# Effectors and potential targets selectively upregulated in human KRAS-mutant lung adenocarcinomas

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# **Abstract**

Genetic and proteomic analysis of human tumor samples can provide an important compliment to information obtained from model systems. Here we examined protein and gene expression from the Cancer Genome and Proteome Atlases (TCGA and TCPA) to characterize proteins and protein-coding genes that are selectively upregulated in *KRAS*-mutant lung adenocarcinomas. Phosphoprotein activation of several Raf/MAPK signaling components was considerably stronger in *KRAS*-mutants than any other group of tumors. However, both *KRAS*-mutants and tumors with other activating mutations in the Raf/MAPK pathway showed comparable activation of mTOR. Co-occurring mutations in *KRAS*-mutants were associated with differential activation of PDK1 and PKC-alpha. Genes showing strong activation in RNA-seq data included negative regulators of Raf/MAPK signaling along with potential oncogenic effectors including activators of Rac and Rho proteins and the receptor protein-tyrosine phosphatase gene *PTPRE*. These results corroborate Raf/MAPK signaling as an important therapeutic target in *KRAS*-mutant lung adenocarcinomas and pinpoint new potential targets.

# Introduction

How mutationally activated *KRAS* and other canonical *RAS* genes malignantly transform cells and how to block this process for therapeutic benefit has been a subject of intense investigation for over thirty years. The majority of efforts to date have relied on model systems, using established cell lines and mouse models. These studies have identified signaling pathways that are directly stimulated by biochemically active Ras proteins, including the Raf/MAPK and PI3K/Akt pathways<sup>1</sup>. They also have identified pathways or processes further downstream from Ras proteins that are involved in malignant phenotypes induced by mutant *RAS* genes, including the NF- $\kappa$ B pathway<sup>2, 3</sup>, transcriptional activity of the oncogene  $YAP^4$ , generation of reactive oxygen species<sup>5</sup>, and anabolic glucose metabolism<sup>6</sup>. A third class of proposed oncogenic mediators of mutant *RAS* genes are induced secreted proteins including TGF- $\alpha^7$ , Vegf<sup>8</sup>, IL-8<sup>9</sup>, IL-6<sup>10</sup>, CXCL1<sup>11</sup>, and CCL5<sup>12</sup>. Several members of these three classes of

mediators of oncogenic Ras have been explored as potential therapeutic targets but as of yet there hasn't been a clinically successful treatment developed for cancers with mutant *RAS* genes.

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The recent completion of large-scale human cancer sample characterizations such as the Cancer Genome Atlas (TCGA) and the Cancer Proteome Atlas (TCPA) has enabled an altogether different approach for discovery of protein targets that are upregulated by mutant *RAS* genes. This approach uses a direct comparison of tumor samples containing mutant *RAS* genes with either corresponding normal tissue samples or with tumor samples that contain wild-type *RAS* genes. One chief advantage of this approach is the analysis is done using the correct *in vivo* physiological context, whereas with model systems there is no guarantee that the physiological context is correct although it is thought that mouse models are superior to cell culture models<sup>13, 14</sup>. Another advantage is that direct analysis of human cancer takes into account the enormous diversity of co-occurring genomic alterations. This diversity requires very large panels of human cancer cell lines to be representative and presents very difficult challenges for mouse modeling.

Recently, three discrete subtypes of human *KRAS*-mutant lung adenocarcinomas were discovered in a breakthrough of our understanding of the variability within this type of lung cancer. These three subtypes have both distinct RNA expression profiles and different patterns of co-occurring mutations in *TP53, STK11, KEAP1*, and others<sup>15</sup>. Furthermore, based on experiments with appropriate cell lines, there appears to be consistent differences in responses to HSP90-inhibitors, suggesting that this classification scheme could be useful in guiding treatment strategies<sup>15</sup>.

In this study, we analyzed genomic and proteomic data from human lung adenocarcinomas to examine the strength and variability of mutant-*KRAS* activation of proteins and genes. *KRAS*-mutants were compared to other lung adenocarcinoma samples and were also examined for the effects of commonly co-occurring mutations.

