Mapping the unique and shared functions of oncogenic KRAS and RIT1 with proteome and transcriptome profiling

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19 Abstract

20

21 Aberrant activation of RAS oncogenes is prevalent in lung adenocarcinoma, with somatic 22 mutation of KRAS occurring in ~30% of tumors. Recently, we identified somatic mutation of the 23 RAS-family GTPase RIT1 in lung adenocarcinoma, but relatively little is known about the 24 biological pathways regulated by RIT1 and how these relate to the oncogenic KRAS network. 25 Here we present quantitative proteomic and transcriptomic profiles from KRAS-mutant and 26 *RIT1*-mutant isogenic lung epithelial cells and globally characterize the signaling networks 27 regulated by each oncogene. We find that both mutant KRAS and mutant RIT1 promote S6 28 kinase, AKT, and RAF/MEK signaling, and promote epithelial-to-mesenchymal transition and 29 immune evasion via HLA protein loss. However, KRAS and RIT1 diverge in regulation of 30 phosphorylation sites on EGFR, USO1, and AHNAK proteins. The majority of the proteome 31 changes are related to altered transcriptional regulation, but a small subset of proteins are 32 differentially regulated by both oncoproteins at the post-transcriptional level, including 33 intermediate filament proteins, metallothioneins, and MHC Class I proteins. These data provide 34 the first global, unbiased characterization of oncogenic RIT1 network and identify the shared 35 and divergent functions of oncogenic RIT1 and KRAS GTPases in lung cancer. 36

37 Introduction

38 Somatic mutation of the KRAS proto-oncogene is a prevalent feature of human cancers,

39 particularly in lung adenocarcinomas where KRAS is mutated in up to 30% of tumors. Cancer-

- 40 associated variants such as G12V and Q61H alter the normal regulation of KRAS GTPase
- 41 activity by disrupting GTP hydrolysis activity or physical interaction with GTPase-activating
- 42 proteins (GAPs)^{1,2}, resulting in heightened downstream cellular signaling through the canonical
- 43 RAS effector pathways RAF/MEK and PI3K/AKT as well as others. Following the discovery of

cancer-associated RAS mutations in the 1980s^{3,4}, thousands of studies have delineated the
critical pathways involved in RAS-mediated cellular transformation, metastasis, and metabolism.

47 Recently, another RAS-family GTPase gene, RIT1, was found to harbor somatic mutations in lung cancer⁵ and myeloid leukemias⁶. Interestingly, germline *RIT1* mutations are found in 48 49 families with Noonan Syndrome, a developmental "RAS"-opathy involving altered craniofacial morphology and cardiac abnormalities⁷, and which can also be caused by germline mutations in 50 51 KRAS itself or other RAS-pathway genes such as SOS1, SOS2, LZTR1 and SHOC2 52 (https://omim.org/). In cancer and development, RIT1 mutations are found in cases that lack 53 canonical KRAS mutations, suggesting that RIT1 may impart the same phenotypes conferred by 54 activation of RAS.

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Prior studies have characterized the role of RIT1 in neural development⁸ and we and others 56 have described the role of mutant RIT1 in cellular transformation^{5,9,10}, knowledge of the function 57 58 of cancer- and Noonan-associated RIT1 variants is relatively limited. Unlike KRAS, RIT1 59 mutations are rarely observed near the critical glycine residues involved in GTP hydrolysis (e.g. 60 G12 and G13 in KRAS or G30 and G31 in RIT1). Instead, *RIT1* mutations occur most frequently near the switch II domain, also targeted by Q61 KRAS variants (Fig. 1a). Nevertheless, these 61 mutations may enhance GTP-bound levels of RIT1^{11,12}. The molecular consequences of RIT1 62 63 switch II domain mutations may additionally be linked to the loss of RIT1's physical interaction with LZTR1, a ubiguitin-conjugating enzyme responsible for degradation of RIT1¹¹. Cancer- and 64 65 Noonan-associated RIT1 variants lose the ability to interact with LZTR1 and consequently are highly overexpressed, resulting in increased signaling activity through the RAF/MEK pathway¹¹. 66

68 Prior studies of RIT1 function focus on candidate cellular signaling pathways based on RIT1's 69 homology to KRAS. To our knowledge, unbiased mapping of downstream RIT1-regulated 70 pathways has not been performed to date. Here we sought to broadly describe the proteome, 71 phosphoproteome, and transcriptome changes induced by wild-type RIT1 and RIT1^{M90I}, a 72 cancer- and Noonan-associated variant, and to compare these changes to those induced by oncogenic KRAS variants. With a particular interest in the consequences of RIT1^{M901} in lung 73 cancer, we profiled the effects of RIT1^{M901} mutation in AALE cells, a non-transformed, 74 75 immortalized, human lung epithelial cell line¹³. 76 77 By comparing the downstream pathways regulated by oncogenic KRAS and RIT1, we uncover 78 previously unknown consequences of RIT1 activation, such as induction of the epithelial-to-79 mesenchymal transition (EMT) and post-translational regulation of HLA protein expression. In 80 addition, we uncover additional functional differences between KRAS and RIT1 including a 81 distinct and unique role of KRAS mutants in regulation of EGFR and USO1 phosphorylation. These data provide the first systems-level view of RIT1 and RIT1^{M901} function. 82 83 **Results** 84

85

Multi-omic profiling of RIT1- and RAS-transformed human lung epithelial cells
We previously demonstrated that RIT1^{M901} and other cancer-associated RIT1 variants can
promote cellular transformation of NIH3T3 cells in vitro and in vivo⁵. To determine whether
RIT1^{M901} was capable of transforming human lung epithelial cells, we expressed mutant RIT1 or
KRAS in the human lung epithelial cell line, AALE. Similar to our previous findings in rodent
cells, both RIT1^{M901} and KRAS^{G12V} enabled AALE cells to form colonies in soft agar (Fig. 1b).

93	The canonical function of oncogenic RAS variants is the downstream activation of the
94	RAF/MEK/ERK cellular signaling cascade ¹⁴ , and RIT1 shares the ability to bind C-RAF and
95	induce transcription of ERK target genes activity ¹¹ . To determine if such regulation is active in
96	AALE cells, we stably expressed wild-type RIT1 or KRAS, or the mutant forms RIT1 ^{M90I} ,
97	KRAS ^{G12V} , and KRAS ^{Q61H} in AALE cells. KRAS ^{Q61H} was included since this mutant more closely
98	resembles the switch II domain mutants observed in <i>RIT1</i> in cancer (Fig. 1a). RIT1 ^{M90I} ,
99	KRAS ^{G12V} , and KRAS ^{Q61H} all enhanced ERK phosphorylation compared to their respective wild-
100	type protein or vector control (Fig. 1c). Interestingly, wild-type RIT1 overexpression also
101	modestly enhanced ERK phosphorylation whereas wild-type KRAS suppressed basal ERK
102	phosphorylation.
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104	To systematically characterize the signaling networks perturbed by mutant RIT1 and KRAS in
105	lung cancer, we expressed each variant in AALE cells and performed both RNA-seq and deep
106	proteome and phosphoproteome profiling by liquid chromatography tandem mass spectrometry
107	(LC-MS/MS). Following trypsin digestion, peptides were labeled with tandem mass tag (TMT)
108	reagents in two overlapping 10-plex sets for relative quantification of proteome and
109	phosphopeptides by LC-MS/MS (Fig. 1d and Supplementary Fig. 1a-b). Following basic
110	reverse phase chromatography, fractions were either directly subjected to LC-MS/MS for total
111	proteome quantification, or subjected to immobilized metal affinity chromatography (IMAC) to
112	enrich for phosphopeptides and then subjected to LC-MS/MS, or. In total, we identified 10,131
113	proteins, 9002 of which were detected and quantified in every sample, and 29,140
114	phosphopeptides, 12,325 of which were identified in common in every sample (Supplementary
115	Tables 1 and 2 and Supplementary Files 1 and 2).
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117 In parallel, we generated deep transcriptome profiles of the same isogenic cell lines. 118 Transcriptome profiling was performed in triplicate on the Illumina NovaSeg platform to a 119 median read-depth per replicate of 70.1 million reads (Fig. 1e, Supplementary Table 3 and 120 Supplementary Fig. 1e). No compensatory feedback regulation of RIT1 to KRAS or vice versa 121 was observed (Fig. 1f). Despite relatively low protein expression of KRAS variants in the AALE 122 lines (Fig. 1c), the majority of KRAS transcripts in each isogenic cell line corresponded to G12V 123 or Q61H variants, respectively, with 84.1% of reads harboring the G12V variant in KRAS^{G12V} 124 cells, and 73.3% of reads corresponding to the Q61H allele in KRAS^{Q61H} cells (Fig. 1g). As 125 expected, known KRAS-regulated gene sets were strongly up- and down-regulated in KRAS-126 mutant cells (Supplementary Fig. 1d). 127 Multi-omic profiling identifies global similarity between signaling regulated by RIT1^{M901} 128 129 and oncogenic KRAS 130 Differentially abundant proteins were identified by comparison to the vector control cells using a 131 two-tailed moderated t-test (Fig. 2a). Selected proteins observed to be significantly modulated 132 by LC-MS/MS were cross-validated by Western blot. FOSL1, also known as FRA1, is a basic leucine zipper transcription factor in the FOS family¹⁵. Activation of RAS is known to promote 133 transcriptional upregulation and protein stabilization of FOSL1^{16,17}. By LC-MS/MS, FOSL1 was 134 markedly overexpressed in KRAS^{G12V}, KRAS^{Q61H}, and RIT1^{M90I}-mutant cells compared to wild-135 136 type cells or vector control cells (Fig. 2b). Consistently, Western blot of independently-derived 137 AALE isogenic lines demonstrated greater abundance of FOSL1 in KRAS- or RIT1-mutant cells 138 compared to wild-type expressing cells (Fig. 2b). TXNIP is an inhibitor of thioredoxin involved in both redox regulation and glucose metabolism^{18,19}. Prior literature identified HRAS^{G12V}-induced 139 suppression of TXNIP transcription and protein translation^{20,21}. TXNIP was among the top down-140 141 regulated proteins in KRAS- and RIT1-mutant proteomes, and was decreased in Western blot

