Targeting Endogenous K-RAS for Degradation through the Affinity-Directed Protein Missile System

Highlights

- Generation of A549 cells with a homozygous knockin of GFP tag on the KRAS gene
- Proteasomal degradation of endogenous GFP-K-RAS using a VHL-GFP-nanobody fusion
- Proteasomal degradation of endogenous H/K-RAS using VHL-H/K-RAS-monobody fusion

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In Brief
By using the affinity-directed protein missile (AdPROM) system comprised of VHL and high-affinity binders of GFP and H/K-RAS, Röth et al. demonstrate the targeted proteasomal degradation of GFP-K-RAS, which was knocked in using CRISPR/Cas9, and endogenous H/K-RAS.
Targeting Endogenous K-RAS for Degradation through the Affinity-Directed Protein Missile System

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SUMMARY

K-RAS is known as the most frequently mutated oncogene. However, the development of conventional K-RAS inhibitors has been extremely challenging, with a mutation-specific inhibitor reaching clinical trials only recently. Targeted proteolysis has emerged as a new modality in drug discovery to tackle undruggable targets. Our laboratory has developed a system for targeted proteolysis using peptidic high-affinity binders, called “AdPROM.” Here, we used CRISPR/Cas9 technology to knock in a GFP tag on the native K-RAS gene in A549 adenocarcinoma (A549 GFPKRAS) cells and constructed AdPROMs containing high-affinity GFP or H/K-RAS binders. Expression of GFP-targeting AdPROM in A549 GFPKRAS led to robust proteasomal degradation of endogenous GFP-K-RAS, while expression of anti-HRAS-targeting AdPROM in different cell lines resulted in the degradation of both GFP-tagged and untagged K-RAS, and untagged H-RAS. Our findings imply that endogenous RAS proteins can be targeted for proteolysis, supporting the idea of an alternative therapeutic approach to these undruggable targets.

INTRODUCTION

The three RAS oncogenes, H-RAS, K-RAS, and N-RAS, represent the most frequently mutated genes in cancer (Cox et al., 2014; Hobbs et al., 2016). They encode four highly similar proteins, namely H-RAS, N-RAS, K-RAS4A, and K-RAS4B, which undergo C-terminal farnesylation (Reiss et al., 1990; Schaber et al., 1990). Farnesylation, in combination with palmitoylation in the hypervariable region (HVR) (N-RAS, H-RAS, and K-RAS4A) or with a polybasic signal in the HVR (K-RAS4B), mediates the plasma membrane interaction (Ahearn et al., 2012). RAS proteins are small GTPases, which cycle between the GTP-bound (active) and GDP-bound (inactive) states, controlled by guanosine nucleotide exchange factors and GTPase activating proteins (GAPs) (Vigil et al., 2010). Activation of RAS proteins by various extracellular growth factors initiates activation of numerous downstream signaling networks, including BRAF/mitogen-activated protein kinase (MAPK) and phosphatidylinositol 3-kinase pathways (Khan et al., 2019a), which are critical for cell proliferation and viability. Many pathogenic mutations in RAS genes impair GAP-mediated GTP hydrolysis, thereby favoring the persistence of the active RAS-GTP state, which triggers constitutive activation of downstream signaling resulting in unchecked proliferation of cancer cells (Hobbs et al., 2016; Marcus and Mattos, 2015).

As the oncogenicity of RAS mutations has been known for over three decades, intensive efforts have been made toward drugging them. These efforts are yet to result in effective RAS-inhibitor therapies (Cox et al., 2014; Papke and Der, 2017). This has promoted the perception that RAS proteins are undruggable. Several factors make RAS proteins difficult targets to engineer selective small-molecule inhibitors. First, the relatively high concentrations of GTP and GDP in cells and picomolar affinity to binding RAS proteins makes it almost impossible to develop GTP/GDP analogs as inhibitors (Cox et al., 2014; John et al., 1990). Second, structural analysis of RAS proteins revealed few sufficiently large and deep hydrophobic pockets on the surface for small-molecule binding (O’Bryan, 2019; Pai et al., 1989). Recently, a covalent inhibitor targeting a cysteine in K-RAS G12C was developed to target this specific mutation (Ostrem et al., 2013). However, these barriers and failure to directly target RAS have prompted researchers to explore targeting upstream regulators, or downstream effectors of RAS proteins (Cox et al., 2014; Papke and Der, 2017; Waldmann et al., 2004), as well as altering levels of RAS protein, for example, by inducing targeted degradation of RAS (Nabet et al., 2018).

Most targeted protein degradation approaches harness the cellular proteolytic pathways that naturally maintain proteostasis, with the ubiquitin proteasome system (UPS) being frequently exploited (Röth et al., 2019). Protein degradation by the UPS is triggered by conjugation of ubiquitin chains onto the target protein, which is achieved through a sequential action of three enzymes: the ubiquitin-activating enzyme (E1), which activates
the C-terminal glycine residue of ubiquitin in an ATP-dependent manner; a ubiquitin-conjugating enzyme (E2), which conjugates the activated ubiquitin to its active site cysteine; and a ubiquitin ligase (E3), which facilitates the transfer of ubiquitin from E2 to primarily lysine residues on substrate proteins (Pickart and Eddins, 2004; Roos-Mattjus and Sistonen, 2004). Further ubiquitylation on one or more lysine residues within ubiquitin then triggers polyubiquitylation, followed by degradation by the proteasome (Akutsu et al., 2016; Komander and Rape, 2012; Yau and Rape, 2016). Targeting RAS for proteolysis relies on the engagement of the cellular proteolytic systems for its ubiquitylation and degradation. In this context, it has been shown that the heterobifunctional molecule dTAG-13, which recruits FKBP12<sup>F36V</sup>-tagged proteins of interest (POIs) to the CRBN/CUL4A E3 ubiquitin ligase for their degradation, can degrade FKBP12<sup>F36V,-K-RAS</sup> overexpressed in cell lines (Nabet et al., 2018). However, FKBP12<sup>F36V</sup> itself can be targeted for ubiquitylation when using heterobifunctional small-molecule binders (Winter et al., 2015). Therefore, it remains unclear, whether using dTAG13 on FKBP12<sup>F36V,-K-RAS</sup> results in the ubiquitination of K-RAS or FKBP12<sup>F36V</sup>. Such information is not only key to evaluate proteolysis as a druggable approach for targeting RAS proteins but also to inform on the development of effective heterobifunctional RAS degraders.

We have previously developed an effective proteolytic affinity-directed protein missle (AdPROM) system for UPS-mediated POI degradation (Fulcher et al., 2016, 2017). AdPROM consists of a fusion of von Hippel-Lindau (VHL) protein, a substrate recruiter of the CUL2-RING E3 ligase complex, and high-affinity binders, such as nanobodies and monobodies, of POIs. Deliver AdPROM into multiple cell lines through retroviral transductions led to efficient degradation of endogenous target proteins, including SHP2 and ASC (Fulcher et al., 2017). Furthermore, to target POIs for which no high-affinity polypeptide binders exist, we utilized CRISPR/Cas9 genome editing to rapidly introduce GFP tags on endogenous VPS34 and PAWS1 genes, and used the AdPROM system consisting of anti-GFP nanobody fused to VHL to achieve near complete degradation of the endogenous GFP-VPS34 and PAWS1-GFP proteins (Fulcher et al., 2016). In this study, we explore the use of the AdPROM system, and demonstrate its efficacy, for targeted degradation of endogenously GFP-tagged K-RAS and untagged, endogenous K-RAS from cells.