#### Results

Raf/MAPK signaling proteins are selectively activated in KRAS-mutant lung adenocarcinomas

230 lung adenocarcinoma samples have been comprehensively characterized for mutations and gene expression by TCGA and these same samples have been characterized by reverse-phase protein array (RPPA) for their levels of 160 different proteins and modified proteins<sup>16</sup>. To determine the selective effects of mutational activation of *KRAS* on these proteins in lung adenocarcinomas, we compared *KRAS*-mutant tumors to other tumors with wild-type *KRAS*. We further divided tumors with wild-type *KRAS* into those with other mutations in components of the mitogenic Raf/MAPK pathway (e.g. *EGFR*, *BRAF*, *RET*, *ALK*) and those without. The number of tumors in these three categories were 75 (*KRAS*-mutant), 85 (other Raf/MAPK mutants), and 70 (all other). These three tumor subsets were compared to each other by pairwise analysis.

We wanted to examine the differences with the greatest potential biological impact, therefore we ranked the differences based on the effect size, rather than *p*-value which can over emphasize very small changes with little variation. We used Cohen's *d* statistic that is highly related to the signal-to-noise statistic used in gene expression studies<sup>17, 18</sup>. Of all the 160 measured modified proteins and native proteins, the top ranking change in *KRAS*-mutant tumors when compared to either group of wild-type *KRAS* tumors is activation of MEK1, as judged by increased levels detected by the anti-phosphoserine 218,222 MEK1 antibody used by TCPA (Figure 1A). Also top-ranking in both comparisons is phosphorylation activation of MAPK (Figure 1B) and a direct kinase target of MAPK, p90-S6-RSK (Figure 1C). Significant large effects on activation of other direct kinase targets of MAPK (YB-1) and further downstream targets (S6) were also observed (Supplementary Table 1).

Also increased in *KRAS*-mutant tumors was activation of mTOR as determined by phosphorylation at serine 2448 (Figure 1D). mTOR activation levels were comparable between *KRAS*-mutant tumors and tumors with other mutations in the Raf/MAPK pathway (Figure 1D). This was somewhat surprising since PI3-kinase is a direct downstream target activated by mutant-Kras proteins, and two kinases activated by PI3-kinase, PDK1 and AKT, did not show any significant phosphorylation activation in either *KRAS*-mutant or other Raf/MAPK mutant tumors (Supplementary Table 1).

Analysis of a well-validated mouse model of *KRAS*-mutant lung adenocarcinoma revealed that tumors showed significant activation of NF-κb and furthermore that these tumors were dependent upon NF-κb activity for tumor maintenance<sup>3</sup>. We did not observe any activation of NF-κb in human *KRAS*-mutant lung adenocarcinomas; in fact, NF-κb activation was significantly lower in KRAS-mutant tumors compared to other Raf/MAPK mutant tumors (Supplementary Table 1).

#### Effect of co-occurring mutations on protein levels in KRAS-mutant lung adenocarcinomas

We next examined the effects of co-occurring mutations in TP53, STK11, and KEAP1 on protein levels in mutant-KRAS tumors. Using cBioPortal analysis of TCGA lung adenocarcinomas, we confirmed that mutations in TP53 and STK11 tend to be mutually exclusive, whereas mutations in KEAP1 and STK11 tend to co-occur. Amongst KRAS-mutant tumors, there were 22 tumors with mutations in TP53 but not STK11 or KEAP1, 20 tumors with mutations in STK11 but not in TP53, and 13 tumors with mutations in KEAP1 but not in TP53 (8 of these tumors also had mutations in STK11). There was only one KRAS-mutant tumor with mutations in both TP53 and STK11 or KEAP1. Excluding this single sample, we divided KRAS-mutant tumors into three groups: those with TP53 mutations (n = 20), those with STK11 and/or STK11 mutations (n = 25), and those without any of these mutations (n = 27). Pairwise analysis of these three groups revealed that the phosphorylation activation status of STK11 pathway components was not significantly affected (Supplementary Table 2).