analysis of independently derived cells (Fig. 2c). These validation data demonstrate the utility of
 LC-MS/MS to describe protein expression changes and additionally suggest the mechanism of
 RAS-mediated modulation of FOSL1 and TXNIP is shared with RIT1^{M90I}.

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146 Next we compared the global effects of RIT1^{WT} and RIT1^{M901} to that of KRAS^{WT} and KRAS

147 variants. Proteome and phosphoproteome data from RIT1^{M90I}-expressing cells were highly

148 correlated with KRAS^{G12V} and KRAS^{Q61H} profiles, suggesting largely similar downstream

149 consequences (*r* = 0.70-0.80 and 0.72-0.75 for proteome and phosphoproteome, respectively;

150 **Fig. 2d**). Despite differences in KRAS protein abundance, KRAS^{G12V} and KRAS^{Q61H} proteomes

and phosphoproteomes were highly correlated (proteome r = 0.85 and phospho r = 0.79; Fig.

152 **2d**). In contrast, wild-type KRAS replicates were the most divergent of all profiles, showing

153 limited correlation to either the KRAS-mutant profiles or RIT1 profiles.

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155 A recent study found that RIT1 variants, including M90I, may function by relieving negative 156 regulation of RIT1 by a LZTR1-dependent proteasomal degradation mechanism¹¹. Accordingly, overexpression of wild-type RIT1 should largely phenocopy expression of RIT1^{M901}. Consistent 157 with this idea, RIT1^{WT} cells more closely resembled both RIT1^{M90I} and KRAS-mutant cells than 158 KRAS^{WT} cells (**Fig. 2d**). These data highlight a critical divergence between KRAS and RIT1: 159 160 expression of wild-type KRAS is not capable of activating downstream oncogenic pathways. 161 whereas expression of wild-type RIT1 in part resembles activation of RIT1 or KRAS by 162 mutation. We confirmed this observation in a principal component analysis of transcriptome data, which further revealed a high degree of similarity between RIT1^{WT} and RIT1^{M901}-regulated 163 164 gene expression (Fig. 2e and Supplementary Table 4).

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166

167 Oncogenic RIT1 promotes epithelial-to-mesenchymal transition

168 To identify the downstream pathways regulated by oncogenic KRAS and RIT1, we performed gene set overlap analysis using MSigDB Hallmark Pathway gene sets²² (**Fig. 3a**). The epithelial-169 170 to-mesenchymal transition (EMT) pathway was the most significant gene set enriched among up-regulated proteins for both KRAS^{G12V}/KRAS^{Q61H} and RIT1^{WT}/RIT1^{M90I} cell lines. EMT is a 171 172 cellular transdifferentiation process promoted by cell-extrinsic signaling proteins and 173 orchestrated by activation of transcription factors such as Twist. Snail. and Zeb family 174 transcription factors²³. It has long been observed that oncogenic RAS proteins, including KRAS mutants, promote EMT. An EMT-signature is associated with KRAS dependence²⁴, which has 175 been functionally linked to activation of FOSL1²⁵. Interestingly, we find both RIT1^{M90I} and 176 177 KRAS^{G12V}/KRAS^{Q61H} are capable of promoting expression changes of key EMT markers, 178 including up-regulation of Vimentin, N-Cadherin, and Fibronectin, and downregulation of Keratin 179 19 (Fig. 3b and Supplementary Fig. 2a). Although canonical EMT transcription factors Snail 180 (SNA1) and Slug (SNA2) were not detected by proteomic analysis, transcriptomes from RIT1-181 and KRAS-mutant cells showed increased activity of these EMT transcription factors as 182 determined by ChEA3 transcription factor enrichment analysis (Fig. 3c-d and Supplementary 183 Fig. 2b). To our knowledge, this is the first demonstration of mutant RIT1 promoting EMT in any 184 cell type.

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Oncogenic KRAS and RIT1 suppress Class I MHC expression via a post-transcriptional mechanism

Among the top suppressed proteins with differential abundance in both mutant KRAS and
 RIT1^{M90I} cells, were major histocompatibility complex (MHC) proteins. Class I MHC proteins
 HLA-A, HLA-B, HLA-C, and HLA-F were potently suppressed by KRAS^{G12V}, KRAS^{Q61H}, and
 RIT1^{M90I} (**Fig. 4a-b and Supplementary Fig. 3a**). Recently there has been a renewed interest

in expression of immune modulatory proteins in cancer due to the appreciation of the potent role
of the immune system in shaping cancer evolution. Further understanding the regulation of HLA
expression in cancer is particularly critical in metastatic *KRAS*-mutant lung adenocarcinoma,
where chemotherapy combined with immune checkpoint blockade is often used in the first-line
setting.

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Class I MHC genes *HLA-A*, *HLA-B*, and *HLA-C* harbor loss-of-function mutations in cancer²⁶,
demonstrating selective pressure to lose MHC function during tumorigenesis. Both MHC
expression loss and upregulation of the immune suppressive protein PD-L1 enable tumor
evasion of T-cell recognition of aberrant cancer cell proteins²⁷. Moreover, expression loss of
HLA proteins or B2M, another MHC Class I complex protein, is associated with resistance to
immunotherapy in cancer²⁸. We found that RIT1^{M90I}, KRAS^{G12V}, and KRAS^{Q61H} cells all promoted
loss of B2M protein abundance in addition to HLA protein loss (**Fig. 4c**).

