# *EIF1AX* and *RAS* mutations cooperate to drive thyroid tumorigenesis through ATF4 and c-MYC

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- 5 Gnana P. Krishnamoorthy<sup>1</sup>, Natalie R. Davidson<sup>2</sup>, Steven D. Leach<sup>1</sup>, Zhen Zhao<sup>3</sup>, Scott
- W. Lowe<sup>3</sup>, Gina Lee<sup>6</sup>, Iňigo Landa<sup>1</sup>, James Nagarajah<sup>1</sup>, Mahesh Saqcena<sup>1</sup>, Kamini
- <sup>7</sup> Singh<sup>3</sup>, Hans-Guido Wendel<sup>3</sup>, Snjezana Dogan<sup>5</sup>, Prasanna P. Tamarapu<sup>1</sup>, John Blenis<sup>6</sup>,
- 8 Ronald A. Ghossein<sup>5</sup>, Jeffrey A. Knauf<sup>1,4</sup>, Gunnar Rätsch<sup>2</sup>, James A. Fagin<sup>1,4\*</sup>.
- <sup>9</sup> <sup>1</sup>Human Oncology and Pathogenesis Program, <sup>2</sup>Computational Biology Program, <sup>3</sup>Cancer
- <sup>10</sup>Biology and Genetics Program, <sup>4</sup>Department of Medicine, <sup>5</sup>Department of Pathology, Memorial
- <sup>11</sup> Sloan Kettering Cancer Center, New York, NY, U.S.A. and the <sup>6</sup>Department of Pharmacology,
- 12 Meyer Cancer Center, Weill Cornell Medicine, New York, NY, U.S.A.
- 13 Running title: EIF1AX and RAS cooperate to generate mTOR kinase dependency.

#### 14 Correspondence:

- 15 James A. Fagin, MD
- 16 Department of Medicine and Human Oncology and Pathogenesis Program,
- 17 Memorial Sloan-Kettering Cancer Center, 1275 York Avenue, New York, NY 10065, U.S.A.
- 18 Email: faginj@mskcc.org
- 19
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#### 21 Abstract

22

Translation initiation is orchestrated by the cap binding and 43S pre-initiation complexes (PIC).
 Eukaryotic initiation factor 1A (EIF1A) is essential for recruitment of the ternary complex and

- for assembling the 43S PIC. Recurrent *EIF1AX* mutations in papillary thyroid cancers are
- 26 mutually exclusive with other drivers, including RAS. *EIF1AX* is enriched in advanced thyroid
- cancers, where it displays a striking co-occurrence with RAS, which cooperates to induce
- tumorigenesis in mice and isogenic cell lines. The C-terminal *EIF1AX-A113splice* mutation is
- the most prevalent in advanced thyroid cancer. EIF1AX-A113spl variants stabilize the PIC and
- induce ATF4, a sensor of cellular stress, which is co-opted to suppress EIF2α phosphorylation,
- enabling a general increase in protein synthesis. RAS stabilizes c-MYC, an effect augmented
- by EIF1AX-A113spl. ATF4 and c-MYC induce expression of aminoacid transporters and
- enhance sensitivity of mTOR to aminoacid supply. These mutually reinforcing events generate
- therapeutic vulnerabilities to MEK, BRD4 and mTOR kinase inhibitors.

#### 35 Significance

- 36 Mutations of *EIF1AX, a* component of the translation preinitiation complex, co-occur
- with *RAS* in advanced thyroid cancers and promote tumorigenesis. *EIF1AX*-
- A113splice drives an ATF4-induced dephosphorylation of EIF2α, resulting in increased
- 39 protein synthesis. ATF4 also cooperates with c-MYC to sensitize mTOR to aminoacid
- 40 supply, thus generating vulnerability to mTOR kinase inhibitors.

#### 42 Introduction

Papillary carcinomas (PTC) are the most common type of thyroid cancer. They are 43 usually indolent tumors that harbor mutually exclusive mutations in BRAF, RAS or 44 fusions of RET, NTRK or BRAF (1-3). The TCGA study of PTC identified additional 45 driver alterations present at lower frequency, including EIF1AX, PPM1D and CHEK2 46 (4). Poorly differentiated (PDTC) and anaplastic thyroid cancers (ATC) are the most 47 48 aggressive forms of the disease and are characterized by distinct genomic profiles. Although BRAF and RAS mutations are also the main drivers, as compared to PTC they 49 are more frequently associated with mutations of the TERT promoter, TP53, genes 50 encoding PI3K/AKT/mTOR pathway effectors or chromatin modifiers (5-7). They are 51 52 also markedly enriched for *EIF1AX* mutations.

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54 Translation initiation in higher eukaryotes is orchestrated by the tight regulation of the cap binding and the 43S pre-initiation complexes (PIC). Formation of the PIC involves 55 56 recruitment of the ternary complex (EIF2-GTP-tRNAi(Met)) onto the 40S ribosomal subunit. The PIC component EIF1A, which is encoded on human chromosomes X and 57 Y by *EIF1AX* and *EIF1AY*, respectively. Their protein products are highly conserved, 58 and expression of EIF1A is biallelic irrespective of gender (8.9). EIF1A is required for 59 recruitment of the ternary complex and for assembling the 43S PIC (10), which after 60 recruitment onto capped mRNAs scans their 5'UTR and localizes the AUG to initiate 61 translation (11,12). Deregulation of translation initiation is common in tumorigenesis. 62 Increased expression of components of the EIF4F cap binding complex (EIF4E, EIF4A, 63 *EIF4G*) is seen in many cancers. The expression of these genes is under transcriptional 64 control by c-MYC (13,14). EIF4E is limiting for translating the mammalian genome, and 65 is frequently found in excess in cancer cells, where it may help drive a translational 66 output supporting tumorigenesis (15). 67 68

To our knowledge *EIF1AX* is the only example of a PIC subunit recurrently mutated in

cancer. Mutations of *EIF1AX* were first reported in uveal melanomas (8).

Comprehensive genomic profiling of these tumors revealed that *EIF1AX*-mutant tumors

- 72 mark a comparatively low risk form of the disease. EIF1AX and GNA11/GNAQ
  - 3

73 mutations frequently co-occurred and were mutually exclusive with c-MYC amplification

74 (16). EIF1AX mutations have been reported in benign thyroid adenomas (17), follicular

carcinomas (18) as well as in ~1% of PTC in a mutually exclusive manner with other

drivers (4). By contrast, they are present in 11% of PDTC and ATC, and are almost

invariably associated with oncogenic RAS (5,7). The striking co-evolution of EIF1AX

and RAS mutations in advanced disease suggests that they may cooperate to drive

- 79 tumor progression.
- 80

The core RNA binding domain of *EIF1A* is universally conserved from archaea to 81 eukaryotes, whereas eukaryotes differ from bacteria by the addition of the unstructured 82 amino-terminal (NTT) and carboxy-terminal tails (CTT) (19,20). EIF1AX mutations 83 identified in several cancers encode somatic substitutions in the first 2-15 amino acids 84 of the NTT (8,21,22), whereas thyroid cancers harbour an additional hotspot splice site 85 mutation (EIF1AX-A113splice) in the CTT that is private to this disease (4,5,7). EIF1AX 86 mutations are always heterozygous, suggesting that full occupancy of PICs by mutant 87 88 EIF1AX would be detrimental to viability. Structure-function studies in yeast revealed that mutating any of the NTT residues between 7 and 16 amino acid were lethal and 89 90 resulted in leaky scanning of the initiation AUG codon (23,24), and discriminated against AUGs with poor context (25). By contrast, CTT mutants enhanced non-91 92 canonical AUG initiations. Of note, the experimental CTT substitutions tested in yeast did not appropriately model the structural defects of the EIF1AX splice site mutation 93 (A113splice). 94

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96 Here we describe the identification of the key signaling drivers of transformation by

97 *EIF1AX* mutants, particularly *EIF1AX-A113splice*, alone and in the context of RAS, and

98 the therapeutic dependencies they confer.

#### 99 Results

## *EIF1AX* mutation is a strong co-operating event with *RAS* in advanced thyroid cancer; the hotspot A113splice mutation induces aberrant splice variants.