The strongest effect of co-occurring *TP53* mutations was seen with increased levels of Annexin I protein (Figure 2A). As expected, the group with mutations in the protein kinase gene *STK11* had significantly lower levels of activation of its direct target, AMPK, than the other two groups (Figure 2B). More surprising however were the significantly lower levels of phosphorylation activated PDK1 and PKC-alpha proteins in this same group (Figure 2C and 2D).

# Top ranked KRAS-mutant induced genes by RNA-seq data

We then looked at RNA-seg data to find genes with the greatest induction specifically in KRASmutant lung adenocarcinomas. We ranked genes based on the average effect size when KRASmutant tumors were compared to tumors with other mutations in genes of the Raf/MAPK pathway, and when compared to tumors with no mutations in genes of the Raf/MAPK pathway. KRAS itself was the top-ranked induced gene (Figure 3; Supplementary Table 3). Within the top-100 ranked induced genes were four negative regulators of Raf/MAPK signaling (DUSP4, DUSP6, SPRY4, and NF1); the canonical Raf/MAPK transcriptional targets ETV4 (an ETS-family member) and FOS (Figure 3; Supplementary Table 3). Also within this top-ranked group are several candidate transcriptional oncogenic effectors for mutant KRAS, including the receptor tyrosine kinase genes INSR and IGFR1 and the oncogenic signaling adaptor gene IRS2; the receptor tyrosine phosphatase genes PTPRE and PTPRM; genes encoding guanine-nucleotide exchange-factors that activate Rac/Rho/Cdc42 proteins DOCK5, DOCK1, DNMBP, and PLEKHG2; genes encoding epigenetic regulators CHD2, MLL2, MYST4, BAZ2A (Figure 3). Also included in the top-100 ranked induced genes is the gene CXorf61 which encodes a tumor antigen termed Kita-Kyushu lung cancer antigen 1 (Figure 3).

Approximately 50% of the aforementioned genes that are selectively induced in *KRAS*-mutant lung adenocarcinomas were also significantly affected by co-occurring mutations in *TP53*, *STK11*, or *KEAP1*. For the most part, co-occurring mutations in *STK11* and/or *KEAP1* were associated with significantly stronger expression of these genes. In only one case (*DUSP6*) was co-occurrence of mutant *TP53* associated with stronger expression (Figure 4).

#### **Discussion**

Several components of the Raf/MAPK pathway show strong selective activation in *KRAS*-mutant lung adenocarcinomas, providing further corroboration that this pathway is a key therapeutic target for *KRAS*-mutant lung tumors. However, a potentially important finding of this study is that many of the signaling proteins and pathways thought to be activated in *KRAS*-mutant human lung adenocarcinomas based on studies with models systems are not activated, at least not at steady state levels as assayed by immunoblotting of tissue samples. This makes it less likely that these pathways or proteins correspond to *KRAS*-selective dependencies. These proteins include PI3-kinase (as judged by

activation of AKT), AKT, and NF-κB, all of which have been proposed to be important mediators of mutant *KRAS* in lung adenocarcinomas<sup>3, 19</sup>. Since mTOR is the only component of the PI3-kinase pathway that was selectively activated in *KRAS*-mutant tumors, a greater therapeutic window could conceivably be achieved by combining Raf/MAPK inhibitors with highly selective mTOR inhibitors that do not inhibit PI3-kinase.

All three subtypes of *KRAS*-mutants showed approximately equivalent activation of Raf/MAPK components. However, there were significant differences in protein and phosphoprotein levels detected in the three subgroups. The subgroup comprised of *KRAS/STK11* and *KRAS/KEAP1* double mutants, along with *KRAS/STK11/KEAP1* triple mutants, had significantly lowers levels of phosphorylated AMPK, which is to be expected since AMPK is a direct kinase target of STK11. However this group was more unexpectedly associated with lower levels of phosphorylation activation of PDK1 and PKC-alpha, both of which have been actively pursued targets for developing potentially clinically effective inhibitors. However in this case, it would appear that loss of STK11 function is likely driving this decrease in activity, and that as such this would not represent a potential selective dependency in any of the *KRAS*-mutant subgroups.