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206 Class I MHC expression is known to be dynamically regulated by upstream signals controlled by 207 interferon gamma exposure, NF-kB signaling, and chromatin regulators such as EZH2^{29,30}. Each 208 of these mechanisms involves transcriptional regulation of class I MHC genes. However, there 209 were no transcriptional differences in HLA genes in the KRAS-mutant and RIT1-mutant cells nor 210 were any transcriptional differences observed in the upstream regulators of MHC Class I 211 expression NLRC5 and IRF1 and IRF2 (Supplementary Fig. 3b). Moreover, we excluded the 212 possibility that lentiviral transduction or expression of a foreign antigen was responsible for the HLA suppression, because HLA protein expression was maintained or enhanced in RIT1^{WT}-213 214 expressing cells as well as vector control cells, which express the Renilla luciferase gene. 215

To identify the possible mechanism of RIT1^{M90I}- and KRAS-mediated MHC suppression, we 216 identified other proteins that, like HLA, were upregulated in RIT1^{WT} cells but suppressed in 217 218 RIT1-mutant and KRAS-mutant cells (Fig. 4d). This analysis revealed the pervasive 219 downregulation of the Rab-mediated ER/Golgi vesicle-trafficking pathway that controls MHC 220 Class I processing and presentation as well as the MHC Class I complex proteins themselves 221 (Fig. 4e). In addition, expression of the proteasomal subunit PSMB9 correlated with loss of the 222 MHC processing machinery (Fig. 4f and Supplementary Fig. 3c). Loss of PSMB9, also known 223 as LMP2, has been previously linked to loss of MHC expression after oncogenic transformation³¹. We conclude that RIT1^{M90I} and KRAS^{G12V}/KRAS^{Q61H} suppress MHC Class I 224 225 expression through a post-transcriptional mechanism possibly involving PSMB9. Further 226 investigation of MHC Class I expression loss driven by these oncogenic RIT1 and KRAS is 227 critical to better understand the role of RAS and RIT1 signaling on immune evasion in cancer. 228

229 The identification of a major class of proteins regulated at the post-transcriptional level in RIT1-230 and KRAS-transformed lung epithelial cells brought to our attention the possibility of other post-231 transcriptional regulation by RIT1 and KRAS. Indeed, oncogenic RAS signaling profoundly alters cap-dependent translation via activation of the p90 ribosomal S6 kinases (RSKs)³² and 232 PI3K/mTOR³³, so differential protein translation could significantly contribute to altered protein 233 234 abundance in RAS-transformed cells. To determine whether there were other protein classes in 235 addition to MHC Class I proteins with significant post-transcriptional regulation, we performed a 236 global correlation analysis of the transcriptome and proteome. Significant linear correlations 237 between transcript and protein abundance were observed for RIT1 and KRAS variants, with the correlation highest for cells expressing mutant KRAS^{G12V} (r = 0.3725) or KRAS^{Q61H} (r = 0.3620) 238 239 (Fig. 4g). While expression of the majority of genes were correlated at the RNA and protein 240 levels, the metallothionein protein family including MT1E, MT1F and MT1X was highly

upregulated in the proteome but not transcriptome of KRAS-mutant cells (Fig. 4g). In addition,
intermediate filament proteins were also substantially regulated post-transcriptionally; both
alpha-internexin (INA) and vimentin (VIM) were expressed more highly in the proteome than
expected from RNA-seq data (Fig. 4g). These data highlight the utility of LC-MS/MS to identify
protein abundance changes that would not be predicted from transcriptome analysis.

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Phosphoproteome profiling illuminates shared and unique signaling by RIT1 and KRAS 247 248 Protein phosphorylation is a reversible and dynamic mechanism of intracellular signaling that 249 enables rapid intracellular transduction of signals controlling cell proliferation, survival, and 250 metabolism. Although both RIT1 and KRAS act as GTPase switches, they both stimulate 251 activation of cellular protein kinases such as BRAF. We therefore evaluated protein 252 phosphorylation regulated by wild-type and mutant RIT1 and KRAS. Phosphosite abundance 253 was expressed as a relative abundance normalized to the total protein abundance for each 254 phosphoprotein. Unsupervised hierarchical clustering of the phospho-signatures identified the 255 RIT1^{M901} phosphoproteome as most similar to KRAS^{G12V} and KRAS^{Q61H} phospho-signatures 256 (Supplementary Fig. 4a). We performed Kinase-Substrate Enrichment Analysis (KSEA³⁴), which uses kinase-substrate pairings from PhosphoSitePlus³⁵ and NetworKIN³⁶ to identify 257 258 differential phosphorylation of kinase-substrate families (**Supplementary Table 5**). These data 259 further confirmed the similarity in phosphorylation state between RIT1-mutant and KRAS-mutant 260 cells. The top kinases with increased substrate phosphorylation in RIT1-mutant and KRAS-261 mutant cells were ribosomal S6 kinase (RPS6KA1), Protein kinase C (PRKCA), AKT1, and 262 MAPKAPK2 (Fig. 5a-c, Supplementary Fig. 4b-e, and Supplementary Table 5). The levels of 263 phosphorylation of RPS6KA1 and MAPKAPK2 substrates were enhanced most strongly in the 264 mutant cells and less in RIT1 WT and KRAS WT-expressing cells (Fig. 5b-c). Substrates of 265 Aurora kinase B and CDK1 and PAK1 were suppressed in RIT1- and KRAS-mutant cells (Fig.

266 **5a** and **Supplementary Fig. 4b-c and Supplementary Fig. 4f**). Although the total

phosphorylation of each substrate reflects the balance between kinases and phosphatases in
the cell, these data suggest that RIT1^{M90I}, like oncogenic KRAS, can activate the canonical RAS
effector pathways involving S6 kinase and AKT.

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Next we assessed the divergent functions of RIT1^{M90I} and KRAS^{G12V}/KRAS^{Q61H} by identifying 271 272 proteins with differential phosphorylation in KRAS-mutant versus RIT1^{M901}-mutant cells. 902 273 differentially phosphorylated sites were identified by two-tailed t-test and multiple hypothesis 274 correction (Fig. 5d; FDR < 0.05). Interestingly, the top site with lower phosphorylation in KRAS^{G12V} and KRAS^{Q61H} cells was EGFR serine 1026 (Fig. 5d). In lung adenocarcinoma, KRAS 275 276 mutations and EGFR mutations are mutually exclusive, suggesting a powerful genetic 277 interaction between these two genes. Recent work demonstrated that mutant KRAS and EGFR display synthetic lethality³⁷. However the mechanism underlying this lethality is unknown. 278 279 Further inspection of the phospho-proteome signatures revealed extensive alteration of EGFR 280 phosphorylation by KRAS^{G12V} and KRAS^{Q61H}, but not by RIT1^{M90I}. 11 of 12 EGFR sites detected 281 by LC-MS/MS occur in the cytoplasmic carboxy-terminal tail of EGFR (Fig. 5e). Five of these 282 sites (S991, S991/T993 double phosphorylation, S1026, S1039, and T1041/S1045 double phosphorylation) were significantly depleted of phosphorylation in KRAS^{G12V} and KRAS^{Q61H}-283 expressing cells but not in RIT1^{M901}-expressing cells. Interestingly, these sites lie in a region of 284 285 EGFR that is involved in receptor internalization and endocytosis³⁸ and a phosphorylationdeficient mutant at S991 is defective at internalization³⁹. Consistently, EGFR protein abundance 286 287 was increased in KRAS-mutant cells (Supplementary Fig. 4g) Although the specific regulatory 288 mechanisms leading to this depletion remain unknown, these data point to the existence of 289 feedback regulatory signaling from oncogenic KRAS to EGFR.

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Examining phosphorylation uniquely promoted by KRAS^{G12V} and KRAS^{Q61H}, we identified USO1 291 292 phosphorylation at S48 as one of the top most significantly increased phosphorylation events in 293 KRAS-mutant cells. USO1, also known as p115, is a vesicle tethering factor involved in ER-294 Golgi intracellular trafficking⁴⁰. Although wild-type KRAS and KRAS-mutant proteomic 295 signatures were largely divergent, USO1 serine 48 phosphorylation was promoted by both 296 KRAS^{WT} and mutant KRAS (Fig. 5f). KRAS relies on vesicle trafficking to ensure proper post-297 translational farnesylation and palmitovlation, which are required for targeting of KRAS to the 298 plasma membrane⁴¹. We hypothesized that USO1 S48 phosphorylation was therefore 299 correlated with KRAS expression rather than activity. Indeed, a significant correlation was 300 observed between overall KRAS expression and USO1 phosphorylation (Fig. 5g). In contrast, 301 USO1 S48 phosphorylation was only modestly changed in RIT1-mutant cells (Fig. 5d). Notably, RIT1 lacks the farnesylation and palmitovlation signals present in RAS isoforms⁴², so the 302 303 differential regulation of USO1 by KRAS and RIT1 may be related to differences in RIT1 and 304 KRAS trafficking.