RESULTS

Generation of a GFP-KRAS Knockin Non-small Cell Lung Cancer A549 Cell Line

The high degree of amino acid sequence similarity between the four RAS proteins, i.e., K-RAS4A, K-RAS4B, H-RAS, and N-RAS (Figure 1A), and the subsequent difficulty in generating selective antibodies against individual isoforms pose substantial challenges in studying specific RAS proteins (Waters et al., 2017). To explore targeted proteolysis of K-RAS using the AdPROM system, we used CRISPR/Cas9 technology to generate an A549 non-small cell lung carcinoma (NSCLC) cell line harboring a homozygous knockin of green fluorescent protein (GFP) cDNA at the N terminus of the native K-RAS gene (Figure S1). As K-RAS4A and K-RAS4B are splice variants differing only in their extreme C terminus (Figure 1A), this approach allowed us to simultaneously tag both isoforms with GFP. The homozygous GFP knockins on the native KRAS locus (A549<sup>GFPKRAS</sup>) were verified by genomic sequencing (Figure S1). Moreover, by western blot analysis using both panRAS and K-RAS4B antibodies, the appearance of higher-molecular-weight GFP-K-RAS species with a concurrent disappearance of the native-molecular-weight K-RAS species was evident in the A549<sup>GFPKRAS</sup> cell line compared with wild-type (WT) A549 control cells (Figure 1B). The use of a panRAS antibody resulted in the detection of two distinct bands in A549 WT cells (Figure 1B). As the lower band remained intact in A549<sup>GFPKRAS</sup> cells, it most likely corresponds to H- and/or N-RAS (Figure 1B). However, in A549 cells we were unable to detect any endogenous signals with most commercially available H-RAS-, N-RAS-, or K-RAS4A-specific antibodies (listed in the STAR Methods). As K-RAS is an integral part of the MAP kinase signaling pathway, we wanted to analyze the effect GFP fusion would have on the MAPK signaling pathways. Under cell culture conditions, both MEK1/2 and ERK phosphorylations were strongly decreased in A549<sup>GFPKRAS</sup> cells compared with A549 WT cells, while levels of BRAF were slightly decreased (Figure 1B). Interestingly, phosphorylation of EGFR receptor at Tyr1068 and AKT phosphorylation at Ser473 was higher in A549<sup>GFPKRAS</sup> cells than in WT cells (Figure 1B). By qRT-PCR, we showed that levels of H- and N-RAS transcripts were slightly reduced in A549<sup>GFPKRAS</sup> cells compared with WT A549 cells, while transcript levels of K-RAS were reduced by roughly 50% (Figure S2). We were able to efficiently immunoprecipitate GFP-K-RAS from A549<sup>GFPKRAS</sup> but not WT A549 cell extracts (Figure 1C).

A number of RAS antibodies have been evaluated for selective recognition of the different RAS proteins by western blotting (Waters et al., 2017), but none of these have been selective for use in immunofluorescence studies. Consequently, studies evaluating subcellular distribution of RAS proteins have been restricted to overexpression systems. Validation of A549<sup>GFPKRAS</sup> cells allowed us to investigate the subcellular distribution of endogenous GFP-K-RAS driven by the native promoter. Endogenous GFP-K-RAS displayed predominantly plasma membrane distribution, which was confirmed by co-staining with P120 catenin, which is known to localize to the plasma membrane (Reynolds et al., 1994) (Figures 1D and S3). In addition, we also observed some weak cytoplasmic localization of GFP-K-RAS. However, no co-localization of GFP-K-RAS was observed with mitochondrial marker ATPB (Schatz and Butow, 1983) (Figures 1D and S3).

Finally, we compared turnover of WT K-RAS and GFP-K-RAS proteins by adding cycloheximide to the respective cell line and analyzing protein levels over the course of 12 h. We could not find any remarkable differences in protein stability between WT and GFP-K-RAS (Figures 1E and S4). However, as expected, a robust degradation of c-myc was observed within 2–4 h (Figure S4).

Targeted Degradation of GFP-K-RAS by the Proteolytic AdPROM System

We sought to test whether endogenously expressed GFP-K-RAS protein in A549<sup>GFPKRAS</sup> cells could be targeted for degradation by AdPROM (Fulcher et al., 2016, 2017). We have previously shown
that fusion of VHL to an aGFP16 nanobody recruits GFP-tagged proteins, such as VPS34 and PAWS1, to the CUL2-RBX1 E3 ligase machinery for target ubiquitination and subsequent proteasomal degradation (Fulcher et al., 2016). Therefore, we postulated that GFP-K-RAS could be recruited in a similar manner to the CUL2-RBX complex for ubiquitination and degradation (Figure 2A).

Indeed, expression of VHL-aGFP16 AdPROM resulted in near complete clearance of GFP-K-RAS from A549GFPKRAS cells compared with the untransduced controls, while the low-molecular-weight band corresponding to H- and/or N-RAS was unaffected (Figure 2B). In contrast, neither VHL nor the aGFP16 nanobody alone, serving as controls, caused any apparent changes in the steady-state levels of GFP-K-RAS from A549GFPKRAS cells compared with the untransduced controls, while the low-molecular-weight band corresponding to H- and/or N-RAS was unaffected (Figure 2B).

Treatment of VHL-aGFP16 AdPROM expressing A549GFPKRAS cells with the Cullin neddylation inhibitor MLN4924 partially rescued the degradation of GFP-K-RAS compared with DMSO-treated controls (Figure 2C). The neddylation of CUL2 allows a conformational change of the CUL2-RBX E3 ligase machinery so that the RBX E3 ligase can ubiquitinate substrates recruited by VHL. In line with this notion, the levels of HIF1α protein, a bona fide substrate of VHL (Yu et al., 2001), were stabilized upon MLN4924 treatment compared with DMSO control (Figure 2C).

