1 binding site strength (signal-over-1079 background, SOB) increases with length of associated A-stretch. Mean and standard 1080 deviation of MKRN1 binding sites associated with A-stretches of increasing length (x-1081 axis). MKRN1 binding sites without associated A-stretches are shown for comparison 1082 on the left. Number of binding sites in each category indicated as barchart above. (D) 1083 The top 20% MKRN1 binding sites show a strong RNA binding preference for AAAA. 1084 Scatter plot compares the frequency of 4-mers within the 9-nt MKRN1 binding sites

and flanking 40-nt windows for the top 20% and bottom 20% MKRN1 binding sites (according to SOB). 4-mer frequencies are displayed as z-scores based on background distribution from binding site permutations.

1088

1089 **Supplemental Figure S5.** Interaction with PABP is required for MKRN1 RNA binding. 1090 (A,B) UV crosslinking experiments to measure the RNA binding capacity of GFP-MKRN1^{wt} and GFP-MKRN1^{PAM2mut}. Autoradiographs (top) and Western blots (bottom) 1091 1092 show GFP-MKRN1/RNA complexes and GFP-MKRN1 protein, respectively, in the eluates from replicates 2 (with 4SU and UV crosslinking at 365 nm) (A) and 3 (with 1093 1094 conventional UV crosslinking at 254 nm) (B). For calibration, input samples for GFP-1095 MKRN1^{wt} were diluted to 75%, 50% and 25% prior to GFP AP. Note that samples were 1096 loaded in different order in (B). Quantifications are given below. Uncropped gel images 1097 are shown in Supplemental Fig. S10.

1098

1099 Supplemental Figure S6. MKRN1 is required to stall ribosomes at K(AAA)₂₀ in 1100 reporter assays. (A) Translation of dual fluorescence reporter plasmids was assessed 1101 by flow cytometry upon MKRN1 and/or ZNF598 KD. Median RFP:GFP ratios (normalised to K₀ in control KD) are shown for the reporter plasmids K₀, K(AAA)₁₂, 1102 1103 K(AAA)₂₀, and R(CGA)₁₀. Error bars represent standard deviation of the mean (s.d.m., 1104 $n \ge 6$ replicates; paired two-tailed t-test, Benjamini-Hochberg correction). Density plot 1105 of median RFP:GFP ratios of one replicate experiment with K(AAA)₂₀ with control or 1106 *MKRN1* KD (two independent siRNAs, KD1 and KD2) or *ZNF598* is shown on the right. 1107 (B) KDs of MKRN1 and ZNF598 were assessed by Western blot (n = 3 replicates). 1108 Black arrowhead indicates ZNF598. Replicates 2 and 3, and uncropped gel images 1109 are shown in Supplemental Fig. S11A,B. (C) MKRN1 KD2 also reduces MKRN2

1110 levels. MKRN1 KD1 and KD2 were performed for 72 h. Expression levels of MKRN1 and MKRN2 were assessed in relation to ß-actin levels by gPCR in MKRN1 KD (siRNA 1111 1112 1 and 2) and control KD. Error bars indicate s.d.m. (n = 2 replicates). (D,E) Cross-1113 regulation of MKRN1 and ZNF598. (D) MKRN1 KD1 reduces endogenous ZNF598 1114 protein levels. Effect of MKRN1 KD (KD1, siRNA 1 and KD2, siRNA 2) and ZNF598 KD for 72 h was assessed by Western blot for endogenous MKRN1 and ZNF598. 1115 Quantifications depict MKRN1 or ZNF598 expression levels in MKRN1 or ZNF598 KD 1116 1117 over control KD condition, normalised to tubulin levels (n = 3 replicates). Replicates 2 1118 and 3, and uncropped gel images are shown in Supplemental Fig. S11C,D. (E) 1119 ZNF598 overexpression reduces MKRN1 protein levels. Effect of ZNF598 and MKRN1 (wt and mutants) overexpression was tested after 48 h. Quantification as in (D). 1120 Uncropped gel images for all replicates are shown in Supplemental Fig. S11E,F. 1121

1122

Supplemental Figure S7. Interactome of GFP-MKRN1^{RINGmut} reveals putative 1123 ubiguitylation substrates. Experiments were performed using SILAC-based MS. 1124 Asymmetrical z-scores of combined SILAC ratios (n = 3 replicates) are shown. Proteins 1125 1126 are detected in at least two out of three replicates. (A) Protein interactome of GFP-1127 MKRN1^{RINGmut} in HEK293T cells analysed by quantitative mass spectrometry. Combined SILAC ratios (n = 3 replicates) after z-score normalisation are plotted 1128 1129 against log₁₀-transformed intensities. 1,097 protein groups were quantified in at least 1130 two out of three replicates (Supplemental Table S1). MKRN1 and interesting 1131 candidate ubiquitylation targets are highlighted. (B) Quantitative comparison of the 1132 interactome of GFP-MKRN1^{wt} and GFP-MKRN1^{RINGmut} shows that potential ubiquitylation candidates identified in (A) are enriched in GFP-MKRN1^{RINGmut} over 1133 GFP-MKRN1^{wt}. Comparison reveals 137 proteins to be significantly enriched 1134

1135 (MKRN1^{RINGmut} over MKRN1^{wt} with FDR < 5% and MKRN1^{wt}/GFP z-score > 1). 1136 Combined ratios of three replicates are shown in a scatter plot. Only proteins detected 1137 in at least two out of three replicates are shown. Highlighting as in (*A*).