102 Analysis of our institutional clinical genomics database of 148 advanced thyroid cancers coupled to data from two previously published studies (5,7) showed that 26/246 (11%) 103 104 tumors harbored EIF1AX mutations, 25 of which were associated with mutant RAS (25/26; p=3.15x10E<sup>-13</sup>) (Fig. 1A). The EIF1AX mutations clustered within the first 15 105 106 amino acids of the N-terminal tail (NTT), as reported in uveal melanomas (8), or more frequently at a hotspot splice acceptor site upstream of exon 6 (A113splice) in the C-107 108 terminal tail (CTT) (17/26) (Fig. 1A). The A113splice mutation, not seen so far in any other cancer type, abolished the acceptor site of exon 6, resulting in two alternatively 109 spliced transcripts (Fig. 1B): i) c'splice, through usage of a cryptic site within exon 6, 110 vielding a 132AA protein through an in-frame exclusion of 12 AA. ii) t'splice, which 111 retains intron 5, resulting in a 115AA truncated protein. We confirmed the presence of 112 these alternatively spliced mRNA's in the RNAseg data of A113splice-mutant PTCs 113 from the TCGA study (4) (Supplementary Fig. S1A). Western blotting of HTH83 and 114 C643 thyroid cancer cell lines harboring the A113splice mutation showed EIF1AX 115 protein products corresponding to c'spl and t'spl mRNAs, with c'spl as the 116 predominantly expressed isoform (Fig. 1C). Their predicted AA sequences are shown in 117 Supplementary Fig. S1B. 118

### Aberrant splice variants of *EIF1AX-A113splice* mutation induce transformation *in vitro.*

- 121 To explore the biological consequences of the *EIF1AX-A113splice* mutation, either
- alone or in the context of mutant *RAS*, we generated isogenic thyroid cancer cell lines
- 123 by CRISPR-Cas9 knock-in of heterozygous mutations of A113splice into KRAS<sup>G12R</sup>-
- mutant CAL62 or *RAS*-wild type TTA1 cells, and by reversing the endogenous
- 125 A113splice mutation in HRAS<sup>G13R</sup> and EIF1AX<sup>A113spl</sup>-mutant C643 cells (Fig. 2A,
- 126 Supplementary Fig. S2A and S2B). Introduction of *A113splice* into CAL62 or TTA1 cells
- 127 markedly increased their colony formation in soft agar, whereas transformation
  - 5

- 128 efficiency was decreased by editing out *A113splice* in C643 cells (Fig. 2A). We then
- 129 tested the effects of the two *EIF1AX-A113splice* products, c'spl and t'spl, independently
- and in combination on transformation of Nthy-Ori 3-1 cells, an SV40 large-T-antigen
- immortalized human thyroid cell line. The c'spl product markedly increased colony
- formation, whereas the t'spl did not, and dampened the effects of c'spl when both were
- 133 co-expressed through a bicistronic vector, consistent with EIF1AX-c'spl being the
- 134 functionally active variant (Supplementary Fig. S2C).

## EIF1AX-c-splice cooperates with oncogenic *Ras* to induce disease progression in genetically engineered mice.

137 We next investigated the interaction of oncogenic Hras and EIF1AX-c'spl *in vivo*. For

- this we generated mice with thyroid-specific, doxycycline (dox)-inducible expression of
- 139 EIF1AX-c'spl by targeting a *TRE-EIF1AX-c'spl* construct into the mouse *ColA1* locus
- and crossing the resulting animals with *Tg-rtTA* mice (Fig. 2B). Dox-fed *Tg-rtTA/TRE-*
- 141 *EIF1AX-c'spl* mice expressed the c'spl protein in thyroid tissue (Fig. 2C), which
- developed thyrocyte hyperplasia with atypical features (18/19), with one animal
- developing a low-grade classical PTC (Fig. 2D and 2E). These findings closely
- phenocopy the human thyroid histologies associated with an isolated *EIF1AX* mutation
- 145 (4,17). *TPO-Cre/FR-Hras<sup>G12V</sup>* mice express Hras<sup>G12V</sup> upon Cre-induced recombination in
- the thyroid (Fig. 2B), which is insufficient to drive tumorigenesis (26). By contrast, the
- 147 quadri-transgenics (TPO-Cre/FR-Hras<sup>G12V</sup>/Tg-rtTA/TRE-EIF1AX-c'spl) displayed
- neoplasms along the spectrum of disease progression with a penetrance of 30%,
- including Hurthle cell adenoma, PTC and PDTC (Fig.2D and 2E, Supplementary Fig.
- 150 S2D), consistent with the histological characteristics of human thyroid tumors harboring
- the combined genetic lesions (17).

# *EIF1AX* mutants have higher affinity to components of the translation PIC and increase protein synthesis.

- 154 EIF1AX is an essential subunit of the translation PIC (10,12). We performed co-
- immunoprecipitation experiments to probe for possible aberrant interactions of EIF1AX
- mutants with components of the ternary complex (TC) and the PIC. IP of HEK293T cells
  - 6

expressing HA-tagged wild-type (WT) EIF1AX, NTT mutants (G8R, G9R, G15V) or 157 EIF1AX-c'spl with an antibody to HA showed pulldown of the TC component eukaryotic 158 initiation factor  $2\alpha$  (EIF $2\alpha$ ) by all EIF1AX proteins, with EIF1AX-c'spl showing greater 159 affinity (Fig. 3A). IP of the HEK293T lysates with EIF5, a component of the PIC, did not 160 detect EIF1AX-WT in the immunoprecipitate, likely because of the known labile 161 interactions between these PIC subunits (10). By contrast, EIF1AX mutants, particularly 162 G8R, G9R and c'spl, exhibited increased affinity for EIF5 (Fig. 3B), consistent with 163 stabilization of the PIC. This was confirmed in the isogenic EIF1AX-splice cell lines (Fig. 164 3C), and in thyroid cancer cells with endogenous *EIF1AX* mutations (Supplementary 165 Fig. S3A). These data suggest that EIF1AX mutants result in a more stable 43S 166 ribosomal complex. As translation initiation is a rate limiting process, we next tested 167 168 whether the EIF1AX mutants altered nascent protein synthesis in the isogenic lines. Lazidohomoalanine (AHA)-labeled proteins were markedly increased by knock-in of the 169 A113splice mutation into RAS wild type (TTA1) or mutant cells (CAL62), whereas 170 reversion of the mutation in C643-spl-rev cells had the opposite effect (Fig. 3D and 3E). 171 172 NthyOri cells stably expressing EIF1AX-G8R, G9R or EIF1AX-c'spl also showed an increase in nascent protein synthesis compared to WT, with c'spl having the greatest 173 174 effect (Fig. 3D and 3E). The increased protein synthesis by c'spl is comparable to that induced by EIF4E overexpression (Supplementary Fig. S3B) and is blocked by mTOR 175 176 kinase inhibition (Supplementary Fig. S3C).

### 177 Increased global protein synthesis by EIF1AX-splice is mediated by ATF4-

178 induced EIF2α dephosphorylation.

To determine whether the effects on protein synthesis were global or selective, and the candidate mechanisms involved, we performed low pass ribosome footprinting in C643 cells to identify subsets of mRNAs that were translated with greater (TE-high) or lesser (TE-low) efficiency than its isogenic splice reversed control (Supplementary Table S1 and S2). Interestingly *ATF4*, a known translationally regulated gene, scored as a preferentially translated candidate (log<sub>2</sub>FC:0.71; p-val:0.005; p-adj: 0.09) (Supplementary Table S1). Accordingly, polysome profiling by density gradient

186 fractionation showed that ATF4 mRNA was enriched in actively translating polysome

fractions of NthyOri cells expressing EIF1AX-splice (Supplementary Fig. S4A). 187 Additionally, GSEA of RNAseg of CAL62-splice vs. CAL62 cells found an ATF4 188 activation signature (NES:1.7; Nom p-val:0.012) (Supplementary Table S3). ATF4 is a 189 key transcription factor that integrates responses to cell stressors, such as amino acid 190 deficiency or ER protein folding defects. Despite the increase in ATF4, expression of the 191 EIF1AX-splice variants was not associated with significant activation of the ER stress 192 pathway (Supplementary Fig. S4B). ATF4 mRNA contains two upstream open reading 193 frames (uORF) that determine the efficiency of its translation (27). The second uORF 194 (uORF2) overlaps with the canonical ATF4 start codon and is a strong inhibitor of ATF4 195 translation. Under normal conditions, translation starts at uORF1, and the ribosome 196 dissociates at the stop codon. It reassembles at uORF2, which prevents ATF4 197 198 translation. During cellular stress, EIF2 $\alpha$  is phosphorylated at serine 51 by stresssensing kinases. As a result, it remains GDP-bound, dampening formation of the TC 199 (EIF2α -GTP-Met-tRNA). When TC availability is limited, the ribosome fails to 200 reassemble at uORF2, and instead reinitiates translation at the canonical ATF4 start 201 202 codon (28). As EIF1AX is known to impact the fidelity of start codon selection (25,29), we tested whether EIF1AX-splice preferentially translates ATF4 by altering selectivity 203 towards the two upstream (<sup>-3</sup>ACCAUG/<sup>-3</sup>GCCAUG) and the main (<sup>-3</sup>AACAUG) ATF4 204 start codons. For this we engineered reporter constructs in which the firefly luciferase 205 206 protein was under control of the different ATF4 translation initiation contexts (Kozak + start codons). Expression of EIF1AX-WT or EIF1AX-c'spl in HEK293T cells co-207 transfected with appropriate reporters showed that the inhibitory uORF2 led to less 208 efficient translation in EIF1AX mutant-expressing cells (Supplementary Fig. S4C). This 209 210 would conceivably de-repress translation initiation at the ATF4 mORF. Interestingly, reporter activity under control of the ATF4 mORF was also markedly increased in 211 EIF1AX-c'spl cells (Supplementary Fig. S4C). As a complementary strategy, we used a 212 Translation Control Reporter System (TCRS) (30) to test whether increased efficiency of 213 ribosomal re-initiation might explain the effects of the mutant EIF1AX on ATF4 214 translation. Similar to ATF4, the TCRS construct has 3 ORF's: a short uORF, followed 215 by 2 overlapping ORFs encoding a long peptide (L.P) and a short peptide (S.P), 216 respectively, the latter serving as a marker for ribosome re-initiation (Supplementary 217