Despite the high apparent efficiency of GFP-KRAS degradation by VHL-aGFP16 AdPROM, retroviral transduction of A549GFPKRAS cells often generates uneven levels of AdPROM expression in a mixed population of cells. To get a better understanding of the distribution of the cells within this population, we used a flow cytometric analysis based on GFP fluorescence. We used gates to define a GFP-positive population based on the GFP signal from untransduced A549GFPKRAS cells and using WT A549 cells as a GFP-negative control (Figure 2D). In accordance with the western blot results (Figures 2B), 98% of cells transduced with VHL-aGFP16 AdPROM virus showed GFP-KRAS degradation as compared with untransduced A549GFPKRAS cells (Figure 2D), which manifested in an overall reduction of GFP fluorescence of the single-cell population (Figure 2E). The remaining 2% of A549GFPKRAS cells produced GFP signal comparable with untransduced GFP-positive-population, which could be due to low-level AdPROM expression within these cells (Figure 2D). In contrast, A549GFPKRAS cells expressing VHL or aGFP16 alone were defined as GFP positive at 99.3% or 99.8%, respectively (Figures 2D and 2E).
AdPROM-Mediated Degradation of Endogenous RAS Proteins

The AdPROM-mediated degradation of GFP-K-RAS in A549\(^{GFPKRAS}\) cells demonstrated the feasibility of targeted degradation of endogenous K-RAS. However, the presence of the GFP tag raises the possibility of ubiquitination occurring on the GFP moiety, instead of K-RAS. Therefore, we sought to explore whether we could exploit the AdPROM system to degrade endogenous, unmodified K-RAS from A549 cells. At present, there are no reported high-affinity, selective...
polypeptide binders of K-RAS. However, we utilized an anti-H-RAS (aHRAS) monobody that was reported to bind and immunoprecipitate both H-RAS and K-RAS, but not N-RAS (Spencer-Smith et al., 2017). Using this monobody with a FLAG tag, we showed that anti-FLAG immunoprecipitates (IPs) could robustly co-precipitate both GFP-tagged and untagged K-RAS as well as the lower-molecular-weight protein band matching both H- and N-RAS, which is most likely to be H-RAS (Spencer-Smith et al., 2017) (Figure 3A). However, neither of the RAS proteins was completely depleted from flowthrough extracts, suggesting incomplete immunoprecipitation (Figure 3A). In contrast, anti-FLAG IPs from extracts expressing Flag-VHL control did not co-precipitate either protein (Figure 3A).

Figure 3. Degradation of Endogenous RAS Using a RAS-Specific Monobody
(A) Cell lysates of indicated cell lines were subjected to immunoprecipitation with anti-Flag beads. Input (I), flowthrough (Ft), and precipitates (IP) were run on SDS-PAGE and subjected to western blotting with the respective antibodies.
(B) After retroviral transduction and selection, cell lysates of indicated cell lines were separated on SDS-PAGE and analyzed by western blotting using the indicated antibodies.
(C) Indicated cell lines were treated with 1 μM MLN4924 or 0.1% DMSO for 24 h. Cell lysates were separated on SDS-PAGE and analyzed by western blotting using the indicated antibodies.
(D) Indicated cell lines were analyzed on a Canto flow cytometer. Shown populations were preselected for cells and single cells before defining the gate for GFP-positive cells (shown). GFP-A is plotted against PE-A in all cases. Numbers indicate percentage of cells within the respective gate.
(E) Histogram representation of plots in (D); Ki = A549GFPKRAS cells.
(F) Wide-field immunofluorescence microscopy of indicated cell lines treated with anti-GFP antibody and DAPI for staining. Scale bars, 10 μm. Two representative images are shown for each condition. Western blots and immunofluorescence data are representative of at least three independent experiments. Flow cytometry data are representative of two independent experiments.
CUL2-RBX E3 ligase machinery (Figure 3C). As expected, MLN4924 also stabilized endogenous HIF1α (Figure 3C).

We also assessed the relative abundance of GFP-K-RAS in mixed populations of A549GFPKRAS cells transduced with VHL-aHRAS AdPROM in comparison with controls by flow cytometry. We found that 77% of cells showed degradation of GFP-K-RAS, as assessed by the shift of the GFP-positive gated population toward the GFP-negative population (Figure 3D) and the overall reduction of GFP signal (Figure 3E). The remaining 23% of cells transduced with VHL-aHRAS were seemingly unaffected in terms of GFP level (Figures 3D and 3E). Transductions with VHL or aHRAS alone did not induce any noticeable shift of the GFP population or GFP signal intensity (Figures 3D and 3E).

Uneven retroviral transduction of cells could result in unequal expression of the AdPROM constructs in different cells resulting in a mixed, divergent cell population, which may account for the apparent uneven degradation of GFP-K-RAS through VHL-aHRAS. When we analyzed these A549GFPKRAS mixed cell populations by immunofluorescence for GFP signal, in non-transduced and aHRAS-transduced control cells, a predominant plasma membrane GFP-K-RAS signal was evident (Figure 3F). Transduction of A549GFPKRAS cells with either VHL-aHRAS or VHL-aGFP16 AdPROM produced a heterogeneous population comprising cells with missing or severely attenuated GFP signal, and cells with intact GFP-K-RAS staining pattern, localizing mainly to the plasma membrane (Figure 3F). In contrast, we noticed a slight increase in endoplasmic reticulum (ER)/perinuclear GFP-K-RAS signal in cells transduced with the aHRAS monobody alone (Figure 3F). Interestingly, we detected that the majority of the monobody itself was in the nucleus (Figure S5), while we were unable to consistently detect signals for the AdPROM fusion proteins by anti-FLAG immunofluorescence (Figure S5).

AdPROM-Mediated Degradation of Untagged Endogenous RAS Proteins

Having verified that VHL-aHRAS AdPROM recognizes and degrades GFP-K-RAS, we next tested its ability to degrade endogenous K- and H-/N-RAS in WT A549 cells. The transduction of cells with VHL-aHRAS resulted in a substantial reduction in apparent levels of both K-RAS (upper band) and H-/N-RAS (lower band) proteins as detected by the panRAS antibody and compared with the non-transduced controls (Figure 4A). Unlike in A549GFPKRAS cells (Figure 3B), WT cells transduced with VHL-aGFP16 AdPROM did not display any noticeable changes in K-RAS and H-/N-RAS protein levels relative to...
non-transduced cells (Figure 4A), further validating the targeted nature of RAS degradation by AdPROM. Cells transduced
with the aHRAS monobody alone showed a slight increase in abundance of both K-RAS and H-/N-RAS proteins compared
with non-transduced controls (Figure 4A). To ascertain whether AdPROM-mediated degradation occurs via the proteasome,
we treated A549 cells expressing the VHL-aHRAS AdPROM system or A549 WT cells with proteasomal inhibitors MG132 and bortezomib, both of which resulted in a strong accumulation of polyubiquitinated proteins (Figure 4B). In A549 WT cells, RAS protein levels increased only slightly after 14 h of MG132 and bortezomib treatment. In contrast, in VHL-aHRAS transduced cells, both bortezomib and MG132 rescued RAS protein levels, with bortezomib rescuing it to levels comparable with A549 WT cells (Figure 4B). Next, we sought to explore whether RAS protein degradation triggers a change in RAS transcript levels. We transduced A549 WT cells with constructs encoding Flag-VHL-aHRAS, Flag-aHRAS, Flag-VHL-aGFP16, or Flag-VHL and a pBabeD empty construct as a calibrator. While we noticed a slight increase in K-RAS4A transcripts in cells expressing aHRAS or VHL-aHRAS relative to other cells, these changes were not statistically significant (Figure 4C).