1138

1139 Supplemental Figure S8. GO term analysis of MKRN1 ubiquitylation targets and proteome analysis upon MKRN1 KD. (A) GO terms enriched for the 21 MKRN1 1140 1141 ubiguitylation targets. Corrected P values (modified Fisher exact test, Benjamini-1142 Hochberg correction) are depicted for all significant GO terms (corrected P value < 0.05) for Biological Process (BP), Molecular Function (MF) and Cellular Compartment 1143 1144 (CC), together with the number of ubiguitylation targets associated with the respective 1145 term. (B) Proteome analysis of MKRN1 KD in HEK293T cells analysed by quantitative mass spectrometry. Log₂-transformed, combined normalised SILAC ratios (n = 31146 replicates) are plotted against log₁₀-transformed intensities. 6,425 protein groups were 1147 quantified in at least one out of three replicate experiments (Supplemental Table S4). 1148 1149 Selected ubiquitylation targets of MKRN1 are highlighted.

1150

1151 Supplemental Figure S9. Images of full membranes and different exposure times (exp.) for Western blot analyses in Fig. 1C and Supplemental Fig. S2C in the 1152 1153 presence or absence of RNase A and T1. (A-C) PABP interacts with MKRN1^{wt} and 1154 MKRN1^{RINGmut} but not MKRN1^{PAM2mut}. Western blot analysis was performed with 1155 antibodies against PABPC1/3 and GFP. Images of full membranes and different 1156 exposure (exp.) times for both antibodies are shown for replicate 1 (A) which is 1157 presented in Fig. 1C, as well as replicates 2 (*B*) and 3 (*C*). Black and blue arrowheads indicate GFP-MKRN1 and PABPC1/3, respectively. (D-F) Endogenous MKRN1 1158 1159 interacts with GFP-PABPC1 independent of RNA. Western blot analysis was

1160 performed with antibodies against MKRN1 and GFP. Images of full membranes and 1161 different exposure times for both antibodies are shown for replicate 1 (D) which is 1162 presented in Supplemental Fig. S2C, as well as replicates 2 (E) and 3 (F). Black and blue arrowheads indicate MKRN1 and GFP-PABPC1, replicates. (G-I) Endogenous 1163 1164 MKRN1 interacts with GFP-ELAVL1 independent of RNA. Western blot analysis was performed with antibodies against MKRN1 and GFP. Images of full membranes and 1165 different exposure times for both antibodies are shown for replicate 1 (G) which is 1166 1167 presented in Supplemental Fig. S2C, as well as replicates 2 (H) and 3 (I). (J-L) 1168 Endogenous MKRN1 interacts with GFP-IGF2BP1 independent of RNA. Western blot 1169 analysis was performed with antibodies against MKRN1 and GFP. Images of full 1170 membranes and different exposure times for both antibodies are shown for replicate 1 (J) which is presented in Supplemental Fig. S2C, as well as replicates 2 (K) and 3 1171 1172 (*L*).