- Fig. 4D). Expression of EIF1AX NTT or c'spl mutants in HEK293T cells co-transfected
- with TCRS showed higher S.P. levels as compared to EIF1AX-WT, indicative of higher
- 220 efficiency of ribosome re-initiation. ATF4 is believed to feed forward to induce
- expression of its own transcript (31). Consistent with this, ATF4 mRNA was induced by
- ~ 3-fold in all EIF1AX-splice isogenic contexts (Supplementary Fig. S4E). The
- 223 coordinate robust increase of ATF4 gene expression may have dampened the
- sensitivity of the ribosome profiling experiments.
- 225 Serine 51 phosphorylation of EIF2α in response to cellular stress represses global
- translation, but increases translational efficiency of ATF4 (27). Therefore, under
- 227 physiological conditions ATF4 is downstream of p-EIF2α. By contrast, EIF2α in EIF1AX-
- splice mutant cells is paradoxically underphosphorylated. Hence, EIF1AX-splice co-opts
- this pathway by constitutively activating expression of ATF4, placing it upstream of
- $EIF2\alpha$ , increasing availability of the ternary complex and de-repressing global
- translation (Fig. 4A). This is consistent with ATF4 dephosphorylating pS51-EIF2 $\alpha$
- through a negative feedback loop engaged via ATF4-dependent upregulation of
- GADD34 (Fig. 4B), a EIF2α specific cofactor for protein phosphatase-1 (PP1) (32).
- 234 Moreover, the GADD34/PP1 phosphatase inhibitor salubrinal blocked EIF2α
- 235 dephosphorylation and preferentially repressed global protein synthesis in EIF1AX-
- splice vs splice-reverted C643 cells (Fig. 4C).

#### 237 EIF1AX activates mTOR through aberrant expression of ATF4 and c-MYC

- 238 We performed Gene Set Enrichment Analysis (GSEA) to identify the oncogenic
- signaling pathways activated by EIF1AX-splice. In addition to ATF4, the top-ranked
- signatures enriched in EIF1AX-mutant cells in the RNAseq profiles of the isogenic
- 241 CAL62 and C643 models included the following terms: translation, ternary and 43S
- complex formation, mTORC1 signaling and transcriptional targets of c-MYC (Fig. 5A;
- Supplementary Table S3 and S4). Accordingly, the mTOR substrates P70-S6 kinase
- and 4EBP1 were activated by expression of the EIF1AX-splice products in both RAS-
- 245 WT and RAS mutant cell lines (Fig. 5B). The EIF1AX mutant induction of mTOR
- signaling was not associated with PI3K pathway activation in RAS-WT cells, whereas
- 247 AKT and PRAS40 phosphorylation were increased in RAS-mutant cells. Despite this,

the activation of mTOR by the aberrant EIF1AX gene products was neither PI3K- nor

249 RSK-dependent, as treatment with the pan-PI3K inhibitor GDC0941, the pan-AKT

kinase inhibitor MK-2206 or the pan-RSK inhibitor LJI308 did not impair the induction of

- p4EBP1 in the parental C643 compared to splice-reverted cells (Supplementary Fig.
- 252 S5A and S5B).

253 Based on these findings, we hypothesized that c-MYC and ATF4 could be key

oncogenic clients of EIF1AX. Accordingly, c-MYC and ATF4 protein levels were higher

in the isogenic lines and in NthyOri cells expressing the EIF1AX splice variants (Fig.

4B), as well as in mice with dox-inducible thyroid-specific expression of EIF1AX-c'spl

257 (Fig. 5C). The increase in ATF4 and c-MYC was associated with greater abundance of

aminoacid (AA) transporters for glutamine (ASCT2) and leucine (LAT1), which are

known to be regulated by these transcription factors (33,34) (Fig. 4B). The differential

260 expression of these transporters in RAS mutant thyroid cancer cell lines with

261 endogenous *EIF1AX* mutations compared to those that were *EIF1AX* WT was

262 particularly striking (Supplementary Fig. S5C).

We next explored whether ATF4 and/or c-MYC accounted for the increased expression 263 of the aminoacid transporters in cells expressing EIF1AX-splice. Silencing of ATF4 or c-264 MYC alone modestly decreased ASCT2 abundance in C643 cells, with minimal effects 265 on LAT1. However combined ATF4/c-MYC knockdown repressed both transporters, 266 and decreased p4EBP1 (Fig. 5D). This was also the case in NthyOri-splice cells (Fig. 267 5E). The increased expression of ASCT2 and LAT1 in EIF1AX-splice expressing cells 268 could induce mTOR activation through increased influx of glutamine and leucine 269 (34.35). Consistent with this, addition of leucine and glutamine in combination after 3h 270 of aminoacid depletion resulted in a more rapid and robust induction of p4EBP1 in C643 271 cells compared to their isogenic wild-type revertants (Fig. 5F). The mTOR pathway in 272 273 EIF1AX-splice cells was also more sensitive to depletion of glutamine (Supplementary Fig. S5D). 274

### *EIF1AX* and *RAS* mutants converge to stabilize c-MYC, promote mTOR activation and sensitize cells to mTOR, BRD4 and MEK inhibitors.

Deregulated expression of c-MYC in cancer is commonly due to increased protein 278 279 stability. Indeed, the higher c-MYC protein levels in EIF1AX-splice expressing cells not associated with induction of c-MYC mRNA (Fig. 6A) or increased translational efficiency 280 (Supplementary. Fig. S6A). Instead, expression of EIF1AX-splice in KRAS<sup>G12R</sup>-CAL62 281 shifted the half-life of c-MYC from 25 to 60 min as compared to the isogenic parental 282 283 cells (Fig. 6B). Silencing of oncogenic KRAS also decreased c-MYC protein levels (Fig. 6C), although to a lesser extent than in EIF1AX-splice cells, consistent with the latter 284 285 cooperating with oncogenic RAS to further stabilize the protein. The key contribution of RAS and MAPK signaling to c-MYC levels was further confirmed in HTH83 cells, which 286 harbor endogenous HRAS<sup>Q61R</sup> and EIF1AX-A113splice mutations. In these cells HRAS 287 silencing or MEK inhibition decreased c-MYC and ASCT2 expression, whereas the pan-288 PI3K inhibitor GDC-0941 was without effect (Supplementary Fig. S6B and S6C). 289

To explore potential therapeutic dependencies of RAS + EIF1AX mutant thyroid 290 cancers, we investigated the effects of the MEK inhibitor trametinib, the mTOR kinase 291 inhibitor AZD8055 or the BRD4 inhibitor JQ1 (to target c-MYC transcription, with the 292 caveat that JQ1 also inhibits other bromodomain proteins) alone and in various 293 combinations in xenografts of CAL62-splice and parental cells. CAL62-splice xenografts 294 grew to a larger size and were more sensitive to the growth inhibitory effects of 295 AZD8055 or JQ1 than the parental controls, whereas trametinib had equivalent efficacy 296 in both contexts (Fig. 6D; i, ii & iii). The combination of AZD8055 with either trametinib 297 or JQ1 induced tumor shrinkage in CAL62-splice but not in parental cells, and was 298 superior to either drug alone (Fig. 6D; iv, v & vi). Consistent with their effects on growth, 299 Western blots of tumor lysates from mice treated with each condition showed that 300 301 AZD8055 in combination with either trametinib or JQ1 showed the most profound inhibition of c-MYC and mTOR substrates (Fig. 6E). However, despite comparable 302 inhibition of these signaling nodes in EIF1AX WT and mutant cells, the latter show 303 304 preferential tumor shrinkage, consistent with heightened dependency on the pathways 305 activated by these mutant proteins.

EIF1AX is the only PIC component that is recurrently mutated in cancers (4,5,8,21,22).

#### 306 Discussion

307

EIF1AX mutations have been presumed to result in a change- or gain-of-function 308 because of their predilection for specific substitutions in the N- and C-terminal tails. 309 However, functional insights so far have been primarily confined to how EIF1AX 310 mutants alter usage of initiation codons with varying contexts in yeast (25,29). In uveal 311 312 melanomas, *EIF1AX-NTT* mutants are associated with relatively indolent disease. Isolated *EIF1AX* mutations are also found in low-risk thyroid tumors (4,17). When 313 coupled to RAS mutations they mark aggressive and often lethal forms of PDTC and 314 ATC. This is phenocopied in the mouse models we described here. However, the 315 penetrance of these cancers in *EIF1AX-c'spl/Hras<sup>G12V</sup>* mice was relatively low, 316 suggesting that other factors may be required for transformation. In that respect, most 317 human EIF1AX/RAS PDTCs and ATCs harbored either TERT promoter or TP53 318 mutations, which are major drivers of tumor progression in this disease (5,6,36). 319 Phosphorylation of Ser51 of the  $\alpha$  subunit of EIF2 is a central common conduit for many 320 cellular stress pathways, including nutrient/amino acid starvation, ER stress and 321 oxidative insults. This modification of EIF2 $\alpha$  prevents its recycling into the TC, thereby 322 inhibiting global protein synthesis, which helps cells to adapt by conserving nutrients 323 and relieving ER stress. However, a subset of mRNAs, most prominently ATF4, are 324 preferentially translated. ATF4 induces a transcriptional program that includes genes 325 involved in aminoacid transport, metabolism and protection from oxidative stress, which 326 allow cells to orchestrate a more sustained adaptation to these stressors (37-43). The 327 328 induction of ATF4 in *EIF1AX-A113splice* cells is independent of EIF2 $\alpha$  phosphorylation, and takes place by modulating ATF4 translation efficiency as well as inducing its 329 330 transcription. This illegitimate ATF4 activation then hijacks a negative feedback pathway that leads to dephosphorylation of Ser51 of EIF2 $\alpha$ , by inducing expression of GADD34, 331 332 a co-factor for the EIF2a phosphatase PP1 (32,43), thus increasing global protein synthesis. Hence, this mutated translation initiation component attains a gain-of-function 333 334 by deregulating the stringent control of the rate of global protein synthesis by EIF2a

dephosphorylation. This in itself has significant oncogenic potential, as expression of
 hypo-phosphorylated EIF2α is sufficient to transform NIH3T3 cells (44).