Next, we looked at the global quantitative proteomic changes upon targeted degradation of RAS proteins through the AdPROM system. To this end, we used puromycin-selected cells transduced with pBabeD empty vector, a plasmid encoding Flag-VHL-aHRAS, or a plasmid encoding for Flag-VHL, and performed tandem mass tag-labelled total proteome analysis (Figures 4D and 6). We found that H-RAS was significantly reduced by more than 50% in VHL-aHRAS samples, compared with both VHL alone and the empty vector controls (Figure 4D, Table 1). K-RAS was significantly reduced by ~15% in VHL-aHRAS samples compared with empty vector; however, the reduction was not significant when compared with VHL alone control. N-RAS was unchanged in all conditions, further consolidating the specificity of the monobody toward the H- and K-RAS isoforms (Spencer-Smith et al., 2017). Interestingly, in VHL-aHRAS samples the only proteins significantly downregulated by a factor higher than 2 when compared with VHL alone were H-RAS and NPTX1, the latter, however, most likely stemmed from upregulation by VHL, as the increase was observed in the “VHL alone” sample and it was not changed when compared with the empty vector (Table 1). To our surprise, M-RAS protein abundance increased by 2-fold in cells expressing VHL-aHRAS. In addition, transcription initiation factor TFIID subunit 4B (TAF4B) was more than 3-fold more abundant in the VHL-aHRAS-expressing cells compared with controls (Table 1). The abundance of transcription factor LBH, Annexin-A8-like protein 1 (ANXA8L1), NEDD9, and Transgelin also increased >1.5-fold in VHL-aHRAS-expressing cells compared with controls (Table 1). VHL abundance was roughly 9-fold higher in the VHL-aHRAS-transduced samples compared with the empty vector control and 2-fold higher compared with VHL alone samples (Table 1). While we could not detect HIF1α, we found that overexpression of VHL, either alone or when fused to the aHRAS monobody, did not significantly change protein levels of other VHL substrates (Zhang and Yang, 2012) MYBBP1A (Lai et al., 2011) or RNA polymerase II subunit RPB1 (Kuznetsova et al., 2003).

Expansion of the RAS-Targeting AdPROM System in Different Cell Lines

Having demonstrated for the first time that the VHL-aHRAS AdPROM system could target endogenous, unmodified H- and K-RAS for degradation in A549 cells, we sought to explore whether the system would work in other cell lines. First, we compared different cell lines for their endogenous RAS protein expression (Figure 5A) relative to A549 cells. All cells tested displayed K-RAS protein expression similar to, or slightly lower than, A549 cells.

Table 1. Proteins Identified in Total Proteome Analysis in Comparison of VHL-aHRAS versus VHL Transduced Cells, as Either 2-Fold More or Less Abundant

| Classification       | Protein                                      | Uniprot ID   | VHL-aHRAS/VHL Fc | p Value  | VHL-aHRAS/EV Fc | p Value  | VHL/EV Fc | p Value  |
|----------------------|----------------------------------------------|--------------|------------------|----------|-----------------|----------|-----------|----------|
| VHL-aHRAS >2-fold decrease | Neuronal pentraxin-1 (NPTX1) | Q15818 | 0.348 | 0.00607 | 0.961 | 0.77092 | 3.028 | 0.01113 |
|                      | H-RAS                                       | P01112       | 0.464 | 0.01404 | 0.466 | 0.01391 | 0.998 | 0.84955 |
| VHL-aHRAS >2-fold increase | TFIIID subunit 4B (TAF4B) | Q92750 | 4.875 | 0.012 | 3.598 | 0.03235 | 0.801 | 0.25714 |
|                      | LBH                                          | Q53Q2V       | 2.847 | 0.00527 | 1.942 | 0.08367 | 0.682 | 0.38491 |
|                      | Annexin A8-like protein 1 (ANXA8L1)          | Q5VT79       | 2.434 | 0.03474 | 1.487 | 0.20081 | 0.511 | 0.68722 |
|                      | Enhancer of filamentation 1 (NEDD9)          | Q14511       | 2.268 | 0.00515 | 1.508 | 0.01131 | 0.72 | 0.04447 |
|                      | Transgelin (TAGLN)                           | Q01995       | 2.176 | 0.00251 | 1.823 | 0.00281 | 0.838 | 0.26401 |
|                      | M-RAS                                        | Q14807       | 2.079 | 0.00114 | 2.017 | 0.05143 | 1.013 | 0.48323 |
| K/N-RAS              | K-RAS                                        | P01116       | 0.909 | 0.48915 | 0.841 | 0.01985 | 0.886 | 0.59775 |
|                      | N-RAS                                        | P01111       | 0.984 | 0.84292 | 1.082 | 0.82715 | 1.036 | 0.86126 |
| VHL and targets      | VHL                                          | P40337       | 1.952 | 0.00117 | 9.313 | 0.00018 | 4.686 | 0.00104 |
|                      | MYBBP1A                                      | Q9BGQ0       | 0.988 | 0.73087 | 0.886 | 0.47528 | 0.857 | 0.60345 |
|                      | RPB1                                         | P24928       | 1.028 | 0.78989 | 0.991 | 0.75666 | 0.987 | 0.24496 |

Values of identified proteins are given for VHL-aHRAS versus empty vector (EV) and VHL versus EV-transduced cells as well. UniProt ID is given, as well as fold change (Fc) values and p values for the respective comparison. In addition, results are shown for K- and N-RAS, as well as VHL and two described VHL substrates.
cells. SW620 cells, which harbor the G12V mutation on K-RAS (Morandi et al., 2012), displayed similar levels of expression to A549 cells; however, we noticed that K-RAS in this cell line produced a slight but noticeable molecular weight shift when probed with panRAS and K-RAS4B antibodies (Figure 5A). Protein levels corresponding to the lower H- and/or N-RAS band were similar in all lines tested but overall, much lower in intensity than that seen for K-RAS. We tested the ability of VHL-aHRAS

Figure 5. Degradation of RAS in Different Cell Lines and Effects on Proliferation
(A–C) Lysates of untreated (A) or retrovirally transduced cell lines (indicated expression constructs) (B) were separated by SDS-PAGE and analyzed by western blotting with the indicated antibodies. Comparison of cell lines in (A) was done only once. K-RAS mutation statuses for individual cell lines are indicated in brackets. (C) A total of 5,000 cells from (B) or A549 cells from (Figure 4A) or (Figure 3B) were grown in triplicate in 12-well dishes. After 7 days, cells were fixed and stained with crystal violet. A representative image of the replicates is shown.
(D) Staining from plates in (C) was extracted by methanol and absorbance at 570 nm was measured. Plotted 570-nm values are relative to the respective untreated sample. The number of biological replicates (applies to western blots in (B) as well) is indicated next to the cell line and error bars (SD) are shown. For statistical analysis one-way ANOVA analysis with Dunnett’s multiple comparisons test was done. Comparisons were drawn to the untreated sample. *p < 0.05; **p < 0.01.
AdPROM to degrade RAS proteins from HT-29 and SW620 cells. In HT-29 cells, which express WT RAS proteins but harbor the activating BRAF V600E mutation (Tan et al., 2008), only the levels of H-RAS but not K-RAS proteins were reduced by VHL-aH-RAS AdPROM compared with controls (Figure 5B, left panel). For SW620 cells, which harbor the G12V mutation of K-RAS, we noticed a high K-RAS signal to H-/N-RAS signal ratio, as the latter was barely detectable (Figure 5B, right panel). We observed stabilization of K-RAS with the aHRAS monobody, latter was barely detectable (Figure 5 B, right panel). We noticed a high K-RAS signal to H-/N-RAS signal ratio, as the SW620 cells, which harbor the G12V mutation of K-RAS, we