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306 Also among the top differentially phosphorylated sites were 32 phosphorylation sites in AHNAK 307 proteins 1 and 2. AHNAK and AHNAK2 are large scaffolding proteins that have been implicated as tumor suppressor proteins in breast and lung cancer^{43,44}. Among all phospho-proteins, a 308 309 higher proportion (32/117) of sites on AHNAK and AHNAK2 were differentially phosphorylated 310 than expected by chance (P < 0.0001 by Chi Square test; **Supplementary Fig. 4h**). Intriguingly, 311 two recent proximity-labeling proteomic studies identified AHNAK and AHNAK2 as KRAS-312 interacting proteins^{45,46}, raising the possibility that a direct physical interaction between KRAS 313 and AHNAK proteins may be involved in the differential AHNAK phosphorylation we observe.

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315 **Discussion**

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317 Here we describe quantitative proteomic, phosphoproteomic, and transcriptomic datasets that provide the first systematic view of the RIT1^{M901}-regulated signaling network. These datasets 318 319 were generated from isogenic human lung epithelial cells to provide a physiological view of the 320 consequences of RIT1 activation in the same cellular compartment that is involved in lung 321 adenocarcinoma, a tumor type with prevalent mutations in KRAS and RIT1. Broadly, we find that 'omic signatures from RIT1^{M90I}-expressing cells largely phenocopy those from cells with 322 323 overexpression of wild-type RIT1. This finding lends further support to the notion that oncogenic RIT1 variants function at least in part through increasing RIT1 abundance¹¹. This is in contrast 324 325 to KRAS, where overexpression of wild-type KRAS induces signatures unrelated or opposite to 326 that of oncogenic KRAS variants G12V and Q61H. The opposing functions of wild-type and 327 mutant KRAS is consistent with recent evidence suggesting that KRAS functions as a dimer and 328 that wild-type KRAS directly inhibits the function of oncogenic KRAS variants via physical 329 dimerization⁴⁷. This divergence in the function of wild-type RIT1 and KRAS hints at fundamental 330 differences in molecular regulation of each wild-type GTPase. The ability of RIT1 to promote 331 downstream RAF/MEK/ERK signaling when aberrantly expressed suggests that RIT1 may not 332 be subject to the same tight regulation by GTPase-activating proteins (GAPs) that normally 333 keep RAS in an inactive state. Furthermore, these data raise the possibility that wild-type RIT1 334 overexpression in *RIT1*-amplified cancers may contribute to tumorigenesis. *RIT1*, on 335 chromosome 1q, is frequently amplified in uterine carcinosarcoma, liver hepatocellular cancer, 336 cholangiocarcinoma, breast cancer, lung adenocarcinoma, and ovarian cancer. RIT1 mRNA 337 expression is increased in amplified cases, regardless of tissue type, raising the possibility that 338 RIT1 overexpression could play a role in tumorigenesis in these cancers.

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We find that RIT1^{M90I}, KRAS^{G12V}, and KRAS^{Q61H} share the ability to activate canonical RAS 340 341 effector pathways PI3K/AKT and RAF/MEK. Likely as a consequence of RAF/MEK signaling to FOSL1, RIT1^{M901} also shares the ability to induce EMT markers including Vimentin, N-cadherin, 342 343 and fibronectin. KRAS and RIT1 variants also shared the ability to profoundly suppress HLA-A, -344 B, and -C expression via a posttranscriptional mechanism. Taking advantage of differential expression of HLA proteins between RIT1^{WT} and all other isogenic lines, we identified an entire 345 346 Rab-mediated endocytic network that was lost together with HLA proteins in RIT1- and KRAS-347 mutant cells. This downregulated module also included PSMB9, a subunit of the 348 immunoproteasome that is involved in antigen processing for class I MHC presentation. RAS oncogenes have long been recognized to suppress surface MHC expression⁴⁸, in some cases 349 transcriptionally and in others post-transcriptionally³¹. Our data link both oncogenic RIT1 and 350 351 RAS to modulation of the processing and trafficking of MHC Class I molecules. Further 352 identification of the mechanism of RIT1/RAS-mediated MHC suppression will provide a better 353 understanding of tumor immune evasion which is critically needed to optimize patient 354 stratification of cancer immunotherapy. 355 356 In addition to the largely concordant regulation of proteins by mutant RIT1 and KRAS, we identified several unique phosphoproteins with differential abundance in RIT1^{M901} and KRAS-357 358 mutant cells. These included EGFR, a key oncoprotein in lung adenocarcinoma, which showed 359 reduced phosphorylation of sites involved in receptor internalization and endocytic trafficking. 360 Given the potent genetic interactions between KRAS and EGFR in lung cancer and colon 361 cancer, it is attractive to speculate that feedback regulation of KRAS to EGFR could provide an 362 explanatory mechanism for this phenomenon. Future work is needed to determine the basis of 363 the specific regulation of EGFR phosphorylation by oncogenic KRAS but not RIT1.

364	Together, these results demonstrate the power of quantitative proteomics and transcriptomics to
365	provide global views of cancer oncogene signaling. Our multi-omic analysis validated known
366	consequences of RAS activation such as EMT and activation of RAF/MEK and PI3K signaling.
367	For the first time, we gained a global view of RIT1 function, which confirmed its ability to
368	stimulate canonical RAS signaling. However, phosphoproteomic profiling identified a number of
369	key divergent mechanisms between KRAS- and RIT1-mutant cells, which point to the existence
370	of novel, unique regulators or effectors of KRAS and RIT1 still to be identified. Future work is
371	needed to investigate the mechanisms of these differences between KRAS and RIT1, the
372	results of which will have important implications for cancer therapy and Noonan Syndrome.
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378 Methods

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380 Isogenic Cell Line Generation

381 Plasmid constructs were cloned using Gateway Technology (Invitrogen/ThermoFisher) using 382 pLX301 destination vector (Broad Institute) and pDONR223-RIT1 donor vectors previously 383 described⁵. Lentivirus was generated by transfection of HEK293T cells with packaging and 384 envelope vectors using standard protocols. AALE cells were a kind gift of Jesse Boehm (Broad 385 Institute). Isogenic cells were generated by transduction of lentivirus generated from pLX317-Renilla luciferase or pLX301-RIT1^{WT}, pLX301-RIT1^{M90I}, pLX301-KRAS^{WT}, pLX301-KRAS^{G12V}, or 386 pLX301-KRAS^{Q61H} and selection with puromycin. Stable pools of cells were maintained in small 387 388 airway growth medium (Lonza).

389

390 Soft Agar Assay

 1×10^5 cells were suspended in 1 ml of 0.33% select agar in small airway growth medium without EGF (Lonza) and plated on a bottom layer of 0.5% select agar in the same media in six-well dishes. Each cell line was analyzed in triplicate. Colonies were photographed after 14–21 days and quantified using CellProfiler⁴⁹.

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396 Transcriptome profiling

Three technical replicates per cell line were harvested at ~90% confluence (n = 18 total dishes). Cells were lysed and total RNA was extracted using Direct-zol RNA Miniprep plus (Zymo Research). Libraries were constructed using the non-strand-specific poly-A selection Illumina TruSeq kit for 50bp paired-end reads. Libraries were then pooled and sequenced on the Illumina NovaSeq platform (Fred Hutch Genomics Core). Reads were aligned to the human reference genome build hg19/GRCh37 using STAR v.2.5.3a⁵⁰. Alignments were annotated for duplicates and read groups, and then reordered and indexed, using Picard Tools v.1.114⁵¹.

Read statistics for each RNA-seq sample were calculated using RSeQC⁵². Quantification of
gene transcripts was performed by the featureCounts program within the Subread package⁵³,
using hg19 gene annotation from UCSC Gene level CPM and RPKM values were calculated
with edgeR v.3.22.3⁵⁴, and converted into transcripts per million (TPM values with an in-house
script. In total, 12,887 genes were identified with average logCPM at least 0.1 across all
samples. Differential expression analyses comparing KRAS or RIT1 perturbed cell lines against
vector control lines were performed using edgeR⁵⁴.