337 RAS mutations are found along the entire spectrum of thyroid cancer, although the frequency is markedly enriched in PDTC and ATC (5). Allelic imbalance favoring 338 339 oncogenic Ras gene dosage appears to be critical for transformation, and can be achieved through loss of the WT copy (45) or mutant allele amplification (46-48). 340 341 Intensification of RAS signaling in thyroid cancer can also be mediated through YAPinduced transcriptional activation of RAS, leading to tumor progression (26). Oncogenic 342 RAS acts via ERK to prevent c-MYC degradation, primarily through phosphorylation of 343 serine 62, a site within the c-MYC degron recognized by a specific E3 ubiquitin ligase 344 345 (49,50). pS62-c-MYC primes subsequent phosphorylation at threonine 58, which facilitates dephosphorylation of S62, poising MYC for degradation. We found that 346 EIF1AX-splice increased c-MYC protein half-life, further augmenting the effects of 347 oncogenic RAS signaling on c-MYC protein stability. Interestingly, transcription-348 349 independent MYC-MAX activation was seen in the uveal melanoma (UM) TCGA study in the context of EIF1AX-NTT mutations (16). Consistent with this, the EIF1AX 350 mutations in uveal melanoma display mutual exclusivity to tumors with chromosome 8q 351 gain, which harbors the MYC gene locus (8g<sup>24.21</sup>). Conceivably, by analogy to the RAS-352 353 EIF1AX cooperativity on MYC in thyroid cancer, a potential mechanistic basis of MYC-MAX activation in uveal melanomas may involve interactions with constitutive G-protein 354 oncogenic signaling mediated by the co-occurring GNAQ/GNA11 mutations in EIF1AX-355 mutant uveal melanomas (8,16). 356

357 The intersection of ATF4 and c-MYC on regulation of amino acid transporter expression and mTOR activation is well established (33-35). We found that c-MYC and ATF4 co-358 359 regulated aminoacid transporters in the EIF1AX-RAS context, particularly ASCT2 and LAT1, leading to mTOR activation. Accordingly, expression of ATF4, c-MYC and amino 360 361 acid transporters in a panel of RAS-mutant human thyroid cancer cell lines displayed a striking concordance with *EIF1AX* mutation. These events cooperate to sensitize mTOR 362 363 signaling to aminoacid supply. EIF1AX-splice also slightly augments RAS-induced PI3K signaling in the isogenic cell lines, through unclear mechanisms. However, the 364

activation of mTOR is PI3K-AKT- and RSK-independent, and driven primarily by the
 increased influx of amino acids. Whether increased PI3K contributions to tumorigenesis
 through alternative mechanisms in this context cannot be ruled out.

In summary, *EIF1AX-A113splice*, a mutation commonly encountered in thyroid cancer in association with oncogenic *RAS*, leads to induction of ATF4, which in turn induces a

global increase in protein synthesis through GADD34-dependent dephosphorylation of

- EIF2α. The mutant EIF1AX, in concert with oncogenic RAS, also increases c-MYC
- protein stability. c-MYC and ATF4 cooperate to induce transcription of aminoacid
- transporters, and the resulting aminoacid flux activates mTOR signaling. Our data point
- to mTOR kinase as a primary node for pharmacological targeting, and provides a
- rationale for MEK or c-MYC co-inhibition to maximize therapeutic responses (Fig. 7).

Besides the ~ 11% of advanced thyroid cancers harboring *EIF1AX* and *RAS* mutations,

these two oncogenes also co-occur in low grade serous ovarian cancers (21) and in

some widely invasive Hurthle cell carcinomas. mTOR kinase inhibitors are currently not

- approved for any indication. However, they are still being extensively investigated in
- combination with other agents. The findings reported here provide a strong rationale for
- 381 combined mTOR and MEK inhibitors for tumors harboring these defects.

#### Methods and Materials 383

#### Cell lines. 384

389

Cell lines were maintained at 37°C and 5% CO<sub>2</sub> in humidified atmosphere and were 385 grown in RPMI-1640 for Nthy-Ori 3-1, C643, CAL62 and HTH83, DMEM for HEK293T, 386 HTH7, ACT1 and DMEM:RPMI for KMH2 supplemented with 10% of FBS. 2 mmoL/I 387

glutamine, 50 U/mL penicillin (GIBCO), and 50 µg/mL streptomycin. C643, Hth7 and 388 Hth83 cell lines were obtained from Dr. Nils-Erik Heldin, Uppsala University Hospital,

390 Sweden. CAL62 cells were obtained from Dr. Jeanine Gioanni, Centre Antoine-

Lacassagne, France. The ACT1 line was obtained from Dr. Naoyoshi Onoda, 391

392 Osaka City University Graduate School of Medicine, Japan. KMH2 were obtained from

Japanese Collection of Research Bioresources Cell Bank (JCRB), Japan. All cell lines 393

tested negative for mycoplasma and were authenticated using short tandem repeat and 394

single nucleotide polymorphism analyses. 395

#### 396 Plasmids and constructs.

The cDNAs of the EIF1AX splice variants (c'spl and t'spl) were cloned by PCR 397 amplification from parental EIF1AX-A113splice mutant human thyroid cell line HTH83. 398 399 The full-length cDNAs of human wild type EIF1AX and the EIF1AX splice variants (c'spl and t'spl) were cloned into pLVX-puro and the dox-inducible pLVX-Tight-puro vectors 400 (Clonetech). EIF1AX N-terminal tail (NTT) mutants were generated from pLVX-puro-401 EIF1AX-WT by site-directed mutagenesis (Strategene protocol). The EIF1AX c'spl 402 cDNA was cloned into the pLVX-puro vector or the pLVX-Tight-puro vector. Bicistronic 403 expression constructs for EIF1AX-WT and the two EIF1AX splice variants were 404 generated by sequential cloning into pLVX-Tight-puro, as schematically shown in 405 Supplementary Fig. S2C. The PCR primers used to amplify EIF1AX-WT, to generate 406 NTT mutants by site-directed mutagenesis, or to amplify the specific splice variants in 407 cell lines harboring endogenous EIF1AX mutations are shown in Supplementary Table 408 S5. The TCRS and the TCRS<sup>ΔuORF</sup> vector systems were provided by Dr. Cor Calkhoven 409 (ERIBA, Groningen, NL). The targeting plasmid CoIA1-TRE (CoIA-CHC system for 410

- 411 cDNA expression) used to clone the EIF1AX-c'spl cDNA was provided by Dr. Luke Dow
- 412 (Weill Cornell College of Medicine, NY). All constructs were sequence-verified.

### 413 Development of isogenic EIF1AX-splice-expressing and EIF1AX-splice-repaired 414 thyroid cancer cell lines by CRISPR-Cas9.

CAL62 and TTA1 cells, which are WT for *EIF1AX*, were modified to endogenously 415 express EIF1AX splice variants (cryptic and truncated splice) by targeted disruption of 416 the splice acceptor site by CRISPR-Cas9. A 20bp CRISPR guide sequence (sgRNA) 417 was designed to span the splice site of exon 6, exploiting the endogenous PAM 418 sequence within the splice acceptor site (Supplementary Table S5). The sgRNA was 419 420 annealed and cloned into the pLentiCRISPR vector (Addgene # 49535) that transcribes Cas9 and the CRISPR guide/tracer RNA. Parental CAL62 and TTA1 cells were 421 transfected with pLentiCRISPR-SqRNA using FuGENE HD, followed by selection in 1 422 µg/mL puromycin for 3 days. Cells were then grown for 5 to 7 days, and then plated as 423 single cells into 96-well plates. The propagated clones were tested for disruption of the 424 EIF1AX exon 6 splice site by PCR-based sequencing of genomic DNA encompassing 425 the targeted region. The substitutions introduced in CAL62 and TTA1 cells effectively 426 disrupted the splice acceptor site but were not identical to the naturally occurring 427 endogenous EIF1AX-A113spl mutations. Positive clones of CAL62-splice and TTA1-428 splice were confirmed for expression of EIF1AX c'splice and t'splice mRNAs and their 429 encoded proteins by immunoblotting. 430

The parental C643 cell line, which harbors an endogenous EIF1AX-A113splice 431 mutation, was used to revert the mutant allele by CRISPR-Cas9 knock-in of the 432 433 corresponding WT sequence through use of a homologous directed repair template 434 (HDR). The sgRNA targeting intron 5 was cloned into the pX330 vector (Addgene# 42230). The homologous recombination donor vector was designed using vector 435 builder (https://www.vectorbuilder.com/design.html). Specifically, the donor vector 436 (Vector ID: VB160426-1022eqv) was designed to integrate into the intron 5 locus upon 437 438 recombination directed by homologous arms of 629bp (right) and 701bp (left) flanking a puromycin resistance cassette. The two arms encompassed the 3' end of intron5, the 439 entire exon6 and a fragment of intron6. C643 cells were co-transfected with pX330-440

sgRNA and the donor vector harboring the HDR template. After 72h cells were

subjected to puromycin selection. Surviving cells were seeded as single cells in a 96-

- 443 well plate for testing. Clones that were positive for the targeted locus by PCR-based
- sequencing were tested for expression of the splice variant mRNAs by RT-PCR and by
- immunoblotting. Clone C643-spl-rev (C643-Xcl18) was used in this manuscript.