Finally, we wanted to explore whether targeted degradation of K- and H-RAS proteins from WT A549, HT29, and SW620 cells using the VHL-aH-RAS AdPROM, and GFP-K-RAS from A549GFPKRAS cells using the VHL-aGFP16 AdPROM would impact cell proliferation. No significant differences in proliferation could be observed for either WT A549 or A549GFPKRAS cells following AdPROM-mediated degradation of the respective RAS proteins compared with controls after 7 days, as measured by crystal violet staining (Figures 5C and 5D). Although A549 cells harbor the oncogenic KRASG12S mutation, they also harbor over 250 genetic mutations (COSMIC cell lines project) (Tate et al., 2018), including some known oncogenes and tumor suppressors reducing the likelihood that these cells are solely dependent on the K-RASG12S oncogene for their proliferation. Interestingly, proliferation of HT-29 cells was reduced by about 50% by the aHRAS monobody alone (Figures 5C and 5D), while the VHL-aHRAS and VHL-aGFP16 constructs reduced growth to a lesser extent (Figure 5D). Both the aHRAS monobody alone and the VHL-aHRAS AdPROM, but not VHL-aGFP16 AdPROM were able to reduce the proliferation of SW620 cells significantly by about 50% (Figures 5C and 5D).

**DISCUSSION**

Overexpression of GFP-tagged or other epitope-tagged K-RAS has been used frequently to investigate RAS localization (Schmick et al., 2014; Spencer-Smith et al., 2017; Tsai et al., 2015). This overcomes the difficulty in the study of RAS proteins in the absence of robust reagents to reliably detect specific RAS proteins at the endogenous levels, especially by immunofluorescence (Waters et al., 2017). Our homozygous A549GFPKRAS NSCLC cell line, generated using CRISPR/Cas9, has allowed us not only to assess localization of endogenously driven GFP-K-RAS protein, but its mobility shift has allowed us to test the utility of panRAS and K-RAS antibodies in detecting K-RAS by western blotting. However, our results also prompt cautious use of GFP-tagged K-RAS, as A549GFPKRAS exhibit a drastically altered phosphorylation status for downstream targets MEK1/2, ERK1/2, and AKT, as well as EGFR. This, of course, might be related to clonal variation; however, in screening for GFP-positive cells following CRISPR-mediated GFP-K-RAS knockin, we only obtained one viable clone, perhaps hinting at a low tolerance for the presence of GFP on endogenous K-RAS. Beyond the plasma membrane localization, we observed additional disperse cytoplasmic signals of endogenous GFP-K-RAS, but no mitochondrial localization. When overexpressed, K-RASG12V has been suggested to be transported into mitochondria, leading to alterations of membrane potential, a decrease in respiration, and an increase in glycolysis (Hu et al., 2012). Potential compartments for the observed cytosolic signal for K-RAS could be Golgi, as seen for H- and N-RAS (Goodwin et al., 2005), which could correspond to K-RAS4A signal, or ER. However, this remains to be verified.

In this report, we demonstrate that endogenous K-RAS and H-RAS proteins can be targeted for degradation using the proteolytic AdPROM system. RAS proteins have remained elusive targets for anti-cancer therapies, primarily due to their undruggability (Cox et al., 2014). Research into obtaining small-molecule inhibitors of K-RAS has been carried out for over 30 years without much success (Cox and Der, 2010). Recently, RAS-targeting small molecules have emerged, with specificities to (1) a specific mutation status of K-RAS (G12C), i.e., ARS-1620 (Janes et al., 2018), and ARS-853 (Patricelli et al., 2016); (2) K-RAS, independent of the mutation status (McCarthy et al., 2019); or (3) RAS proteins in either nucleotide bound state (Kessler et al., 2019). Two compounds targeting K-RASG12C mutation, AMG510 and MRTX849, are currently undergoing clinical trials (Lindsay and Blackhall, 2019). An alternative approach has been the development of high-affinity polypeptide binders of RAS that neutralize the RAS function. A class of binders based on ankyrin repeat proteins (Guillard et al., 2017) can bind and neutralize specific nucleotide loading states of RAS proteins (Guillard et al., 2017). Similarly, a fibronectin type III domain-based RAS binding monobody (Khan et al., 2019b; Koide et al., 1998; Spencer-Smith et al., 2017, 2019) was shown to bind and inhibit the dimerization of both K- and H-RAS, and the overexpression of this monobody was shown to suppress tumor growth in mice (Khan et al., 2019b). Besides inhibition, RAS degradation offers another alternative approach at inhibiting RAS function to target RAS-dependent cancer cells. In this context, the dTAG-13 proteinase targeting chimera (PROTAC) was used to degrade FKBP12G38K-tagged K-RAS (Nabet et al., 2018) through the UPS, albeit when overexpressed in cells. Our AdPROM system demonstrates that endogenous RAS proteins can be targeted for proteolysis through the UPS and suggests that pharmacological targeting of RAS proteins for proteosomal degradation is a viable option for intervention. Although targeted delivery of polypeptide binders of RAS proteins or the proteolytic AdPROM system into RAS-dependent cancer cells remains challenging and currently offers limited therapeutic potential, these are excellent tools to provide the proof of concept. Further optimization of efficient AdPROM gene or protein delivery technologies might enable the study of short-term responses in downstream signaling of RAS. In the clinic and for a thorough analysis of degradation kinetics, cell-permeable small-molecule PROTACs are more viable options than the current AdPROM system, as it relies on long antibiotic selection for transduced cells and in its current form is not tractable. Recently two allosteric small-molecule binders were described for K-RAS with micromolar and nanomolar binding affinities (Kessler et al., 2019; McCarthy et al., 2019). It would be important to test these binders’ capabilities as K-RAS targeting warheads in a PROTAC approach; similar to the recently published ones (Bond et al., 2020; Zeng et al., 2020). In this context, a re-evaluation of RAS binding molecules, with or without inhibitory function, might prove successful for PROTAC design. Work published while this study was under review utilizing a KRASG12C-specific PROTAC harboring a CRBN recruiting warhead was able to
demonstrate the degradation of GFP-K-RAS, but the PROTAC was unable to degrade endogenous or untagged K-RAS (Zeng et al., 2020). Based on these results, this study suggested that targeting endogenous K-RAS for degradation by the proteasome would be difficult, if not impossible. However, our study here demonstrates that VHL-aHRAS AdPROM clearly targets both endogenous H- and K-RAS proteins for degradation through the proteasome. Moreover, a very recent preprint study has demonstrated that a VHL-recruiting K-RASG12C-specific PROTAC is indeed able to degrade endogenous K-RASG12C in different cell lines, although its impact on the viability of different cell lines after PROTAC treatment over the K-RASG12C inhibitors is yet to be addressed (Bond et al., 2020). It has been shown that cells that undergo long-term PROTAC treatment can gain resistance mutations in the E3 ligases or their receptors, which stop the PROTAC-E3 interaction (Zhang et al., 2019). In the search for other E3 ligases applicable for PROTAC development, the AdPROM system represents a rapid research tool with the potential to screen the efficacy of different E3 ligases, or RAS peptide recruiters in degradation of RAS. In addition, our A549GFPKRAS cells provide an excellent high-throughput screening platform to test the efficacy of either new E3 warheads that are compatible with RAS degradation, or new RAS-recruiting warheads.