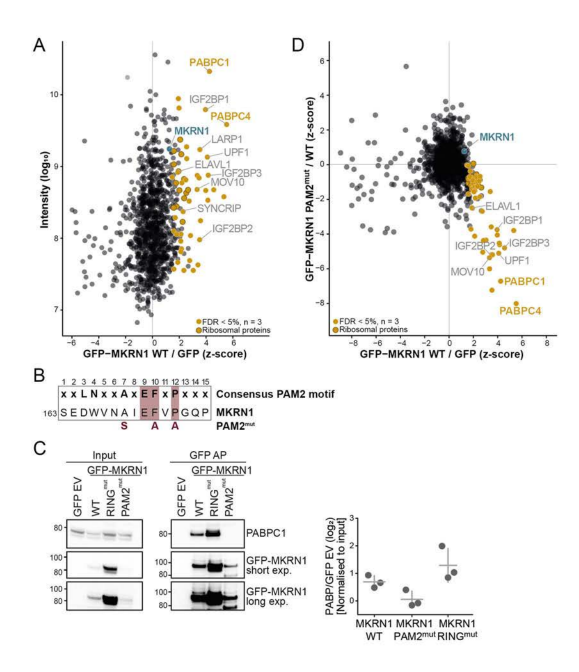
1173

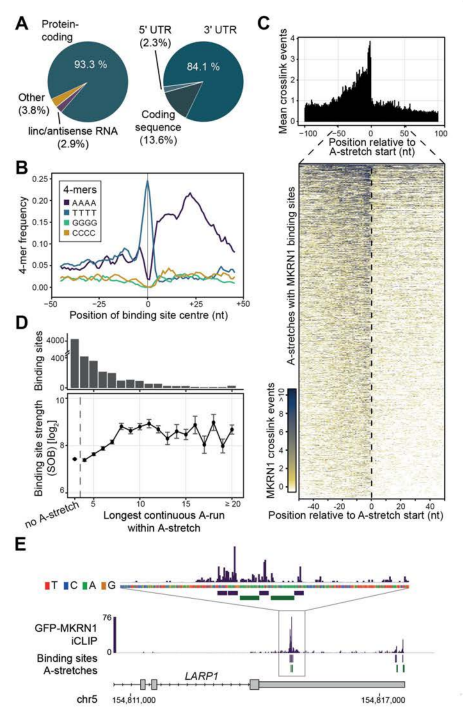
1174 Supplemental Figure S10. Images of full membranes of autoradiographs and Western blot analyses in Fig. 3D (replicate 1) and Supplemental Fig. S5 (replicates 1175 1176 2 and 3). UV crosslinking experiments to measure the RNA binding capacity of GFP-1177 MKRN1^{wt} and GFP-MKRN1^{PAM2mut}. Autoradiographs (A, left; B, top) and Western blots 1178 (A, right; B, bottom) show GFP-MKRN1/RNA complexes and GFP-MKRN1 protein. 1179 respectively, in the eluates from replicates 1 and 2 (with 4SU and UV crosslinking at 1180 365 nm) (A) and 3 (with conventional UV crosslinking at 254 nm) (B). (B) Images of full 1181 membranes of Western blot analyses with both antibodies are shown for replicate 3 1182 (*B*).

1183

1184 **Supplemental Figure S11.** Images of full membranes and different exposure (exp.) 1185 times for Western blot analyses in **Supplemental Fig. S6B,D,E.** (A,B) KDs of MKRN1 1186 and *ZNF598* assessed by Western blot (n = 3 replicates) from **Supplemental Fig.** 1187 **S6B**. Western blot analysis was performed with antibodies against MKRN1, ZNF598. 1188 and tubulin. Black and blue arrowheads indicate MKRN1 (53 kDa) and ZNF598 (99 1189 kDa), respectively. Uncropped gel images of replicates 1 and 2 (A) and 3 (B). (C,D) Images of full membranes are shown for cross-regulation between MKRN1 and 1190 1191 ZNF598 KD from Supplemental Fig. S6D. MKRN1 KD1 reduces endogenous ZNF598 1192 protein levels. Western blot analysis was performed with antibodies against MKRN1, 1193 ZNF598, and tubulin. Coloured arrowheads as in (A). Uncropped gel images of 1194 replicate 1 (C) and replicates 2 and 3 (D). (E,F) Images of full membranes are shown for cross-regulation of MKRN1 and ZNF598 overexpression (OE) from Supplemental 1195 1196 Fig. S6E. ZNF598 OE reduces MKRN1 protein levels. Western blot analysis was 1197 performed with antibodies against MKRN1, ZNF598, and tubulin. Black arrowheads 1198 indicate MKRN1. Images of full membranes and different exposure times (exp.) for 1199 both antibodies are shown for replicates 1, 2 (E), and 3 (F). Note the opposite order of 1200 replicates 1 and 2 (2 left, 1 right) in (E).