#### 446 **EIF1AX overexpression in cell lines.**

- 447 Nthy-Ori 3-1 cells (hereafter referred to as NthyOri, derived from wild-type human
- thyroid cells immortalized with large T-antigen) (51) were used to generate stable and

dox-inducible lines expressing EIF1AX-WT, NTT-mutants or the splice variants. The

- 450 pLVX-puro vector cloned with the respective various EIF1AX cDNA's were used for
- 451 constitutive expression, whereas the pLVX-Tet-On Advanced vector system (Clontech)
- 452 was used to generate dox-inducible EIF1AX or bicistronically expressed EIF1AX-splice
- 453 variants, as described in 'Plasmids and Constructs' section.
- 454 To generate stable EIF1AX-expressing NthyOri clones, the constructs described above
- were used for lentiviral production in HEK293FT cells using the Mission Lentiviral
- 456 Packing Mix (SIGMA). The constitutively expressing NthyOri-EIF1AX stable lines were
- 457 generated by infecting with the corresponding viral particles, and the dox-inducible
- 458 NthyOri-EIF1AX cells by co-infecting with lentiviral-transduced pLVX-Tight-puro-EIF1AX
- and pLVX-Tet-On particles in the presence of 8 µg/mL polybrene (Sigma, St Louis, MO)
- 460 overnight. After 24 hours in complete medium, cells were selected in 1 μg/mL
- 461 puromycin with or without 300 μg/ml G418 as required. The mass culture was then
- cloned into single cells, which were expanded and tested for the expression or induction
- 463 of EIF1AX by immunoblotting.

### 464 **Tg-rtTA/TRE-EIF1AX-c'spl and TPO-Cre/FR-Hras**<sup>G12V</sup> mice.

- The dox-inducible *EIF1AX-c'spl* mouse model was developed by the ESC-GEMM
- 466 method (52). ESC derived from *TPO-Cre/FR-Hras<sup>het</sup>/RIK<sup>het</sup>/CHC<sup>het</sup>* mice were used to
- 467 target the human EIF1AX-c'spl cDNA into the homing cassette that directs site-specific
- 468 integration of the transgene downstream of the Col1a1 locus by recombination-
- 469 mediated cassette exchange. The *TPO-Cre* drives Cre recombinase under the control

of the thyroid peroxidase gene promoter, which is active in thyroid cells at E14.5 (53). 470 ESC clones targeted with TRE-EIF1AX-c'spl into the Cola1-homing cassette (CHC) 471 472 were microinjected into blastocysts produced from NCI C57BL/6-cBrd/cBrd/Cr (C57BL/6 albino) mice and implanted into CD-1 pseudo-pregnant mothers enabling production of 473 chimeric pups. To achieve higher expression of rtTA we bred out the RIK<sup>het</sup> cassette 474 and bred in thyroglobulin-driven rtTA (Tg-rtTA(s)M2) (54). The resulting TPO-Cre/FR-475 *Hras<sup>het</sup>/Ta-rtTA /TRE-EIF1AX<sup>het</sup>* mice were intercrossed to generate the following lines: 476 TPO-Cre/FR-Hras<sup>hom</sup>, Tg-rtTA/TRE-EIF1AX<sup>hom</sup> and TPO-Cre/FR-Hras<sup>hom</sup>/Tg-rtTA/TRE-477 EIF1AX<sup>hom</sup>. All animals were fed doxycycline-impregnated chow (TD01306, Envigo), 478 and the appropriate lines verified by immunoblotting to achieve dox-inducible 479 expression of EIF1AX-c'spl in thyroid follicular cells (Fig. 2C). Animal care and all 480 procedures were approved by the MSKCC Institutional Animal Care and Use 481

482 Committee.

#### 483 Histology and Immunohistochemistry.

484 Mouse thyroids dissected from surrounding tissues were fixed in 4% paraformaldehyde,

embedded in paraffin, sectioned and stained with hematoxylin and eosin (H&E).

486 Histological diagnosis was performed by a thyroid pathologist (RG) blinded to mouse

- 487 genotype. Sections were also immunostained for KI67. Slides were scanned with
- 488 Pannoramic Flash 250 (3DHistech, Budapest, Hungary), and whole thyroid lobes or
- regions of interest viewed using Pannoramic Viewer and exported as tiff images. H&E
- and IHC were performed by the MSK Molecular Cytology Core Facility.

#### 491 Western Blotting and Immunoprecipitation.

Cells were lysed in 1X RIPA buffer (Millipore) supplemented with protease (Roche) and
phosphatase inhibitor cocktails I & II (Sigma). Tumors or xenografts were homogenized
in 1X Lysis Buffer (containing 10 mmol Tris-HCl, 5 mmol EDTA, 4 mmol EGTA, 1%
Triton-X100) with protease/phosphatase inhibitors. Lysates were briefly sonicated to
disrupt the tissue and cleared by centrifugation. Protein concentrations were estimated
by BCA kit (Thermo Scientific) on a microplate reader (SpectraMax M5); comparable
amounts of proteins were subjected to SDS page using NuPAGE 4–12% Bis–Tris

499 gradient gels (Invitrogen) and were then transferred to PVDF membranes. Following

- 500 overnight primary antibody incubation, membranes were incubated with secondary
- antibodies coupled to horseradish peroxidase (HRP) or IRDye fluorophores for 1h at
- 502 room temperature. HRP probed blots were developed using enhanced
- 503 chemiluminescence reagent (Amersham Biosciences), and signal captured using X-ray
- <sup>504</sup> films or with the KwikQuant<sup>™</sup> Imager (<u>http://kindlebio.com/index.php</u>). IRDye-probed
- 505 blots were imaged using the LICOR Odyssey imaging system (Licor Biosciences).
- 506 Immunoblot quantification was done using ImageJ.

507 For co-immunoprecipitation experiments, cells were lysed in buffer containing 75 mM NaCl, 50 mM Hepes, 10 mM MgCl2, 8 mM EGTA, 10 mM β-glycerophosphate, 1 mM 508 509 DTT, 0.5% Triton X-100, along with protease and phosphatase inhibitors. Equal amounts of lysate (500µg) were diluted with lysis buffer in 300µl final volume; 1/10<sup>th</sup> of 510 the volume was denatured and used as input control. Antibodies were incubated with 511 lysates overnight by end-to-end rotation at 4 °C. Antigen-antibody complexes were 512 513 immobilized by incubating with Dyna beads for 1h, the antibody-antigen-beads collected by DynaMag<sup>IM</sup>-spin (Invitrogen), washed with lysis buffer, denatured and subjected to 514 Western blotting.

515 Western blotting

#### 516 Antibodies.

- 517 The following primary antibodies were used for immunoblots at a dilution of 1:1000,
- 518 except where indicated. EIF1AY (sc-84243) was used to immunoblot EIF1AX, EIF5 (sc-
- 519 28309), EIF2α (sc-11386), HRAS (sc-520; 1:500), KRAS (sc-30; 1:500), YB1 (sc-
- 101198), HA-tag (sc-805) were obtained from Santa Cruz biotechnology. c-MYC
- 521 (5605), ATF4 (11815), LAT1 (5347), ASCT2 (5100), pEIF2α-S51 (9721), pAKT-S473
- 522 (4051), pAKT-T308 (4056), AKT (2920), pP70S6K-T389 (9234), P70S6K (2708),
- 523 p4EBP1-S65 (9451), p4EBP1-T37/46 (2855), 4EBP1 (9452), pYB1-S102 (2900),
- 524 pRPS6-S240/244 (2215), pRPS6-S235/236 (2211), RPS6 (2317), pERK (9101), ERK
- 525 (4696), HA-Tag (3724), Biotin (5597), ER stress antibody sampler kit (9956) were from
- 526 Cell Signaling, pPRAS40-T246 (441100G), GADD34 (PA1-139) from Invitrogen,
- 527 ASCT2 (HPA035240) and β-actin (A2228; 1: 10000) from Sigma Aldrich.

- 528 The secondary antibodies were used at a dilution of 1:5000. HRP-conjugated antibodies
- 529 included goat anti-rabbit (Santa Cruz; sc-2004) and goat anti-mouse (Santa Cruz; sc-
- 530 2031). IRDye fluorophore-conjugated antibodies were IRDye® 800CW Goat anti-Rabbit
- 531 IgG (Licor; 926-32211), IRDye® 800CW Goat anti-Mouse IgG (Licor; 926-32210),
- 532 IRDye® 680RD Goat anti-Rabbit IgG (Licor; 926-68071), and IRDye® 680RD Goat anti-
- 533 Mouse IgG (Licor; 926-68070). We also used the following additional reagents:
- 534 Doxycycline (2 µg/mL) and cycloheximide were from Sigma. Salubrinal was from
- 535 Calbiochem. LJI308 was from Selleckchem.