The field of targeted RAS proteolysis is gaining momentum and presents different approaches for different applications. While the dTAG system offers strong, selective, and more importantly inducible degradation of POIs, it relies on the fusion of FKBP12F36V to the N terminus of RAS (Nabet et al., 2018), which is central to the research focus of our laboratory. In our system, it is unclear whether ubiquitination occurs on RAS itself, and/or the monobody, although complex formation would suggest that, within the AdPROM setup, VHL and the monobody are further from the RBX1 ubiquitination zone than bound RAS. In the same line of reasoning, N-terminal GFP of GFP-K-RAS would be even closer to RBX1, which might explain the strong degradation of GFP-K-RAS with VHL-aHRAS. Nonetheless, our study strongly suggests that different high-affinity polypeptide binders that can selectively bind either specific RAS proteins or mutants can be packaged with VHL-AdPROM to target specific RAS proteins for pro teaseomal degradation. At the same time, endogenous VHL substrates are not affected by the apparent overexpression of VHL (Table 1). We also noted that aHRAS monobody alone resulted in a marked stabilization of both H-RAS and K-RAS in multiple cell lines (Figures 4A and 5B). This effect could be caused either by a feedback loop induced by the inhibition of both RAS species imparted by aHRAS binding, or by blocking the natural turnover pathway through binding the RAS dimerization interface at helical structures 4α–5α (Spencer-Smith et al., 2017).

For the cell lines that we used, AdPROM-mediated degradation of H-/K-RAS was not sufficient to induce inhibition of anchorage-dependent cell proliferation. Despite harboring an activating K-RASG12C mutation, A549 cells do not appear to be strictly dependent on K-RAS alone in anchorage-dependent growth. While A549 cells are often discussed to be K-RAS independent (Kazi et al., 2018; Singh et al., 2009; Symonds et al., 2016), expression of MIR-181a-5p, a microRNA targeting the K-RAS 3’ UTR, reduced A549 anchorage-dependent proliferation and migration. However, MIR-181a-5p does not target K-RAS selectively (Ma et al., 2015). Many RAS-dependent cell proliferation assays use anchorage-independent 3D cultures. For example, the K-RASG12C drug ARS-1620 was shown to be effective at inhibiting RAS-dependent cell proliferation in 3D cultures but not in 2D cultures (Janes et al., 2018). In a similar manner, EGFR inhibitors show anti-proliferative effects in A549 cells only in an anchorage-independent growth context (Jaramillo et al., 2008). In contrast, SW620 cells, which are considered to be K-RAS dependent (Kazi et al., 2018; Singh et al., 2012), were inhibited in anchorage-dependent proliferation by aHRAS monobody alone. VHL-aHRAS AdPROM, which caused no detectable degradation of K-RAS in these cells, did not inhibit their proliferation any further. The inhibition of cell proliferation of RAS-dependent cells by aHRAS monobody is consistent with previous reports (Khan et al., 2019b; Spencer-Smith et al., 2017). The lack of degradation proteins, as the initially described binding preference of the mono body to H-/K-RAS, but not N-RAS (Spencer-Smith et al., 2017), is reflected in different levels of degradation when used in the AdPROM system (Table 1). Nevertheless, VHL-aHRAS-mediated RAS degradation was enough to elicit a cellular response to the removal of RAS proteins from a cell. Our data indicate a responsive upregulation of both M-RAS and TAF4B in A549 cells. Overexpression of M-RAS with activating mutations has been shown to lead to ERK signaling activation and transformation of cells (Quilliam et al., 1999). TAF4B, on the other hand, is usually associated with oocyte development (Fa lender et al., 2005a) and spermatogenesis (Falender et al., 2005b), and has not been associated with RAS function before. However, these findings need to be corroborated further.

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of K-RAS by VHL-aHRAS AdPROM could be due to the unusual size shift of K-RAS in these cells, possibly caused by a post-translational modification or a mutation that might allow binding to aHRAS monobody but prevent ubiquitination by the VHL-AdPROM, although this needs to be defined further. However, considering the length of the transduction and the antibiotic selection process the current AdPROM system uses, at the time of proliferation tests, only cells that have overcome the antiproliferative effects of RAS degradation might be selected. Therefore, to assess the effects of AdPROM-mediated degradation of H-/K-RAS on proliferation robustly, it will be essential to first obtain high-affinity polypeptide RAS binders that bind to specific RAS proteins and then use them in RAS-dependent cell lines using a tractable AdPROM system by either delivering AdPROM proteins or a chemically inducible AdPROM system.

SIGNIFICANCE

Our findings demonstrate clearly that endogenous RAS proteins can be targeted for proteasomal degradation by using the AdPROM system. The system unequivocally informs that targeted proteolysis of endogenous K-RAS is a viable strategy to target K-RAS-dependent pathologies. The findings open up exciting opportunities to develop VHL-recruiting K-RAS-specific cell-permeable PROTACs as potential therapeutic agents. Our findings also highlight the need for developing better and more selective RAS binding polypeptides, such as nanobodies or monobodies, to achieve more selective degradation with the AdPROM system.

STAR METHODS

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SUPPLEMENTAL INFORMATION

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AUTHOR CONTRIBUTIONS

T.J.M. generated all plasmids used in this study. S.R., A.K., M.A.Q., K.-H.C., and G.P.S. designed the project. S.R. and G.P.S. drafted the manuscript. S.R. acquired and analyzed the data. S.R. and G.P.S. interpreted the data. H.Z. performed mass spectrometric analysis of samples.

DECLARATION OF INTERESTS

M.A.Q., A.K., and K.-H.C. are employees and shareholders of GlaxoSmithKline.