Genomic studies usually rank differentially expressed genes by p-value or FDR. However, this ranking method does not taking into account the size of the effect. Potential transcriptional effectors of mutant-KRAS would more likely be genes showing the largest consistent increase in RNA. Therefore we used Cohen's d statistic, closely related to the signal-to-noise statistic, to rank genes selectively upregulated in KRAS-mutant tumors. Amongst the top 100 upregulated genes were several negative regulators of Raf/MAPK signaling, consistent with an important known aspect of negative feedback regulation in this pathway<sup>20, 21</sup>, as well as canonical transcriptional targets for Raf/MAPK signaling including FOS and the ETS-family member ETV4. Additionally, several potential oncogenic transcriptionally activated targets were uncovered including IGF1R and IRS2. A connection between mutant KRAS and IGF1R in lung adenocarcionoma has previously been documented<sup>22</sup>. We detected transcriptional activation of several quanine-nucleotide exchange factors that activate different proteins in the Rac/Rho/Cdc42 family. Thus the effect of mutational activation of KRAS on quanine-nucleotide exchange factors for small GTP-binding proteins is broader in lung adenocarcinoma than its biochemical activation of exchange factors that activate Ral proteins. Upregulation of the receptor protein-tyrosine phosphatase gene PTPRE, the third highest ranked upregulated gene, is of particular interest since it has been shown to be upregulated by RAS in a mouse model of mammary carcinoma and to possess on its own the ability to promote mammary tumor formation<sup>23, 24</sup>. Finally, this set of upregulated genes includes a potential immunotherapy target for KRAS-mutant lung adenocarcinomas, the Kita-Kyushu lung cancer antigen 1, encoded by the *CXorf61* gene<sup>25</sup>. This protein has also been shown to be a potential target for T cell based therapies in triple-negative breast cancer<sup>26</sup>.

## Figure legends

**Figure 1**. Relative levels of select phosphoproteins in three subgroups of human lung adenocarcinomas. All values are plotted on an arbitrary log2 scale to highlight relative levels rather than absolute amounts. For each panel, the red colored group represents *KRAS*-mutant tumors, the orange colored group represents other Raf/MAPK pathway mutants, and the blue colored group represents all other tumors. Brackets with associated *p*-values indicate significant differences. (A) levels of MEK1\_pS217\_S221; (B) levels of MAPK\_pT202\_Y204; (C) levels of p90RSK\_pT359\_S363; (D) levels of mTOR\_pS2448.

**Figure 2**. Relative levels of select proteins phosphoproteins in three subgroups of KRAS-mutant adenocarcinomas. All values are plotted on an arbitrary log2 scale to highlight relative levels rather than absolute amounts. For each panel, the green colored group represents *KRAS/TP53* doublemutant tumors, the violet colored group represents other *KRAS/STK11*, *KRAS/KEAP1*, and *KRAS/STK11/KEAP1* compound mutants, and the brown colored groups *KRAS* mutants without those other mutations. Brackets with associated *p*-values indicate significant differences. (A) levels of Annexin I; (B) levels of AMPK\_pT172-R; (C) levels of PDK1\_pS241; (D) levels of PKC-alpha\_pS657.

**Figure 3**. Heatmap showing the relative RNA values of select genes in three subgroups of human lung adenocarcinomas. The three subgroups include *KRAS*-mutant tumors, other Raf/MAPK pathway mutants, and all other tumors. The select genes are indicated on the right. Row values were normalized and scaled and presented as Z-scores.

**Figure 4**. Heatmap showing the relative RNA values of select genes in three subgroups of *KRAS*-mutant human lung adenocarcinomas. The three subgroups are *KRAS/TP53* double mutants; *KRAS/STK11, KRAS/KEAP1*, and *KRAS/STK11/KEAP1* compound mutants; and *KRAS* mutants without those other mutations. The select genes are indicated on the right. Row values were normalized and scaled and presented as Z-scores.