#### 536 L-azidohomoalanine (AHA) labeling.

Isogenic cell lines used to assess nascent protein synthetic rate were grown in 60-mm 537 dishes until ~70% confluent. Prior to labeling cells were incubated with methionine-free 538 media containing 2% FBS for 1h, followed by addition of 50 µm AHA (Life Technologies, 539 C10102) for 20min. Cells were then washed in cold PBS and immediately lysed in buffer 540 containing 50 mM Tris HCI, pH 8.0, and 1% SDS. Complete lysis was achieved by 541 sonication. Comparable amounts of lysates were then subjected to Click-iT reaction for 542 switching azido-modified nascent proteins to alkyne-biotin (Life Technologies, B10185) 543 using the Click-iT<sup>™</sup> Protein Reaction Buffer Kit (Life Technologies, C10276) following 544 the manufacturer's protocol. Biotinylated nascent proteins were subjected to 545 immunoblotting using either anti-biotin, EIF1AX or  $\beta$ -actin primary antibodies and the 546 corresponding IRDye-fluoropore-conjugated secondary antibodies. Images were 547 captured by the LICOR Odyssey imaging system. Biotinylated proteins on the entire 548 549 lane was quantified using Odyssey application software version 3.0 (LICOR 550 Biosciences).

#### 551 **RNA interference.**

- 552 We used SMARTpools (Dharmacon) for ATF4 and c-MYC silencing (Dharmacon). Cells 553 grown without antibiotics at 70% confluence were transfected using 50 nmol/L of
- 554 Smartpools and 6-8 µL of DharmaFECT (Dharmacon). SiRNAs for HRAS and KRAS
- were from ORIGENE and were transfected using SiTran1.0 as per manufacturer's

instructions. Cells were harvested 72h post-transfection and analyzed by Westernblotting.

#### 558 **RNA isolation, cDNA synthesis and qPCR.**

- 559 Total RNA from isogenic cell lines was extracted using the PrepEase Kit (USB
- 560 Corporation). Comparable amounts of RNA (1µg) were subjected to DNase I
- 561 (Invitrogen) treatment and reverse transcribed using SuperScript® III Reverse
- 562 Transcriptase (Invitrogen) following the manufacturer's protocol. qPCR was then
- 563 performed with the Power SYBR Green PCR Master Mix (Applied Biosystems). Primers
- used are shown in Supplementary Table S5. The Ct values of the target genes were
- normalized to β actin and the normalized expression analyzed by the ΔΔCt method.

#### 566 **c-MYC half-life.**

- 567 The relative half-life of c-MYC protein in CAL62 vs. CAL62-splice cells was analyzed by
- <sup>568</sup> measuring c-MYC protein abundance after treatment with cycloheximide (CHX). Cells
- plated in 60-mm dishes and grown 24h in low serum condition (0.5% FBS) were treated
- 570 with 100 µg/ml cycloheximide (Sigma), harvested and lysed at the indicated time points
- 571 and analyzed by Western blotting.

#### 572 Colony formation assay.

- 573 Dishes were first coated with a bottom layer of 0.4% agar in RPMI. Cells were
- resuspended in a top layer of 0.2% agar in RPMI with 10% FBS, and then fed every
- other day by adding drops of media onto the top layer. After 3 weeks, the colonies were
- stained with crystal violet and counted in a GelCount<sup>™</sup> colony counter (Oxford
- 577 OPTRONIX). Minimum diameter of the colonies was set to 100µm.

#### 578 **Tumor xenografts.**

- 579 Approximately 6-8 week-old female SCID mice (NOD.CB17-Prkdc; Envigo RMS.Inc)
- were injected subcutaneously with  $5 \times 10^6$  CAL62 or CAL62-splice cells grown to 70%
- confluence and resuspended in 50% Matrigel (CORNING) into the right and left flanks,
- respectively. Treatments were administered by oral gavage when tumor volume

approached 200 mm<sup>3</sup> as estimated by measuring the length and width with calipers 583 (width<sup>2</sup>  $\times$  length  $\times$  0.52). Tumor-bearing mice were randomly assigned into 5 treatment 584 585 arms: Controls (vehicle-4% DMSO in 30% PEG 300); AZD8055 (10 mg/kg); Trametinib (0.75 mg/kg); JQ1 (40 mg/kg); AZD8055 + Trametinib and AZD8055 + JQ1 586 (all drugs were from Selleckchem). Mice were weighed at the start of treatment and 587 every second day during the treatment period. AZD8055 was dissolved in a mixture of 588 4% DMSO and 30% PEG 300 (SIGMA), trametinib in 4% DMSO in corn oil and JQ1 in 589 2% DMSO, 30% PEG 300 and 5% Tween 80. Treatments were administered by oral 590 gavage in a volume of approximately 200 µL. Tumor volume was measured every 2-3 591 days with calipers. After 21 days mice were humanely killed, and dissected tumors 592 flash frozen for subsequent protein isolation. All animal experiments were performed 593 in accordance with a protocol approved by the Institutional Animal Care and Use of 594 Committee (IACUC) of Memorial Sloan Kettering Cancer Center. 595

#### 596 Statistical analyses.

597 Statistical analysis for animal studies and cell lines was performed using GraphPad 598 Prism 7. P value was determined by two-tailed t-tests. F test was used to compare 599 variances between the groups, if different, Welch correction was applied. Data are 500 shown as mean with SD or mean with 95% CI (n=3 or more biological replicates).

#### 601 Translation Efficiency Analysis by Ribosome profiling.

Ribosome profiling and RNA seg were performed in C643 and C643-spl-rev cells. 602 Triplicates of cells grown in 150mm dishes were treated with cycloheximide for 10 min 603 and ribosome-protected RNA fragments isolated following the published protocol for 604 ribosome profiling (55) with a modification of including unique molecular identities (UMI) 605 in the library reverse transcription primer (Supplementary Table S5) in order to remove 606 PCR duplicates during analysis. Parallel total RNA extraction was performed for RNA-607 seq. Ribosome profiling reads and RNA-Seq reads were aligned using STAR v2.5. (56) 608 using the UCSC human genome reference, hg19 609 (http://hgdownload.cse.ucsc.edu/goldenPath/hg19/chromosomes) with ERCC spike-ins 610

611 included as an extra chromosome. To process the ribosome profiling reads before

alignment, linker sequences (5'-CTGTAGGCACCATCAAT-3') were removed using 612 Trimmomatic v0.32 (57) with the following parameters: number of mismatches between 613 read and adapter: 2; length of alignment between read and adapter: 6. Clipped reads 614 were then filtered to be a minimum length of 25. Reads with unique molecular 615 identifiers (5'-NNNNTGANNNNCC-3') were removed from the sequence and inserted 616 617 into the read name using UMI tools v2.1.1 (58). The set parameters for STAR during alignment were as follows: each read must uniquely map; number of mismatches: 2; 618 maximum intron alignment length: 500000; 3' adapter sequence: CTGTAGGCAC; 619 maximum proportion of mismatches within adapter: 0.1; default values were used for all 620 remaining parameters: 621

- 622 STAR --runThreadN 4 --genomeLoad NoSharedMemory --outSAMtype BAM
- 623 Unsorted --outSAMstrandField intronMotif --outSAMattributes NH HI NM MD AS
- 624 XS --outSAMunmapped Within --outSAMheaderHD @HD VN:1.4 --
- outFilterMultimapNmax 0 --outFilterMultimapScoreRange 1 --
- outFilterScoreMinOverLread 0.33 -- outFilterMatchNminOverLread 0.33 --
- outFilterMismatchNmax 2 --alignIntronMax 500000 --alignMatesGapMax 1000000 -
- -alignSJDBoverhangMin 1 --sjdbOverhang 100 --sjdbScore 2 --readFilesCommand
- 229 zcat --clip3pAdapterSeq CTGTAGGCAC --clip3pAdapterMMp 0.1
- 630 Read alignments are available on NCBI SRA under accession number SRP142722. To
- remove possible rRNA contamination, both the ribosome profiling and RNA-Seq reads
- were aligned to ribosomal sequences gathered from BioMart Ensembl (59) and SILVA
- (60) databases and merged into a single FASTA reference file (GEO accession
- 634 GSE113695). Reads were again aligned using STAR with almost all the same
- parameters, but reads were allowed to align to a maximum of 3 other regions in our
- rRNA FASTA reference. All reads which aligned to the rRNA reference according to the
- criteria above were filtered from the original genome reference aligned reads:
- 638 STAR --runThreadN 4 --genomeLoad NoSharedMemory --outSAMtype BAM
- 639 Unsorted --outSAMstrandField intronMotif --outSAMattributes NH HI NM MD AS
- 640 XS --outSAMunmapped Within --outSAMheaderHD @HD VN:1.4 --
- outFilterMultimapNmax 3 --outFilterMultimapScoreRange 1 --
  - 23

outFilterScoreMinOverLread 0.33 --outFilterMatchNminOverLread 0.33 --

outFilterMismatchNmax 2 --alignIntronMax 500000 --alignMatesGapMax 1000000 -

- -alignSJDBoverhangMin 1 --sjdbOverhang 100 --sjdbScore 2 --readFilesCommand
- 645 zcat --clip3pAdapterSeq CTGTAGGCAC --clip3pAdapterMMp 0.1

For final filtering, all ribosomal reads larger than 35 base pairs were removed and all
reads aligning to the same position with the same UMI were reduced to a single read
using UMI tools. The final alignment files used for quantifications are available on GEO
accession GSE113695.