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412 High performance liquid chromatography tandem mass spectrometry (LC-MS/MS)

413 Cells were washed in ice-cold PBS, scraped into PBS, pelleted, and snap frozen in liquid 414 nitrogen. The experimental workflow for sample processing, TMT-labeling, peptide enrichment, and LC-MS/MS were largely as previously described⁵⁵. Briefly, pellets were lysed in 200 µl of 415 416 chilled urea lysis buffer (8 M urea, 75 mM NaCl, 50 mM Tris (pH 8.0), 1 mM EDTA, 2 µg/ml 417 aprotinin, 10 µg/ml leupeptin, 1 mM PMSF, 1:100 (vol/vol) Phosphatase Inhibitor Cocktail 2, 418 1:100 Phosphatase Inhibitor Cocktail 3, 10 mM NaF, and 20 µM PUGNAc) for each ~50 mg 419 portion of wet-weight tissue. Lysates were reduced with 5mM DTT, alkylated with 10 mM IAM, 420 and digestion performed in solution with 1 mAU LysC per 50 µg of total protein and trypsin at an 421 enzyme/substrate ratio of 1:49. Reactions were quenched with FA and brought to pH = 3 with 422 FA. Peptides were desalted on 200 mg tC18 SepPak cartridges and dried by vacuum 423 centrifugation. 340 µg of peptides were labeled with 10-plex Tandem Mass Tag reagents 424 (TMT10, Fisher Scientific), according to manufacturer's instructions. To enable quantification of 425 peptides across all 12 samples, the samples were labeled in sets of 10 across two different 426 TMT10 pools in a crossover design with 8 of 12 samples analyzed in both TMT10 pools. A 427 50/50 mix of both AALE vector control lysates was used as an internal reference in both TMT10 428 runs (Supplementary Fig. 1b).

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430	Each TMT10-plex was desalted in a 200 mg tC18 SepPak cartridge and fractionated using
431	offline HPLC. 5% of each fraction was collected into an HPLC vial for proteome analysis by LC-
432	MS/MS. The remaining 95% was processed for phospho-peptide enrichment via immobilized
433	metal affinity chromatography (IMAC). IMAC enrichment was performed using Ni-NTA
434	Superflow Agarose beads incubated with peptides solubilized in a final concentration of 80%
435	MeCN/0.1% TFA. Phospho-enriched peptides were desalted and collected into an HPLC vial for
436	analysis by LC-MS/MS.
437	
438	Online fractionation was performed using a nanoflow Proxeon EASY-nLC 1200 UHPLC system
439	(Thermo Fisher Scientific) and separated peptides were analyzed on a benchtop Orbitrap Q
440	Exactive Plus mass spectrometer (Thermo Fisher Scientific) equipped with a nanoflow
441	ionization source (James A. Hill Instrument Services, Arlington, MA). In-house packed columns

442 (20 cm x 75 μm diameter C18 silica picofrit capillary column; 1.9 μm Reprosll-Pur C18-AQ

443 beads, Dr. Maisch GmbH, r119.aq; Picofrit 10 μm tip opening, New Objective, PF360-75-10-N-

(Solvent A) and 90 % acetonitrile /0.1 % formic acid (Solvent B). The 110 min LC-MS/MS

5). Mobile phase flow rate was 200 nL/min, comprised of 3 % acetonitrile/0.1 % formic acid

446 method consisted of a 10 min column-equilibration procedure; a 20 min sample-loading

447 procedure; and the following gradient profile: (min: % B) 0:2; 2:6; 85:30; 94:60; 95:90; 100:90;

448 101:50; 110:50 (the last two steps at 500 nL/min flow rate). Data-dependent acquisition was

performed using Xcalibur QExactive v2.4 software in positive ion mode at a spray voltage of
2.00 kV. MS1 Spectra were measured with a resolution of 70,000, an AGC target of 3e6 and a
mass range from 300 to 1800 m/z. Up to 12 MS/MS spectra per duty cycle were triggered at a

resolution of 35,000, an AGC target of 5e4, an isolation window of 0.7 m/z, a maximum ion time

453 of 120 msec, and normalized collision energy of 30. Peptides that triggered MS/MS scans were

dynamically excluded from further MS/MS scans for 20 sec. Charge state screening was
enabled to reject precursor charge states that were unassigned, 1, or >6. Peptide match was
set to preferred for monoisotopic precursor mass assignment.

457

458 **Protein-peptide identification, phosphosite localization, and quantification**

459 MS data was interpreted using the Spectrum Mill software package v6.0 pre-release (Agilent 460 Technologies, Santa Clara, CA. MS/MS spectra were merged if they were acquired within +/- 45 461 sec of each other with the same precursor m/z. Also, MS/MS spectra that did not have a 462 sequence tag length > 0 (i.e., minimum of two masses separated by the in chain mass of an 463 amino acid) or did not have a precursor MH+ in the range of 750-6000 were excluded from 464 searching. MS/MS spectra searches were performed against a concatenated UniProt human 465 reference proteome sequence database containing 58.929 human proteins including isoforms 466 (obtained 10/17/2014) and 150 additional common laboratory contaminants. ESI-QEXACTIVE-467 HCD-3 scoring parameters were used for both whole proteome and phosphoproteome datasets. 468 Spectra were allowed +/- 20 ppm mass tolerance for precursor as well as product ions, 30% 469 minimum matched peak intensity, and "trypsin allow P" was set as enzyme specificity with up to 470 4 missed cleavages allowed. Carbamidomethylation at cysteine was set as fixed modification 471 together with TMT10 isobaric labels at lysine residues (N-termini would be considered 472 regardless if it was TMT labelled). Acetylation of protein N-termini andoxidized methionine were 473 set as variable modifications with a precursor MH+ shift range of -18 to 64 Da for the proteome 474 searches. For the phosphoproteome searches the precursor MH+ shift range was set to 0 to 475 272 Da and variable modifications of phosphorylation of serine, threonine, and tyrosine. 476 Identities interpreted for individual spectra were automatically designated as confidently 477 assigned using the Spectrum Mill autovalidation module to use target-decoy based false 478 discovery rate (FDR) estimates to apply score threshold criteria. For the whole proteome

datasets, thresholding was done at the spectral (< 1.2%) and protein levels (< 0.1%). For the
phosphoproteome datasets, thresholding was done at the spectral (< 1.2%) and phosphosite
levels (< 1.0%).

482 Replicates across TMT-plexes were highly correlated (**Supplementary Fig. 1c)** with median

483 Pearson *r* = 0.87 and 0.69 for proteome and phosphoproteome, respectively. Technical

484 replicates and biological replicates were merged to generate final total proteome and phospho-

485 proteome profiles for each isogenic cell line (**Supplementary Tables 1 and 2**). Replicate-level

486 profiles are also supplied as JavaScript Object Notation (.json) files that can be visualized and

487 analyzed using the Morpheus Matrix Visualization and Analysis Software at

488 <u>https://software.broadinstitute.org/morpheus</u> (Supplementary Files 1 and 2). Differential

489 protein and phospho-site signatures were generated by computing the mean log₂(fold change)

490 of the abundance of each site in each sample compared to the vector control cells. Statistical

491 significance of differentially abundant proteins and phosphosites was determined by performing

a one sample moderated t-test with multiple hypothesis correction (Supplementary Tables 1

493 and 2).

494

495 Integrative Analysis

496 Correlation of changes in protein expression and changes in RNA expression was modeled

497 using R's Im() function. 95% prediction intervals were calculated to determine genes with weak

498 concordance between protein and RNA expression.

499

500 Gene Set Enrichment Analysis

Analysis of enrichment of KRAS signaling in differential RNA expression profiles was performed
in R with the goseq package⁵⁶. KRAS signaling gene sets were taken from MSigDB hallmark
gene sets^{22,57}.