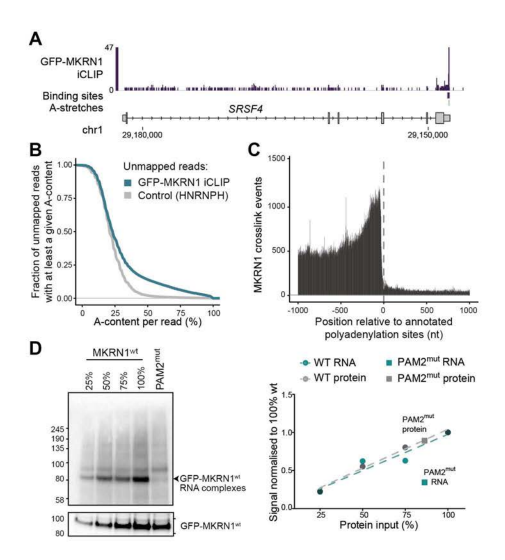
Hildebrandt Fig1



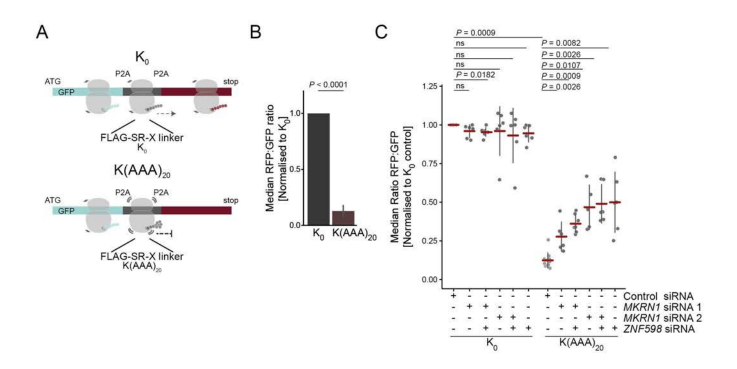


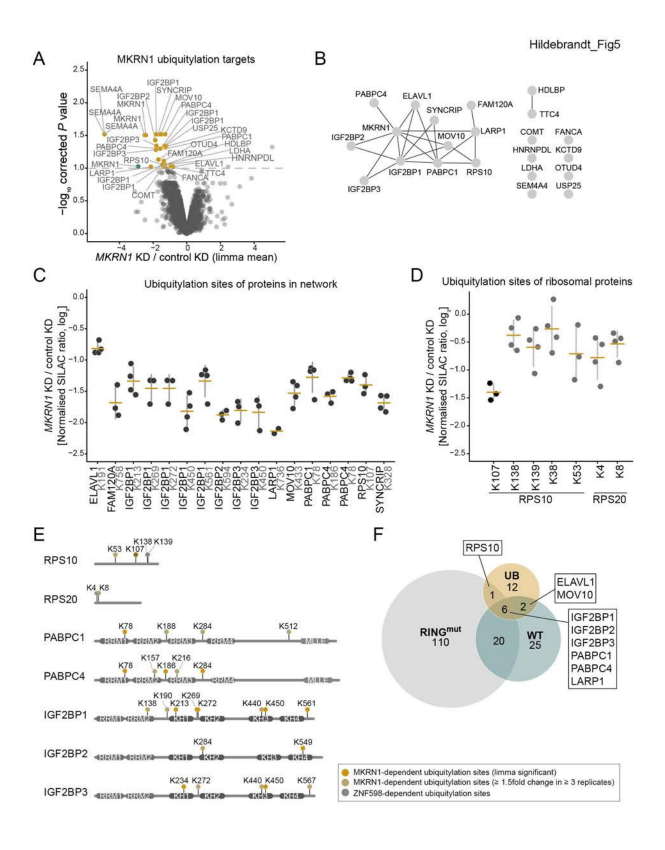
Hildebrandt_Fig2

Hildebrandt Fig3