To estimate abundance on the aligned BAM files, a custom script was used with

651 gencode annotation version 19 and additional ERCC spike-in sequences (ERCC92).

For RNA-Seq samples with spike-ins, the library size normalization factor was estimated

using DESeq2 v3.6 (61). The library size normalization was used in the differential

translation efficiency analysis. RiboDiff v0.2.1 (62) with default parameters was used to

estimate the change in translational efficiency between sample conditions. Only protein

coding genes (Gencode v19) were considered in RiboDiff. The result of RiboDiff is

available in the Supplementary Table S1 and S2. We used the following command line:

658

ribodiff -p 1 -s 10 -m BH

#### 659 **Polysome Fractionation.**

NthyOri-splice dox-inducible cells were treated with or without dox for 72 h, followed by 660 cycloheximide (100µg/ml) for 15 min. Cells were then trypsinized and lysed with buffer 661 containing 15mM Tris-HCl, 300mM NaCl, 15mM MgCl2, 1% Triton X-100, 0.1mg/ml 662 663 cycloheximide and ribonuclease inhibitor (RNasin, PROMEGA). Comparable amounts of lysates (1.5 mg protein) were then layered onto a 10-60% sucrose density gradient 664 prepared in 15mM Tris-HCl, 300mM NaCl, 15mM MgCl2, 0.1mg/ml cycloheximide and 665 RNasin, and fractionated using a SW60Ti rotor in a Beckman ultracentrifuge for 2 h at 666 667 37,000 rpm at 4°C. After centrifugation, the gradients were collected manually from the top into 12 fractions. Fractions were subjected to RNA extraction for ATF4, c-MYC and 668 669  $\beta$ -actin mRNA measurements by real time PCR as described (63).

#### 670 **RNAseq-Gene Set Enrichment Analysis (GSEA).**

- 671 Gene set enrichment analysis (64) was performed with GSEA software
- 672 (http://www.broadinstitute.org/gsea/) using the pre-defined Canonical Pathways and
- 673 Hallmarks Molecular Signatures Database (MSigDB) gene sets
- 674 (http://software.broadinstitute.org/gsea/msigdb/index.jsp). The normalized counts of
- each replicate (GEO accession GSE113695) derived from the RNA-seq of C643 vs
- 676 C643-spl-rev and CAL62-splice vs CA62 cells were used as a dataset to run GSEA
- analysis (Identification-Gene symbol; permutations-1000 and permutation type-gene
- sets). Briefly, GSEA software-derived enrichment scores (ES) identified the functional
- group of genes (pre-defined datasets in the MsigDB) over-represented in the given data
- set. The Normalized Enrichment Score (NES) was used to determine statistical
- significance from the nominal p value after controlling for false positives by calculating
- false discovery rate (FDR). Key top-ranked signatures based on NES, Nom-p-value
- and FDR-q value prompted validation in our experimental models.

#### 684 Luciferase and TCRS assay.

The pGL3-firefly vector was engineered with the ATF4 uORF initiation context by site directed mutagenesis (primers shown in Supplementary Table S5). HEK293T cells were co-transfected with EIF1AX-WT, G9R or c'splice expression vectors along with the engineered firefly vectors using Fugene HD. Renilla luciferase (pRL-null) was cotransfected as a transfection efficiency control. Cells were incubated in 0.5% FBS for 48h and luciferase activity measured using the Dual-Glo Luciferase Assay system on the GloMax-Multi Microplate Reader (Promega).

The efficiency of a ribosomal re-initiation mode of translation in EIF1AX-WT and the mutants was assessed with the translation control reporter system (TCRS) as described previously (30). The TCRS construct has a short uORF and 2 overlapping ORFs encoding a long peptide (L.P) and a short peptide (S.P), respectively, the latter being a marker of ribosomal re-initiation. HEK293T cells co-transfected with TCRS and EIF1AX-WT, -NTT mutants or EIF1AX-c'spl expression vectors. The expression of the short peptide was analyzed by immunoblotting.

#### 699 Data availability.

- Ribosome profiling and RNA sequencing data from this study have been submitted to
- the NCBI Gene Expression Omnibus under accession number GSE113695.
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#### 896 Figure Legend

### Figure 1: *EIF1AX* mutation is a strong co-operating event with *RAS* in advanced thyroid cancer; the hotspot *A113splice* mutation induces aberrant splice variants.

- (A) *Top: EIF1AX* and *RAS* mutations in 151 PDTC and 95 ATC, compiled from MSK
- clinical series (n=148 as of Oct 2017), Landa et al (n=76) (5) and Kunstman et al (n=22)
- 901 (7). Oncoprint shows co-occurrence of *EIF1AX* and *RAS* mutations in 25/26 tumors;
- p=3.15x10E-13; Fisher's exact test. Green: *EIF1AX* or *RAS* missense mutations; Black:
- 903 EIF1AX-A113 splice mutation. Bottom: Distribution of EIF1AX mutations, showing cluster
- of missense mutations in the N-terminal tail and in a hotspot splice acceptor upstream
- of exon 6 in the C-terminal tail (red dots (MSK series); blue (5) and yellow (7)).
- 906 (B) Scheme of aberrant mRNA splicing by the *EIF1AX-A113splice* mutations c.338-
- 907 1G>A/T or c.338-2A>T/G, which abolish the splice acceptor site of exon 6 (dotted line),
- resulting in two alternatively spliced mRNA's: 1. Cryptic splice (c'spl) mRNA, through
- usage of a cryptic splice acceptor site within exon 6, yielding a 132aa protein (Black
- dotted line: normal splice junction; red dotted line: cryptic splice junction; green arrow:
- position of in-frame 12aa loss). 2. Intron 5 retained mRNA: failure to splice out intron 5
- 912 (blue dotted line) yields a115aa truncated protein (t'spl). Red asterisk points to stop
- 913 codon.
- C) EIF1AX Western immunoblot of human *RAS* mutant cell lines with or without the
- 915 endogenous *EIF1AX*-A113splice mutation. Arrows point to EIF1AX-wt and the mutant
- 916 EIF1AX-c'spl and EIF1AX-t'spl proteins.

### 917 **Figure2: Aberrant splice variants of** *EIF1AX-A113splice* **mutation induce**

918 transformation *in vitro* and cooperate with oncogenic *RAS* to promote mouse

- 919 **thyroid tumorigenesis.**
- A) Left: EIF1AX immunoblot of EIF1AX-A113splice knock-in cells (CAL62-splice and
- 921 TTA1-splice) showing de novo expression of cryptic and truncated splice variants, and
- 922 their loss in repaired cells (C643-spl-rev), in which the splice mutation has been
- reverted to WT. *Middle and right panels*: Soft agar colony assays of the corresponding
  - 31

- 924 isogenic cell lines in the EIF1AX-splice and EIF1AX/RAS contexts. \*p<0.0002,
- <sup>925</sup> \*\*p<0.0001, \*\*\*p<0.0001; two tailed unpaired t test; n=3.
- B) Targeting constructs used to develop *Tg-rtTA/TRE-EIF1AX-c'spl/TPO-Cre/FR- Hras*<sup>G12V</sup> mice.
- C) EIF1AX immunoblot of lysates of mouse thyroid tissues in *Tg-rtTA/TRE-EIF1AX-c'spl*
- mice fed with or without dox diet for 40-51 weeks.
- D) *Top:* Representative H&E stained thyroid sections of the indicated genotypes after
- 40-51 weeks of dox. Histology of mice with isolated Hras or EIF1AX-c'spl revealed
- 932 hyperplasia, whereas compound *Tpo-Cre/FR-Hras<sup>G12V</sup>*/Tg-rtTA/*TRE-EIF1AX-c'spl* mice
- 933 developed tumors consistent with PDTC; arrow pointing necrosis. *Bottom*:
- 934 Corresponding KI67 IHC.
- E) Histological characteristics of thyroid tissues from *TPO-Cre/FR-Hras*<sup>G12V</sup>, *TRE-*
- 936 EIF1AX-c'spl and TPO-Cre/FR-Hras<sup>G12V</sup>/TRE-EIF1AX-c'spl mice. Prevalence of thyroid
- 937 cancer: *EIF1AX/Hras* vs. *EIF1AX* p = 0.03; *EIF1AX/Hras* vs *Hras* p = 0.01 (unpaired t
- test). \*Hyperplasia of *Hras*-mutant tumors lacks atypical features, whereas in the
- 939 *EIF1AX* context all hyperplastic lesions had endocrine atypia.