504

505 **Transcription Factor Target Enrichment Analysis** 506 Analysis of over-representation of Transcription Factor targets was performed with ChEA3 by 507 submitting lists of differentially expressed genes (|LFC| > 1 and FDR < 0.05). ChEA3 performs 508 Fisher's Exact Test to compare the input gene set to TF target gene sets in six different 509 libraries⁵⁸. Analysis of the Enrichr Queries library was selected as the focus of the present 510 study. Transcription factors resulting from this analysis were annotated as one of four groups of 511 EMT association. These four groups were the Snail gene family, confirmed EMT genes defined by dbEMT⁵⁹, genes shown to be associated with EMT in at least one study in literature, and 512 513 genes unrelated to EMT. 514 515 Antibodies and immunoblotting 516 Antibodies against FOSL1 (D80B4), TXNIP (D5F3E), and Vimentin (D21H3) were purchased 517 from Cell Signaling Technology. Vinculin (V9264) was purchased from Sigma Aldrich. 518 Secondary antibodies StarBright Blue 700 Goat anti-Rabbit IgG, StarBright Blue 520 Goat anti-519 Rabbit IgG and StarBright Blue 520 Goat anti-Mouse IgG (12005867) were purchased from Bio-520 Rad. Antibody against RIT1 (#53720) was purchased from Abcam. Cell lysates were prepared 521 in RTK lysis buffer with protease (11836153001, Roche) and phosphatase (04906837001, 522 Roche) inhibitors added and quantified by the BCA assay (Thermo Scientific Waltham, MA). 523 Samples were then boiled in Laemmli buffer (1610747, Bio-Rad, Hercules, CA) and 50 ug of 524 protein was loaded onto 4-15% Mini-Protean TGX (4561084, Bio-Rad, Hercules, CA) gels. 525 Protein gels were run and transferred to PVDF membranes (1704274, Bio-Rad, Hercules, CA) 526 according to manufacturer's instructions. Proteins were detected by specific primary antibody 527 and secondary antibody then visualized using the ChemiDoc MP Imaging System (Bio-Rad, 528 Hercules, CA).

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531	KSEA analysis
532	Kinase-substrate enrichment analysis (KSEA) ⁶⁰ was performed using the KSEA App ³⁴
533	(https://casecpb.shinyapps.io/ksea/) using kinase-substrate mappings from PhosphoSitePlus ³⁵
534	and a p value threshold of < 0.05. A minimum of five detected phospho-site substrates were
535	needed for kinases to be included in the analysis. The full list of kinase scores and number of

536 substrates are shown in Supplementary Table 5. 36 kinases had sufficient substrate sites

537 detected to be included in the analysis. Kinase-substrate mappings are shown in

538 Supplementary Table 5.

539

540 DATA AVAILABILITY

The RNA-seq data have been deposited in the NCBI Gene Expression Omnibus database with accession number GSE146479. All mass spectra contributing to this study can be downloaded in the original instrument vendor format from the MassIVE online repository (Accession number to be updated prior to publication.)

545

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554

555 AUTHOR CONTRIBUTIONS

- 556 A.H.B. conceived of and directed the study. S.C. and P.M. supervised the LC-MS/MS
- 557 experiments. F.M. performed the proteomics data analysis, with contributions from
- A.H.B. and K.H. A.L. performed the transcriptome analysis. A.L. and A.H.B. wrote the
- manuscript. A.L., K.H., S.M., I.F., S.F., J.W., and A.H.B. performed experiments. All
- authors discussed results and provided input on the manuscript.

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565 Figure Legends

566 Figure 1. Comparative multi-omic profiling of KRAS- and RIT1-mutant human lung 567 epithelial cells. a, Protein alignment of KRAS-4B (Uniprot #P01116-2) and RIT1 Isoform 1 (Uniprot #Q92963-1) generated by ClustalW2⁶¹. Stars indicate the position of the RIT1^{M901} or 568 KRAS^{G12V} and KRAS^{Q61} variants used in this study. Asterisks indicate fully conserved residues. 569 570 Colons indicate strongly conserved residues. Periods indicate weakly conserved residues. b, 571 Soft agar colony formation assay of isogenic AALE human lung epithelial cells. **, p < 0.01; ****, 572 p<0.0001 by two-tailed t-test. c, Western blot using anti-RAS and anti-RIT1 antibodies (top 573 panels), or antibodies against phosphorylated ERK1/2 or vinculin (loading control). SE = short 574 exposure, LE = long exposure. Isogenic AALE cells were cultured in the presence or absence of 575 EGF for 12 hours. d, LC-MS/MS workflow for generation of proteome and phosphoproteome 576 profiles. bRP, basic reverse phase chromatography. IMAC, immobilized metal affinity 577 chromatography. e, Workflow for Illumina RNA-seg analysis. f, mRNA guantification in 578 transcripts per million (TPM) showing mean ± SD of RIT1 (left) or KRAS (right) in isogenic AALE 579 cells, n = 3 per cell line. *, p < 0.05; ****, p < 0.0001 by two-tailed Student's t-test compared to 580 vector control cells. **g**, RNA-seq quantification of variant allele expression. Data shown is the 581 percentage of reads at the M90I, G12V, or Q61H variant site for the variant allele or wild-type 582 allele.

583 Figure 2. Quantitative proteome and transcriptome profiling identifies similarity in

RIT1^{M901}-mutant and KRAS-mutant signaling networks. a, Volcano plots of global proteome data from isogenic AALE cells showing the log₂(fold change) ("LFC") in protein abundance in each cell line compared to vector control cells. The y-axis displays the negative log₁₀(*p* value) calculated from a one sample moderated t-test with multiple hypothesis correction by the Benjamini-Hochberg method. **b**, Western blot validation of FOSL1 increased protein abundance in RIT1- and KRAS-mutant cells. The chart shows the LFC of FOSL1 as determined by LC-

590 MS/MS. Western blot below shows FOSL1 abundance or Vinculin (loading control). c, Western 591 blot validation of TXNIP protein abundance in RIT1- and KRAS-mutant cells. The chart shows 592 the LFC of TXNIP as determined by LC-MS/MS. d, Correlation heatmap showing pairwise 593 Pearson and Spearman correlations of each proteome and phosphoproteome replicate to every 594 other replicate. To enable correlation of proteome with phosphoproteome, phosphosites were 595 collapsed to the protein level by taking the median of all phosphosites for each protein. e, 596 Principal component analysis (PCA) of RNA-seq data. Circles correspond to control vector or 597 wild-type replicates. Diamonds correspond to RIT1- or KRAS-mutant profiles.

Figure 3. RIT1^{M901} promotes epithelial-to-mesenchymal (EMT) transition. a, Gene set 598 599 overlap analysis of up-regulated ("Up"; LFC>2) and down-regulated ("down"; LFC<-2) proteins using MSigDB Hallmark Pathways²². "K" and "R" indicate analysis based on mean LFC of 600 KRAS^{G12V}/KRAS^{Q61H} cells or RIT1^{WT}/RIT1^{M90I} cells, respectively. Circle size corresponds to the p 601 602 value of gene set overlap analysis determined by MSigDB. b, LFC of protein abundance of EMT 603 marker genes as determined by LC-MS/MS, relative to vector control cells. c, Transcription factor target enrichment analysis of differentially expressed genes in RIT1^{M901}-mutant cells using 604 605 Enrichr libraries. FET, Fisher's exact test. Red = Snail family. Orange = confirmed EMT genes in dbEMT⁵⁹. Pale orange = EMT-associated genes in literature. **d**, Enrichr analysis of KRAS^{G12V}-606 607 mutant proteome data. Annotation is the same as in c.