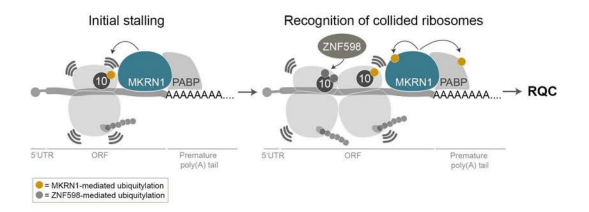


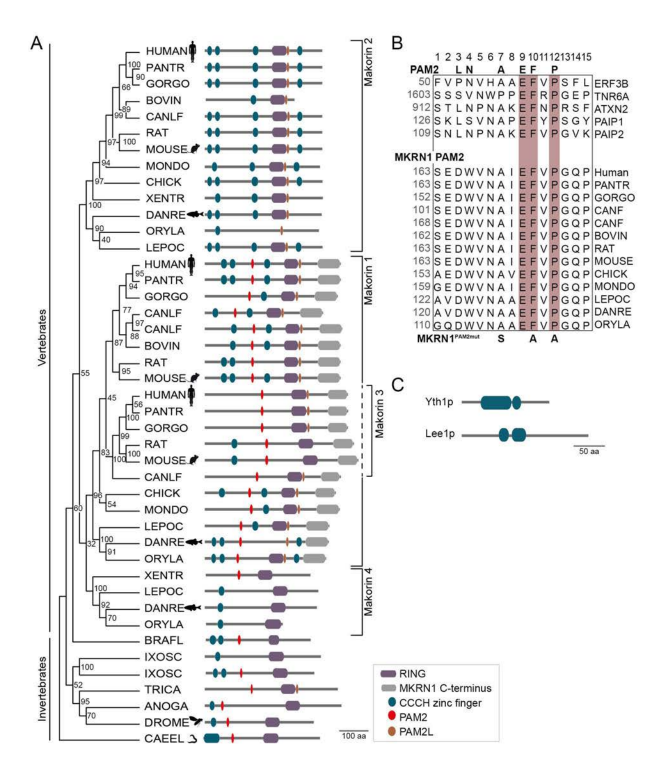
Hildebrandt_Fig4

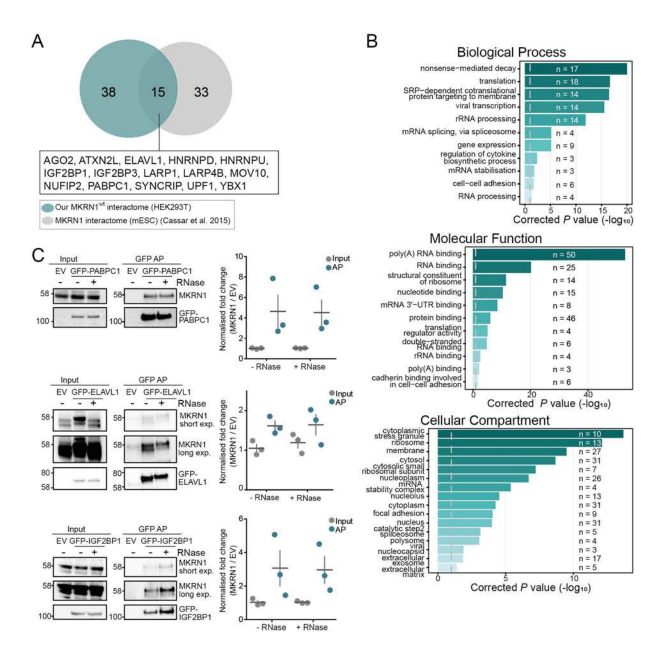


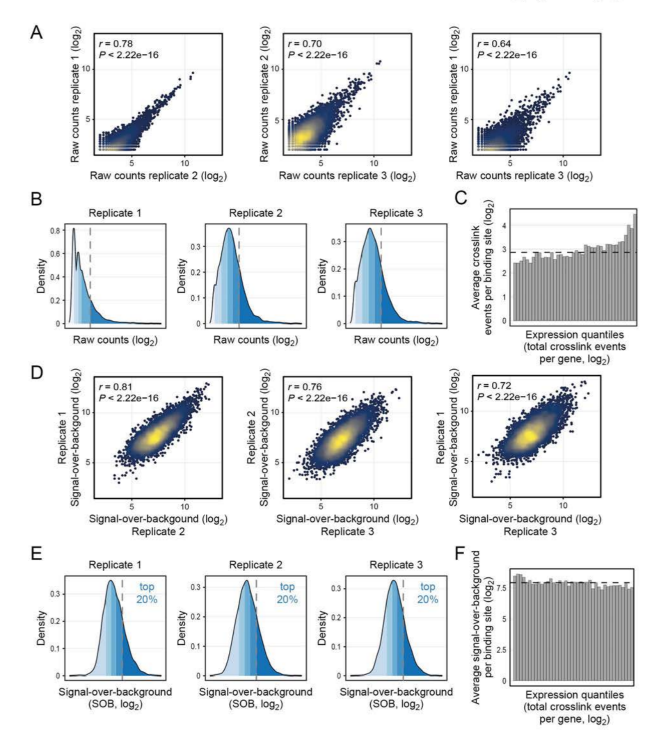