## **Supplementary Table Legends**

**Supplementary Table 1.** Protein levels in three groups of lung adenocarcinomas. Mean levels in three groups are listed along with the p-values and effect sizes of pairwise comparisons. G1 = KRAS-mutants; G2 = other Raf/MAPK pathway mutants; G3 = all others.

**Supplementary Table 2.** Protein levels in three groups of KRAS-mutant lung adenocarcinomas. Mean levels in three groups are listed along with the p-values and effect sizes of pairwise comparisons. G1 = KRAS/TP53 double mutants; G2 = KRAS/STK11 or KRAS/KEAP1 double mutants and KRAS/STK11/KEAP1 triple mutants; G3 = all others.

**Supplementary Table 3.** RNA levels in three groups of lung adenocarcinomas. Mean levels in three groups are listed along with the p-values and effect sizes of pairwise comparisons. G1 = KRAS-mutants; G2 = other Raf/MAPK pathway mutants; G3 = all others.

# Methods

We used mutational, RNAseq and protein data in our analysis from the TCGA lung adenocarcinoma project. All data downloaded was Level 3. Mutation data was downloaded from cBioPortal for Cancer Genomics (<a href="www.cbioportal.org">www.cbioportal.org</a>), under Lung Adenocarcinoma (TCGA, Provisional) with "All Complete Tumors (230)". RNAseq data was downloaded from FireBrowse by Broad Institute (<a href="www.firebrowse.org">www.firebrowse.org</a>), choosing "illuminahiseq\_rnaseqv2-RSEM\_genes". Protein data was downloaded from The Cancer Proteome Atlas (TCPA) project in MD Anderson Cancer Center (<a href="bioinformatics.mdanderson.org/main/TCPA">bioinformatics.mdanderson.org/main/TCPA</a>). To generate heatmaps, the R package "gplots" was used.

We segregated samples using cBioPortal into three mutually exclusive groups: ones with *KRAS* missense mutations, ones with mutations in genes encoding other components of the Raf/MAPK pathway, and other samples. Raf/MAPK pathway mutatations included *BRAF* missense mutations, *EGFR* missense, splice, and inframe mutations, *NF1* mutations, *MET* missense mutations, *ALK* fusions, and *RET* fusions. Those three subgroups were used to perform pairwise comparisons using protein (RPPA) and RNAseq data. We used Cohen's d to measure effect size using the R-package "effsize". Ranking with both effect size and t-test p-value generated top-ranked proteins and genes induced in *KRAS* mutants or Raf/MAPK pathway mutants (Figure 1, Supplementary Table 1). We used a similar approach to divide *KRAS* mutants into three groups depending on *TP53*, *STK11*, or *KEAP1* status.

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#### **Author Contributions**

Conceived and designed the analysis: SP. Computational analysis: JL. Analyzed data: JL, SP, RS. Wrote the paper: JL, SP.

#### **Additional Information**

The authors have declared that no competing financial interests exist.

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Figure 1.

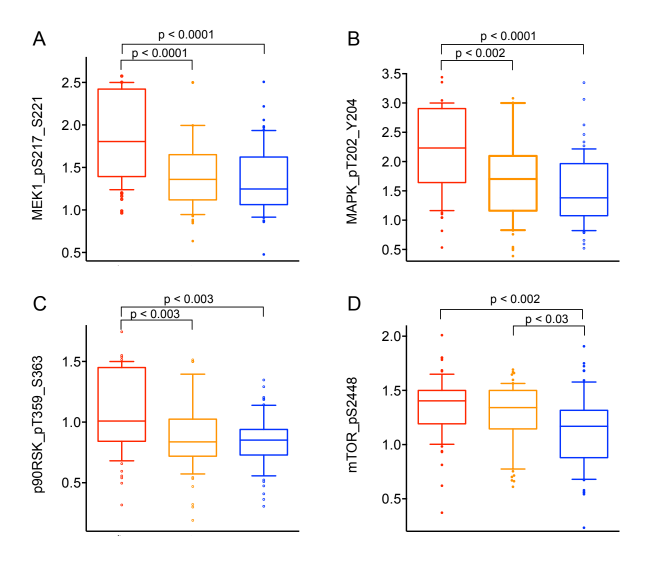


Figure 2.