Figure 4. RIT1- and KRAS-mutant cells suppress Class I MHC expression via global loss
of antigen processing and presenting machinery. a, Rank plot of all protein abundance
changes in KRAS^{G12V}-mutant cells compared to vector control, generated by LC-MS/MS. HLAA,-B,-C, and -F proteins are labeled in blue. b, Heat map showing HLA protein abundance in
each global proteome replicate. Replicates were clustered by unsupervised hierarchical
clustering using all detected proteins. c, Protein abundance of B2M in LC-MS/MS data. d, Top

differentially abundant proteins between RIT1^{WT} cells and all other cell lines. Proteins are 614 615 ranked by the signal-to-noise (S2N) statistic, shown in the bar chart at the right. e, StringDB⁶² 616 network analysis of proteins with S2N>2.5 in analysis shown in d. The network was significantly 617 more connected than expected by chance (p < 1e-16). Disconnected nodes, single connected 618 nodes, and disconnected clusters have been removed from the visualization. Edges represent 619 high confidence interaction scores (>0.9) and network edge thickness indicates the strength of 620 data support from all StringDB active interaction sources. f. Protein abundance of PSMB9 in LC-621 MS/MS data. g, Global proteome-transcriptome correlation analysis. A dashed diagonal line 622 displays the linear regression generated by comparing the LFC of each gene in the 623 transcriptome to its respective protein LFC in the proteome. The resulting Pearson correlation 624 coefficient (r) is shown. Genes outside the 95% prediction interval are plotted in red, and include 625 HLA genes, metallothioneins, and intermediate filament proteins Vimentin (VIM) and alpha 626 internexin (INA).

627 Figure 5. Phosphoproteomic profiling illuminates novel differential post-translational 628 modifications in RIT1^{M901}- and KRAS-mutant cells. a. KSEA analysis of AALE 629 phosphoproteomes. Top differentially phosphorylated kinase substrates are shown. The full 630 KSEA results are shown in **Supplementary Fig 4b-c. b**, Violin plot of phospho-site abundance 631 of phospho-sites that are RPS6KA1 substrates. c, Violin plot of phospho-site abundance of 632 phospho-sites that are MAPKAPK2 substrates. d, Marker selection analysis identifies 633 differentially phosphorylated sites in KRAS-mutant cells compared to RIT1-mutant cells. Phosphosites from KRAS-mutant and RIT1^{M901}-mutant replicate-level phosphoproteome profiles 634 635 (Supplementary File 2) were compared by two-tailed t-test. The top 20 significantly (FDR < 636 0.05) differentially phosphorylated sites in each direction are shown and ranked by t-statistic. A 637 heat map displays the LFC in phosphorylated peptide abundance of each site compared to 638 vector control, after normalizing to total protein abundance. e. LFC of EGFR phosphosites in

639 KRAS-mutant and RIT1-mutant cells. Data shown is the mean + SD of n=8 KRAS-mutant 640 replicates and n=4 RIT1-mutant replicates. *, FDR < 0.01 as determined by two-tailed t-test and 641 two-stage linear step-up procedure of Benjamini, Krieger and Yekutieli. f, Relative 642 phosphorylation of USO1 at serine 48 as determined by LC-MS/MS. Box and whiskers show the 643 25th-75th percentiles and minimum to maximum of the data, respectively. g, Relationship of 644 USO1 S48 phosphorylation to KRAS total protein abundance. A dashed line displays the linear 645 regression fit and grav lines display the 95% confidence interval of the linear model, r = 0.70, p 646 < 0.01.

647

648 SUPPLEMENTARY FIGURE LEGENDS

649 Supplementary Figure 1. Workflow and quality control of proteomic and transcriptomic

650 profiling. a, Replicate-level workflow for tandem mass tag (TMT) labeling and LC-MS/MS.

651 Lysates from duplicate sets of six isogenic cell lines were used to generate two TMT-plex sets,

with control samples used to link the two sets. **b**, TMT 10-plex layout showing mass tags

associated with each replicate. **c**, Average pairwise replicate correlations (Pearson *r*) of all

replicates from each sample group indicated. **d**, Enrichment analysis of differentially expressed

655 genes between KRAS or RIT1 perturbed lines and vector controls using *goseq*⁵⁶. mSigDB

hallmark gene sets specific to KRAS signaling are shown. **e**, RNA-seq run and mapping

657 statistics show total reads, mapped reads, and reads mapped to rRNA, for each sample.

658

659 Supplementary Figure 2. RIT1 and KRAS promote epithelial-to-mesenchymal transition.

660 **a,** Changes in mRNA transcript levels of EMT genes *VIM*, *CDH*2, *FN*1, and *KRT*19, in each

661 isogenic cell line compared to vector control. LFC, log₂(fold-change) compared to vector cells. **b**,

662 Transcription factor target enrichment analysis using Enrichr libraries of differentially expressed

663 genes in RIT1^{WT}, KRAS^{WT}, and KRAS^{Q61H}-mutant cells. FET, Fisher's exact test. Red = Snail

family. Orange = confirmed EMT genes in dbEMT⁵⁹. Pale orange = EMT-associated genes in
literature.

666

Supplementary Figure 3. Post-transcriptional loss of Class I MHC proteins. a, Rank plot of all protein abundance changes in KRAS^{Q61H}-mutant or RIT1^{M90I}-mutant cells compared to vector control, generated by LC-MS/MS. HLA-A,-B,-C, and -F proteins are labeled. LFC, \log_2 foldchange. b, Change in mRNA transcript levels of HLA genes and upstream regulators of MHC Class I, in each isogenic cell line compared to vector controls. LFC, \log_2 fold change compared to vector cells. c, Correlation of protein levels in HLA-A and PSMB9 across each isogenic cell line. A line is the best-fit linear regression with significant non-zero slope (p < 0.05).

675 Supplementary Figure 4. Phosphoproteome profiling identifies enhanced

676 phosphorylation of specific kinase substrates in KRAS- and RIT1-mutant cells. a, Pairwise 677 replicate correlation (Pearson r) heatmap and unsupervised clustering of phosphoproteome data. **b**, KSEA of phosphoproteome data for RIT1^{WT} and RIT1^{M90I}-expressing cells. The kinase 678 679 z-score indicates the overall score for each kinase listed, normalized by the total number of 680 substrates. Significant scores (p<0.05) are indicated in red and blue. Phospho-sites of kinases 681 in red were more highly abundant in the cell line compared to vector control, whereas phospho-682 sites of kinases in blue were more highly abundant in vector control than the indicated cell line. 683 c, KSEA of phosphoproteome data for KRAS-expressing cells. Labeling as in (b). d, Violin plot 684 of phospho-site abundance of AKT1 substrate sites. e, Violin plot of phospho-site abundance of 685 PRKCA substrate sites. f, Violin plot of phospho-site abundance of AURKB substrate sites. g, 686 EGFR protein abundance in LC-MS/MS data compared to vector control. f, Proportion of 687 phosphorylated sites in AHNAK proteins with differential phosphorylation between KRAS-mutant and RIT1^{M901}-mutant cells. Data shown is the percentage of differentially abundant 688

- 689 phosphorylation sites in AHNAK and AHNAK2 compared to all other sites. Significance was
- 690 determined from the analysis in (**b**), FDR < 0.05. ****, p < 0.0001 by two-sided Fisher's exact
- 691 test.