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# STAR★METHODS

## KEY RESOURCES TABLE

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| **Antibodies**      |        |            |
| Alpha tubulin Monoclonal antibody (YOL1/34) | Thermo Fisher | Cat# MA1-80189; RRID: AB_2210200 |
| Goat anti-rat IgG Secondary Antibody (HRP) | Thermo Fisher | Cat# 31470; RRID: AB_228356 |
| B-RAF Rabbit Monoclonal Antibody (7H30L21) | Thermo Fisher | Cat# 702187; RRID: AB_2633065 |
| Donkey Anti-Rabbit IgG, AlexaFluor 488 | ThermoFisher | Cat# A21206; RRID: AB_2535792 |
| Goat anti-Mouse IgG, AlexaFluor 594 | ThermoFisher | Cat# A11005; RRID: AB_2534073 |
| Anti-KRAS+HRAS+NRAS antibody [EPR18713-13] | Abcam | Cat# Ab206969 |
| Anti-HIF-1 alpha antibody [H1alpha67] | Abcam | Cat# Ab1; RRID: AB_296474 |
| Anti-ATPB antibody [3D5] | Abcam | Cat# Ab14730; RRID: AB_301438 |
| Monoclonal Anti-KRAS antibody | SigmaAldrich | Cat# VH0003845M1; RRID: AB_1842235 |
| Monoclonal Anti-Flag M2-Peroxidase (HRP) antibody | SigmaAldrich | Cat# A8592; RRID: AB_439702 |
| Anti GFP from mouse IgG1k (clones 7.1 and 13.1) | SigmaAldrich | Cat# 11814460001; RRID: AB_390913 |
| Monoclonal Anti-FLAG M2 antibody | SigmaAldrich | Cat# F1804; RRID: AB_262044 |
| GAPDH (12C10) rabbit mAb | CST | Cat# 2118S; RRID: AB_561053 |
| Phospho-p44/42 MAPK (E10) Mouse mAb | CST | Cat# 9106S; RRID: AB_331768 |
| MEK1/2 (L38C12) Mouse mAb | CST | Cat# 4694S; RRID: AB_10695868 |
| Phospho-MEK1/2 (Ser221) (166F8) Rabbit mAb | CST | Cat# 2338S; RRID: AB_490903 |
| AKT Antibody | CST | Cat# 9272S; RRID: AB_329827 |
| Phospho-Akt (Ser473) (E7C3) (D9W9U) Mouse mAb | CST | Cat# 12694S; RRID: AB_2797994 |
| Phospho-EGF Receptor (Tyr1068) (D7A5) XP Rabbit mAb | CST | Cat# 3777; RRID: AB_2096270 |
| C-Myc (D84C12) Rabbit mAb | CST | Cat# 5605; RRID: AB_1903938 |
| Anti-rabbit IgG, HRP-linked Antibody | CST | Cat# 7074S; RRID: AB_2099233 |
| Anti-mouse IgG, HRP-linked Antibody | CST | Cat# 7076S; RRID: AB_330924 |
| EGF Fr (1005)-G Antibody | SantaCruz | Cat# sc-03-G; RRID: AB_631420 |
| StarBright Blue 700 Goat Anti-Rabbit IgG | BioRad | Cat# 123014; RRID: AB_2721073 |
| NRAS Rabbit polyclonal antibody | Proteintech | Cat# 10724-1-AP; RRID: AB_2154209 |
| HRAS Rabbit-Polyclonal Antibody | Proteintech | Cat# 18295-1-AP; RRID: AB_2121046 |
| KRAS-2B Rabbit Polyclonal Antibody | Proteintech | Cat# 16155-1-AP; RRID: AB_2134119 |
| KRAS-2A Rabbit Polyclonal Antibody | Proteintech | Cat# 16156-1-AP; RRID: AB_2234477 |
| H-RAS Polyclonal Antibody | Invitrogen | Cat# PA5-22392; RRID: AB_11152295 |
| K-RAS Monoclonal Antibody (9.13) | Invitrogen | Cat# 415070; RRID: AB_2532192 |
| GFP Polyclonal Antibody | MBL/Caltag | Cat# 508; RRID: AB_591819 |
| Mouse Anti-p120 Catenin Antibody [clone 9B9/PP120] | BD Biosciences | Cat# 610133; RRID: AB_397536 |

### Chemicals, Peptides, and Recombinant Proteins

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| Cycloheximide | SigmaAldrich | Cat# C1988 |

(Continued on next page)
| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| Bortezomib          | LC Laboratories | Cat# B-1408 |
| MG132               | abcam   | Cat# Ab141003 |
| PEI MAX – Transfection Grade Linear PEI Hydrochloride MW 40,000 | Polysciences | Cat# 24765 |
| Polybrene (Hexadimethrine bromide) | SigmaAldrich | Cat# 107689 |
| GFP-Trap-Agarose    | Chromotek | Cat# GTA-20; RRID: AB_2631357 |
| Anti-Flag M2 Affinity Gel | Merck | Cat# A2220; RRID: AB_10063035 |
| Vectashield Antifade mounting medium | Vector Labs | Cat# H-1000; RRID: AB_2336789 |
| Lys-C Protease, MS Grade | Alpha Labs | Cat# 125-05061 |
| Pierce Trypsin Protease, MS Grade | ThermoFisher | Cat# 90058 |
| Immobilon Western Chemiluminescent HRP Substrate | Merck | Cat# WBKLS0500 |
| Critical Commercial Assays | | |
| iScript cDNA synthesis Kit | Biorad | Cat# 1708891 |
| SsoFast EvaGreen Supermix | Biorad | Cat# 1725204 |
| TMT10plex Isobaric Label Reagent Set | ThermoFisher | Cat# 90110 |
| Deposited Data | | |
| Results from TMT9plex labelling and MS analysis, see Table S1–A549-ADPROM-TMT9plex-Related to Figure 4D | This paper | N/A |
| Data obtained in this study | This paper | osf.io/zm3dx |
| Experimental Models: Cell Lines | | |
| A549                | ATCC    | Cat# CCL-185; RRID: CVCL_0023 |
| A549GFPKRAS         | This paper | N/A |
| HEK293-FT           | Invitrogen | Cat# R70007 |
| A375                | ATCC    | Cat# CRL-1619; RRID: CVCL_0132 |
| A172                | ATCC    | Cat# CRL-1620; RRID: CVCL_0131 |
| HT-29               | ATCC    | Cat# HTB-38; RRID: CVCL_0320 |
| HPAFII              | ATCC    | Cat# CRL-1997; RRID: CVCL_0313 |
| H460                | ATCC    | Cat# HTB-177; RRID: CVCL_0459 |
| SW620               | ATCC    | Cat# CCL-227; RRID: CVCL_0547 |
| Oligonucleotides | | |
| Primers for qRT-PCR, Screening & Sequencing see Table S2– Primer Sequences – Related to STAR Methods | This paper | N/A |
| Recombinant DNA | | |
| pBabeD P U6 KRAS Nter KI Sense | This paper; MRCPPU Reagents and Services | Cat# DU54976 |
| pX335 KRAS Nter KI AntiSense | This paper; MRCPPU Reagents and Services | Cat# DU54980 |
| pMK-RQ KRAS G12S Nter GFP donor | This paper; MRCPPU Reagents and Services | Cat# DU57406 |
| pBABED P FLAG aHRAS nanobody | This paper; MRCPPU Reagents and Services | Cat# DU57190 |
| pBABED P FLAG VHL aHRAS nanobody | This paper; MRCPPU Reagents and Services | Cat# DU57191 |
| pBABED P FLAG VHL | Fulcher et al. 2017; MRCPPU Reagents and Services | Cat# DU54477 |
| pBABED P aGFP16 | Fulcher et al. 2016, MRCPPU Reagents and Services | Cat# DU54238 |

(Continued on next page)
RESOURCE AVAILABILITY

Lead Contact
Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact, Gopal Sapkota (G.Sapkota@dundee.ac.uk).

Materials Availability
Plasmids generated in this study can be obtained from MRC PPU Reagents and Services (https://mrcppureagents.dundee.ac.uk).