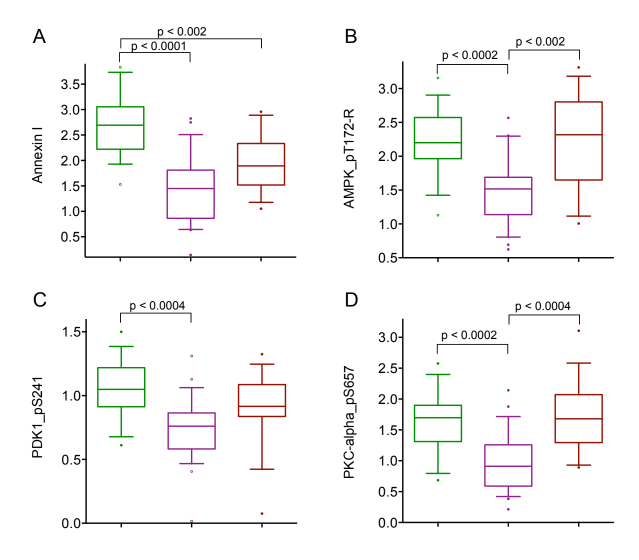
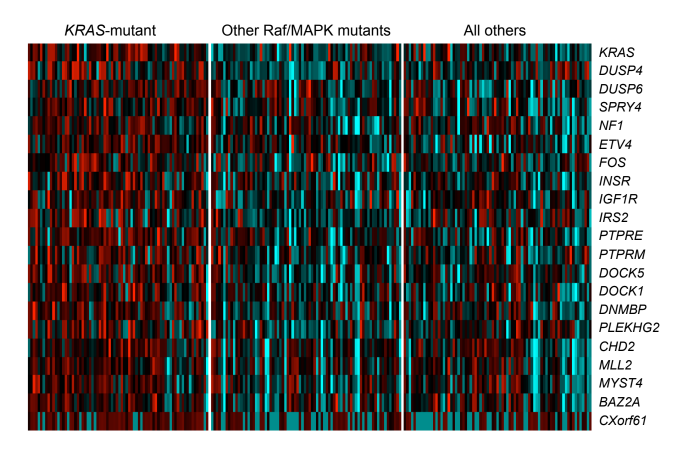
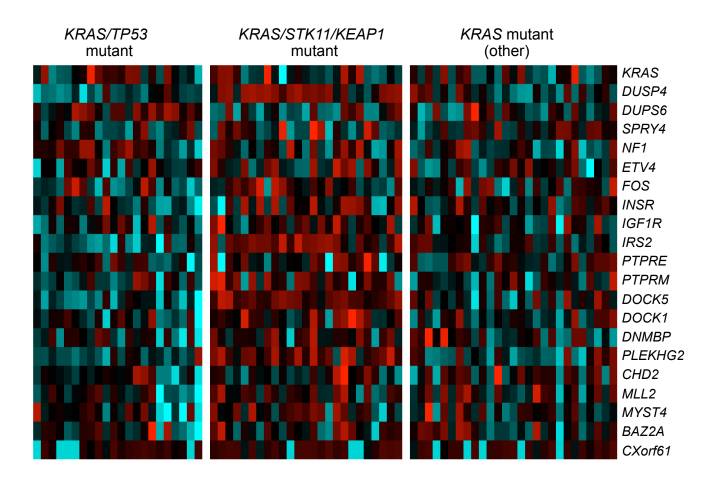


Figure 3.



-2 -1 0 1 2 Row Z-score

Figure 4.



-2 -1 0 1 2 Row Z-score