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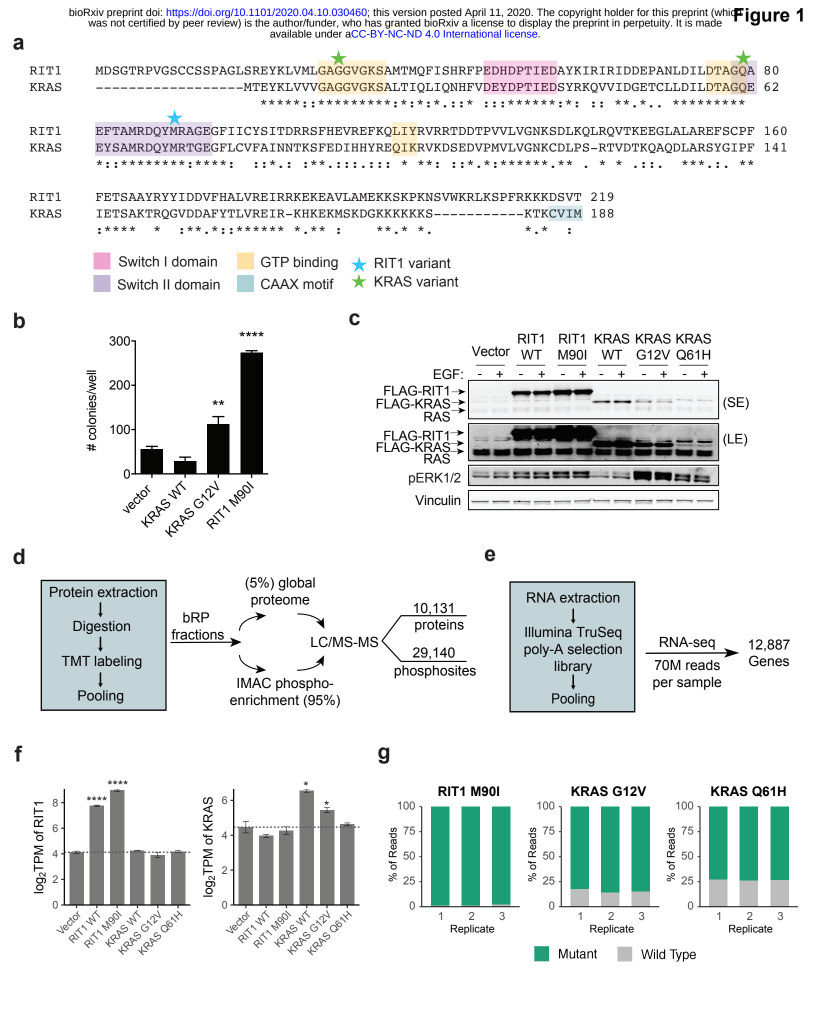
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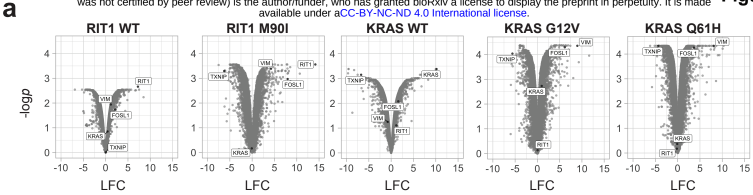
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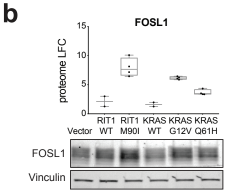
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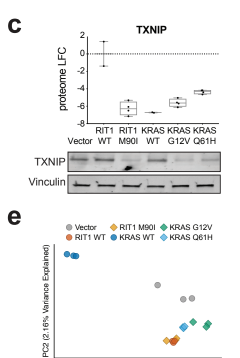
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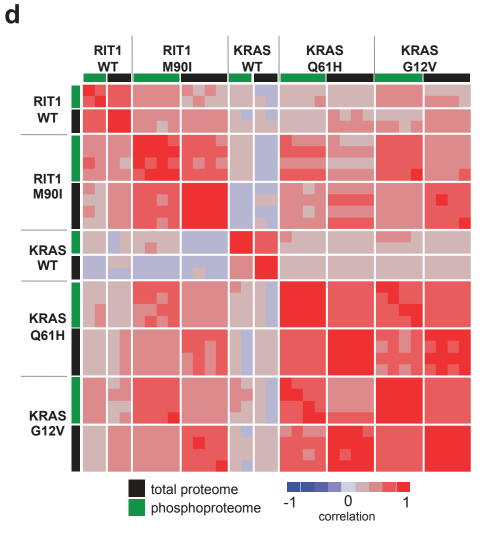
bioRxiv preprint doi: https://doi.org/10.1101/2020.04.10.030460; this version posted April 11, 2020. The copyright holder for this preprint (whi**Figure 2** was not certified by peer review) is the author/funder, who has granted bioRxiv a license to display the preprint in perpetuity. It is made available under aCC-BY-NC-ND 4.0 International license.

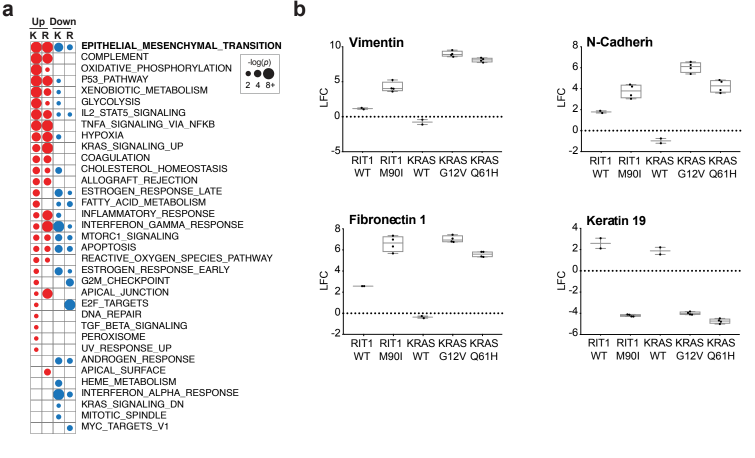






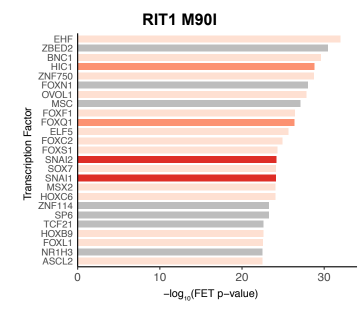
PC1 (96.2% Variance Explained)

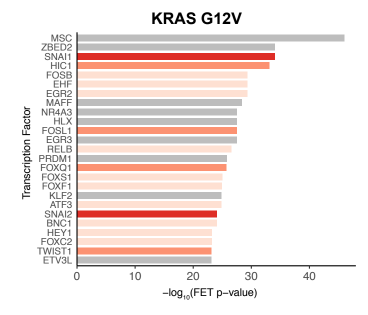


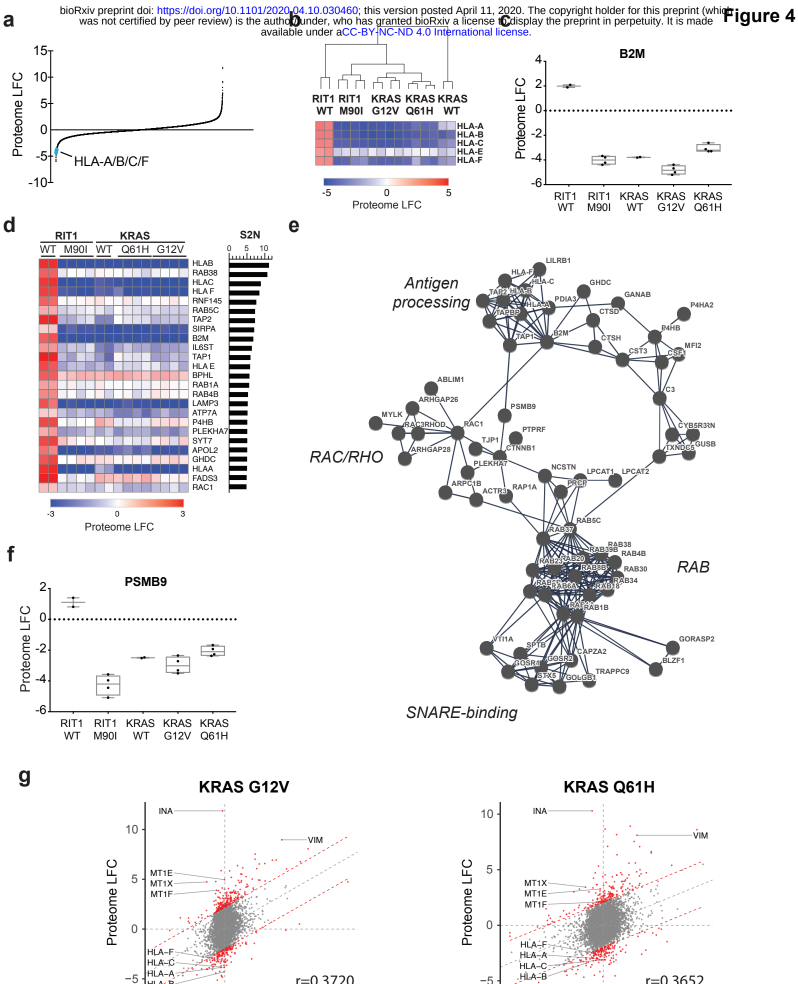


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r=0.3720 HLA-B -2 0 2 4 **RNA LFC**

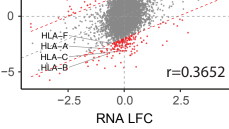


Figure 5