### 940 Figure 3: *EIF1AX* mutants have higher affinity to components of the translation

- 941 **PIC and increase protein synthesis.**
- 942 (A) Input lysates (*left*) and Co-IP (*right*) of HEK293T lysates transfected with empty
- vector or HA-tagged EIF1AX-WT or the following EIF1AX mutants: -G8R, -G9R, -G15V
- or -c'spl. EIF1AX mutants efficiently co-IP the ternary complex component EIF2 $\alpha$ .
- 945 (B) HEK293T lysates transfected with the indicated constructs showed more efficient
- 946 co-IP of EIF1AX mutants vs WT-EIF1AX with an antibody to the PIC subunit EIF5. *Left:*
- 947 Input lysates. *Right:* EIF5 co-IP.
- 948 (C) EIF5 co-IP of EIF1AX in the following contexts: NthyOri cells with dox-inducible
- 949 expression of EIF1AX-c'splice, EIF1AX-splice knock-in (CAL62-splice) and in splice-
- <sup>950</sup> reversed (C643-spl-rev) cells compared to their respective parental isogenic cells.

Splice variants were IP more efficiently than EIF1AX-WT, which was associated with
higher pull-down of the ternary complex protein EIF2α. Arrows point to EIF1AX WT, c'spl and -t'spl proteins.

- (D) *De novo* protein synthesis of *EIF1AX*-splice isogenic lines (left) or NthyOri cells
- transfected with the indicated EIF1AX constructs (right) as determined by L-
- azidohomoalanine (AHA) labeling. Cells were starved of methionine for 1h and
- 957 incubated with AHA for 20min. Lysates were subjected to a click-it chemistry reaction to
- 958 switch azido-modified nascent proteins to alkyne-biotin, and visualized by anti-biotin
- immunoblotting. Bottom panel: Immunoblots for EIF1AX and  $\beta$ -actin.
- 960 (E). Quantification of biotinylated nascent protein in the indicated cells (\*p<0.03,
- <sup>961</sup> \*\*p<0.002, \*\*\*p<0.015, \*\*\*\*p<0.03, \*\*\*\*\*p<0.02; paired t test). Data in C, D and E are
- 962 representative of three independent experiments

## Figure 4: Increased global protein synthesis by EIF1AX-splice is mediated by ATF4-induced EIF2α dephosphorylation.

- 965 (A) Schematic representation of negative feedback inhibition of EIF2α phosphorylation
- by ATF4. ATF4 induces GADD34 mRNA, a component of the protein phosphatase 1
- 967 complex, which dephosphorylates EIF2α. Salubrinal, a selective GADD34/PP1 inhibitor,
- 968 reverses this effect, leading to derepression of protein synthesis.
- (B) Left: Western immunoblot of the indicated proteins in NthyOri-splice and
- 970 RAS/EIF1AX-splice isogenic lines. *Right:* Quantification of GADD34/βactin and
- pEIF2α/tEIF2α in EIF1AX-splice-expressing cells (GADD34: \*p<0.04, \*\*p<0.019,
- 972 \*\*\*P<0.01; EIF2α: \*p<0.001, \*\*p<0.006, \*\*\*p<0.02; paired t test).
- 973 (C) *Top*: Immunoblot for EIF1AX, total and pEIF12 $\alpha$  in vehicle or Salubrinal treated 974 C643 and C643-spl-rev cells. *Bottom*: *De novo* protein synthesis by AHA labeling of 975 C643 vs. C643-spl-rev cells treated with vehicle or the indicated concentration of 976 Salubrinal. *Right*: Salubrinal attenuated protein synthetic rate in EIF1AX-splice cells 977 (\*p< 0.005, \*p<0.001; paired t test). Data in B and C are representative of three 978 independent experiments.

### Figure 5: EIF1AX activates mTOR through aberrant expression of ATF4 and cMYC.

(A) GSEA analysis from RNAseq of CAL62-splice and C643 compared to respective
isogenic WT controls using MsigDB 'hallmarks' and 'canonical pathways' genesets
showed activation of pathways associated with c-MYC, mTORC1, preinitiation complex
formation, translation and tRNA aminoacylation. NES were plotted and Nom pvalue/FDR q-value is indicated for each signature.

(B) Immunoblot for PI3K and mTOR substrates in the indicated isogenic thyroid cell

987 lines modified for expression of EIF1AX-splice in RAS-WT (dox-inducible NthyOri cells,

- 988 TTA1) or *RAS* mutant cells (CAL62 and C643).
- 989 (C) Western blots of thyroid tissue lysates of *Tg-rtTA/TRE-EIF1AX-c'spl* mice fed with
- regular diet or dox for 4 weeks. Membranes were probed with antibodies to the
- <sup>991</sup> indicated proteins. Each lane contains a thyroid lobe lysate from a separate mouse.
- (D) Western blot showing effect of Si-RNA silencing of ATF4, c-MYC, or both on
- abundance of AA transporters and on p4EBP1 in C643 vs C643-spl-rev cells.
- (E) Co-silencing of ATF4 and c-MYC in dox inducible NthyOri-splice cells disrupted theEIF1AX-splice induction of AA transporters and 4EBP1 phosphorylation.
- 996 (F) C643 and C643-spl-rev cells were serum starved overnight, incubated with AA free
- media for 3h followed by stimulation with glutamine (6mM) and Leucine (2mM). QL
- stimulated cells lysed at indicated time points were immunoblotted for p4EBP1. Data in
- B, C, D, E, and F are representative of two independent experiments.

### Figure 6: EIF1AX and RAS mutants converge to stabilize c-MYC, promote mTOR activation and sensitize cells to mTOR, BRD4 and MEK inhibitors.

- 1002 (A) RT-PCR of c-MYC and ASCT2 mRNA levels in CAL62 and CAL62-splice cells (*left*)
- and in dox-inducible NthyOri-splice cells transfected with Hras<sup>G12V</sup> or empty vector
   (*right*).

- 1005 (B) c-MYC protein half-life determined by cycloheximide (CHX: 100µg/ml) chase in
- 1006 CAL62 and CAL62-splice cells. *Left:* Lysates from cells harvested at the indicated time
- points after addition of CHX were immunoblotted for c-MYC and actin. *Right:* Half-life of
- 1008 c-MYC was determined by plotting for its degradation over time after normalizing for
- actin; c-MYC and  $\beta$ -actin band intensity was quantified using ImageJ software; data
- 1010 points are from two independent experiments.
- 1011 (C) *Left:* Western blot of CAL62 and CAL62-splice cells transfected with two
- 1012 independent SiRNAs to Kras or a scrambled control (Scr). Knockdown of oncogenic
- 1013 Kras decreased c-MYC abundance and expression of ASCT2.
- 1014 (D) Effects of the mTOR kinase inhibitor AZD8055, trametinib, or the BRD4 inhibitor
- JQ1 on CAL62 or CAL62-113splice xenografts in SCID mice. Mice were orally gavaged
- 1016 once a day with vehicle or the indicated drugs at the following concentrations: vehicle:
- 1017 4% DMSO + 30% PEG300, 10 mg/kg AZD8055, 0.75 mg/kg trametinib or 40 mg/kg JQ1
- alone (i,ii,iii) or in combinations (iv,v) for 21 days. Data represent mean with SD of 5
- 1019 mice/group. vi) Average tumor volume on day 21 with respect to vehicle-treated mice (\*
- 1020 p<0.0009, unpaired t test with Welch's correction; \*\*p< 0.003, unpaired t test)
- (E) Western blots of the indicated xenograft lysates for pERK, c-MYC, and p4EBP1.
- 1022 Each replicate lysate came from a separate tumor sample.

#### 1023 Figure 7: Mechanism of EIF1AX/RAS co-operation and nodes for targeting

- 1024 Mechanisms of mutant *EIF1AX* co-operation with *RAS* in thyroid tumorigenesis.
- 1025 EIF1AX-splice activates ATF4 expression, inducing a GADD34/PP1-mediated negative
- 1026 feedback dephosphorylation of EIF2α, leading to enhanced TC/PIC loading and an
- increase in global protein synthesis. RAS in turn stabilizes c-MYC, an effect augmented
- 1028 by EIF1AX-splice. ATF4 and c-MYC induce expression of AA transporters, and
- 1029 cooperate to activate mTORC1. The potential targetable nodes that disrupt the
- 1030 oncogenic drive of *EIF1AX* + *RAS* are indicated.

### Figure 1





	Mice #	Age (wk)	Dox (wk)	Normal Histology	Hyperplasia with Endocrine Atypia	Hurthle Cell Adenoma	PTC		PDTC		
Genotype							Variant	#	Variant	#	
TPO-Cre/FR- Hras <sup>G12V</sup>	14	29-81	NA	10	4*	0	NA	0	NA	0	
Tg-rtTA/TRE- ElF1AX-c'spl	19	46-62	35-51	0	18	0	classical, encapsulated	1	NA	0	
TPO-Cre/FR- Hras <sup>G12V</sup> /Tg- rtTA/TRE- EIF1AX-c'spl	15	46-62	35-51	0	8	2	Classical/low grade PTC, encapsulated, solid and follicular growth	2	encapsulated/ invasive, PTC phenotype, microfollicular	3	
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Figure 3



### Figure 4



### Figure 5 B



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2.5 5 7.5 10 15

**EIF1AX** 

p4EBP1(S65) Total-4EBP1 β-ACTIN

2.5 5

0

QL stimul. min

AA depriv.

7.5 10 15

- 0

Α





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# **CANCER DISCOVERY**

# EIF1AX and RAS mutations cooperate to drive thyroid tumorigenesis through ATF4 and c-MYC

Gnana P. Krishnamoorthy, Natalie R Davidson, Steven D Leach, et al.

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