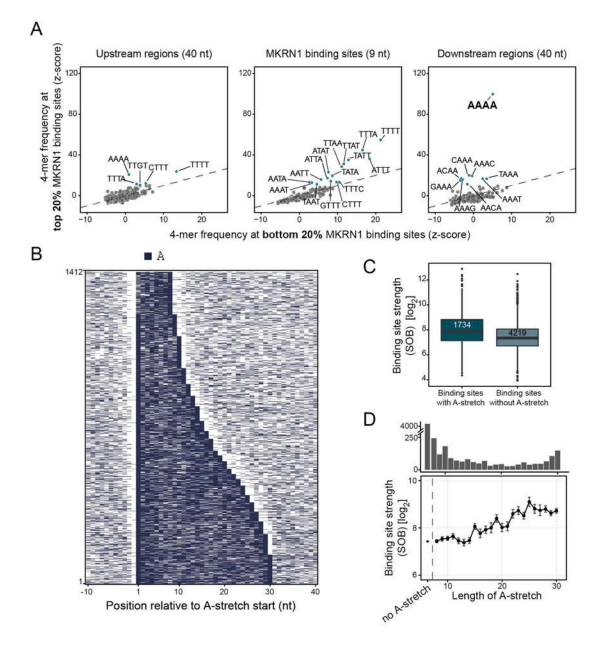
Hildebrandt_Fig6

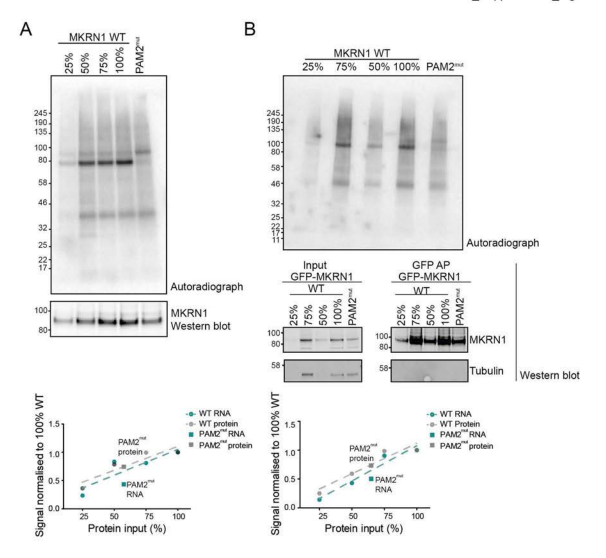


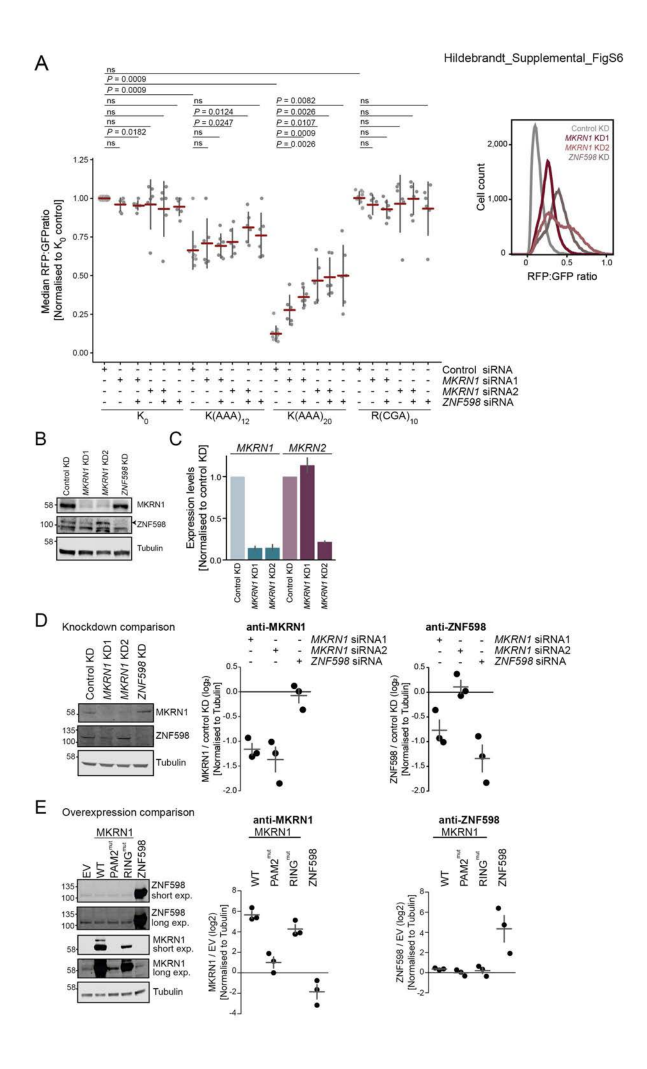




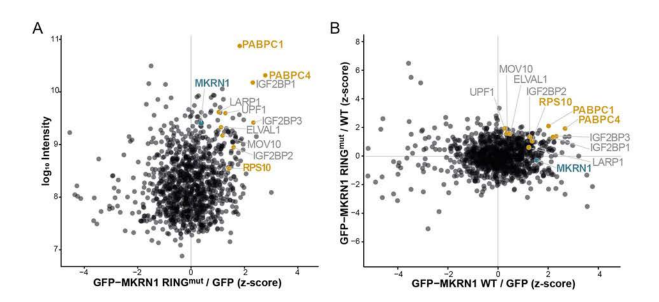


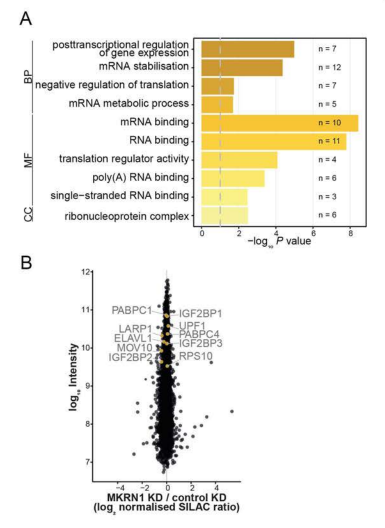


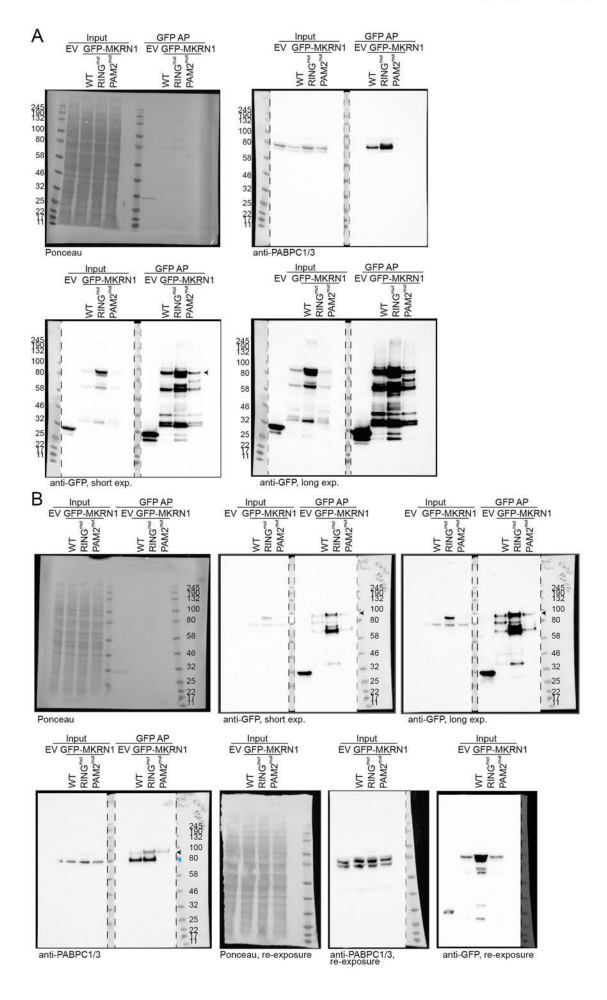


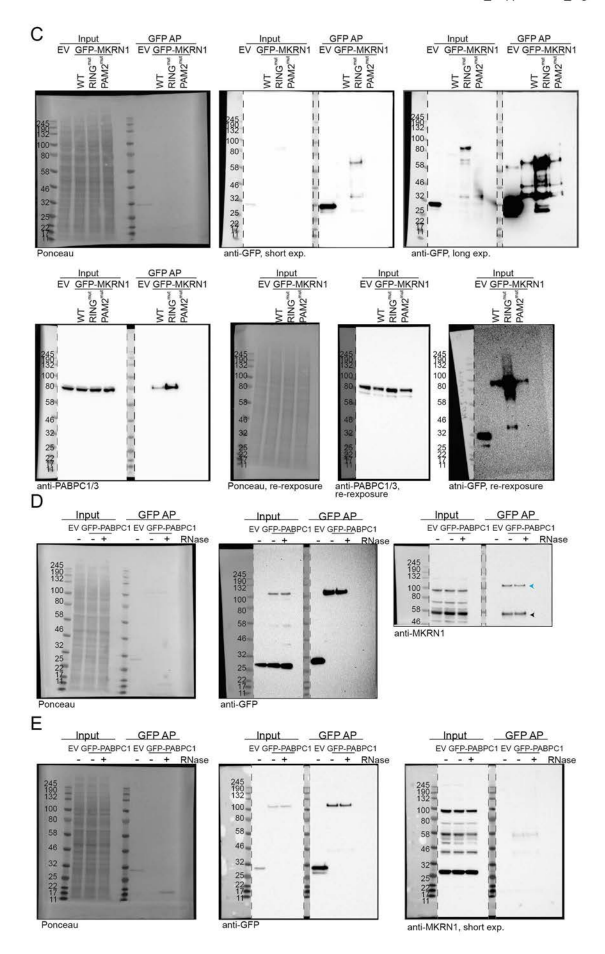


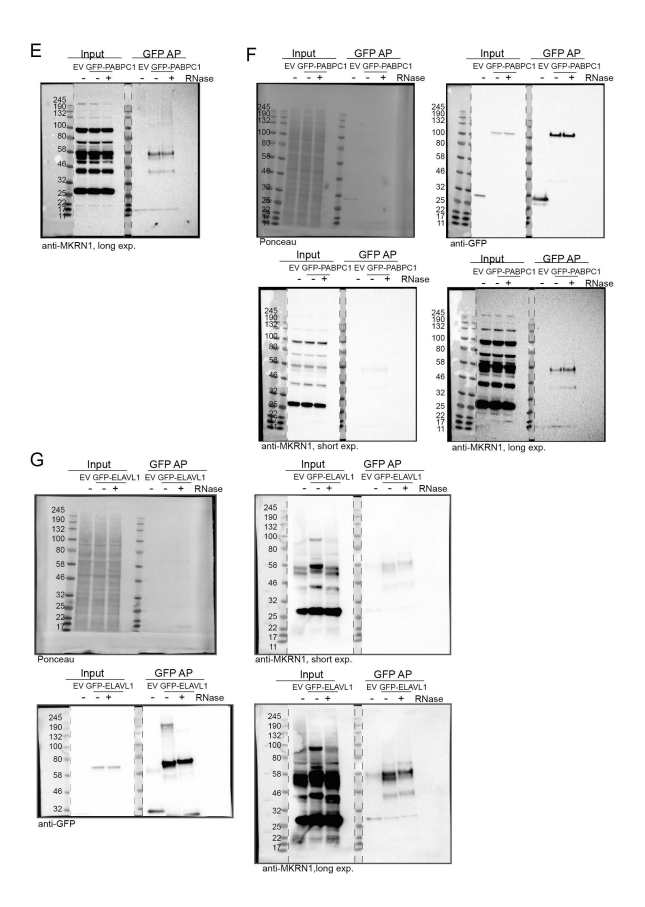
Hildebrandt_Supplemental_FigS7

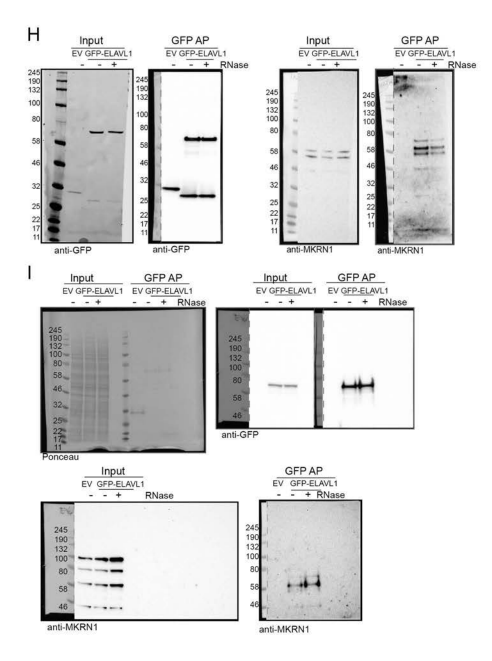


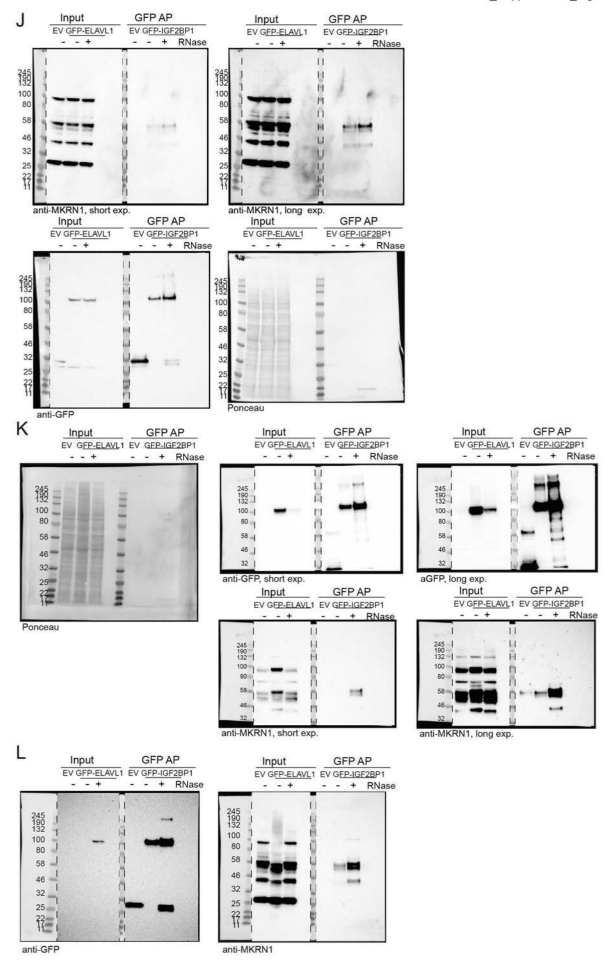




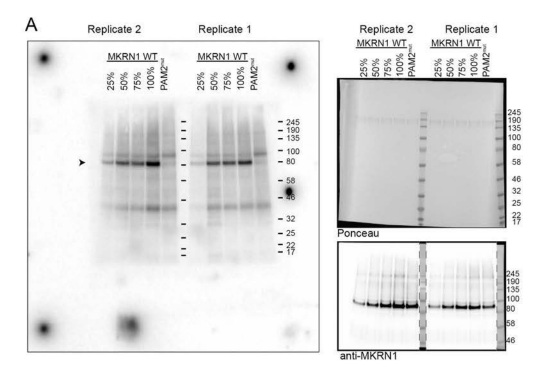








Hildebrandt_Supplemental_FigS10



В