Data and Code Availability
Original data have been deposited to the Center for Open Science repository: osf.io/zm3dx

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Cell Lines
A549GFPKRAS cells were derived from the epithelial lung cancer cell line A549 by CRISPR/Cas9 mediated knock-in of GFP CDS to the KRAS locus. A549 cells were derived from a 58 year old Caucasian male. A375 cells are a malignant melanoma cell line from a 54 year old female. A172 cells are glioblastoma cells from a 53 year old male. SW620 cells are Duke’s type C colorectal adenocarcinoma cells derived from the lymph node metastatic site of a 51 year old Caucasian male. HT-29 cells are colorectal adenocarcinoma cells derived from primary tumor from a 44 year old Caucasian female. HPAFII cells are pancreatic adenocarcinoma cells derived from peritoneal ascitic fluid of a 44 year old Caucasian male. H460 cells are large cell lung cancer cells derived from pleural effusion of a male patient. HEK293-FT cells are a clonal isolate of HEK293 human embryonic kidney cells transformed with the SV40 large T antigen. All cells were cultured in humidified incubators at 37°C and 5% CO2. A549, HEK293-FT, A375, A172 and SW620 cells were cultured in Dulbecco’s modified Eagle’s medium (DMEM; Gibco) with 10% FBS (Sigma), 1% penicillin/streptomycin (Lonza) and 2 mM L-glutamine (Lonza). HT-29, HPAFII and H460 cells were cultured in RPMI1640 medium (Gibco), with the same supplements as DMEM.

METHOD DETAILS

Sequence Alignment
Protein sequences of K-RAS4A/B, H-RAS and N-RAS were taken from Uniprot (The UniProt Consortium, 2019) and aligned in Clustal Omega (Madeira et al., 2019). The alignment was further processed in JalView (Waterhouse et al., 2009) to highlight percent sequence identity.

RNA Extraction, cDNA Synthesis and qRT-PCR
For RNA extraction, 2x10^6 cells were seeded in a 6-well dish and harvested the next day with the RNeasy Micro Kit (Qiagen, #74004) according to the manufacturer’s protocol. 1 μg of RNA was reverse transcribed with the iScript cDNA synthesis Kit (BIORAD, #1708891) according to the manufacturer’s protocol. For qRT-PCR 1 μl of diluted cDNA (1:20 or 1:80) was mixed with forward and reverse primers (Custom primers from Invitrogen, 300 nM final concentration each) and SsoFast EvaGreen Supermix (BIORAD, #1725204) in a 384-well plate (Axygen, #321-22-051) and run on a BIORAD CFX384.
Primer sequences:

K-RAS4A fw: GAGGGAGATCCGACAAATACAG;
K-RAS4A rev: TCTCGAATCTGATAGAAGGATC;
K-RAS4Bfw: TTGCTCTCTAGACACAGTACAC;
K-RAS4B rev: CATCGTCAAACCTGCTTGTG;
Total K-RAS fw: GGAGTACAGTGAACTGAGGG;
Total K-RAS rev: CCATAGTGATCTTCAAGTCC;
H-RAS fw: GAAACAGTGGACAGCTTGG;
H-RAS rev: ACCAAGTGTAAGGAGCAGC;
N-RAS fw: AATACATGAGACAGGAGCAGAAG;
N-RAS rev: GTCGCTCTAGACAGAACTAG;
GAPDH fw: CTGTGTAAGCTATTCTCGT;
GAPDH rev: CTCTCCCTTGTGCTCTTGC.

Following the PCR, melting curves were generated with default settings between 65°C and 95°C in 0.5°C steps at 5 sec intervals. Melting curves were manually analysed for purity of the PCR product, i.e. consistency of amplicon melting temperature between different samples and peak distribution. Fold changes of transcripts were calculated by the 2-ΔΔCt method (Livak and Schmittgen, 2001).

Cell Line Transfection and Transduction

For retrovirus production, 3.2 μg pCMV-gag-pol (Cell Biolabs, RV-111), 2.2 μg pCMV-NSV-G (Cell Biolabs, RV-110) and 6 μg of respective pBabeD plasmids (Flag-aHRAS, DU57190; Flag-VHL-aHRAS, DU57191; Flag-VHL, DU54477; aGFP16, DU54238; Flag-VHL-aGFP16, DU54295) were co-transfected in roughly 70% confluent HEK293-FT cells cultured on a 10-cm dish. Plasmids were mixed with 600 μl Opti-MEM (Gibco) and 24 μl of 1 mg/ml polyethyleneimine (Polysciences) dissolved in 25 mM HEPES pH 7.5. The mixture was vigorously vortexed for 15 s and incubated for 20 min at room temperature. The volume was adjusted to 10 ml with DMEM and added to FT cells. After 24 h, medium was exchanged to DMEM or RPMI, depending on the target cell growth medium. After an additional 24 h, the medium was harvested and filtered through a 0.45 μM Minisart syringe filter (Sartorius). The supernatant was added to a plate of roughly 70% confluent target cells in a 1:10–1:4 dilution (in respective medium) in the presence of 8 μg/ml polybrene (Sigma). For retrovirus production, 2 μg pCMV-gag-pol (Cell Biolabs, RV-111), 2.2 μg pCMV-NSV-G (Cell Biolabs, RV-110) and 6 μg of respective pBabeD plasmids (Flag-aHRAS, DU57190; Flag-VHL-aHRAS, DU57191; Flag-VHL, DU54477; aGFP16, DU54238; Flag-VHL-aGFP16, DU54295) were co-transfected in roughly 70% confluent HEK293-FT cells cultured on a 10-cm dish. Plasmids were mixed with 600 μl Opti-MEM (Gibco) and 24 μl of 1 mg/ml polyethyleneimine (Polysciences) dissolved in 25 mM HEPES pH 7.5. The mixture was vigorously vortexed for 15 s and incubated for 20 min at room temperature. The volume was adjusted to 10 ml with DMEM and added to FT cells. After 24 h, medium was exchanged to DMEM or RPMI, depending on the target cell growth medium. After an additional 24 h, the medium was harvested and filtered through a 0.45 μM Minisart syringe filter (Sartorius). The supernatant was added to a plate of roughly 70% confluent target cells in a 1:10–1:4 dilution (in respective medium) in the presence of 8 μg/ml polybrene (Sigma). After 24 h, growth medium was exchanged with fresh medium containing 2 μg/ml puromycin, to select transduced cells. Puromycin was removed from the medium after 48 h. For inhibitor experiments cells were treated with cycloheximide (100 μg/ml; Sigma-Aldrich, C1888), MLN4924 (1 μM, 24 hours, MRC-PPU Reagents and Services), MG132 (40 μM, 14 hours, abcam, ab141003), Bortezomib (10 μM, 14 hours, LC Laboratories, B-1408) or DMSO (adjusted to match respective inhibitor; Sigma-Aldrich, D2650).

Cells were lysed on ice, by washing once with PBS and scraping in lysis buffer (50 mM Tris–HCl pH 7.5, 0.27 M sucrose, 150 mM NaCl, 1 mM EGTA, 1 mM EDTA, 1 mM sodium orthovanadate, 1 mM sodium β-glycerophosphate, 5 mM sodium fluoride, 5 mM sodium pyrophosphate, 1% (v/v) Triton X-100 and 0.5% Nonidet P-40) supplemented with protease inhibitors (Roche; 1 tablet/20 ml of lysis buffer). Protein content from cleared cell lysates was determined with Pierce Detergent Compatible Bradford Assay Kit (Thermo Fisher). Lysates were processed further or frozen and stored at -20°C.

CRISPR/Cas9

For generation of N-terminal GFP knock-in A549 cell lines the KRAS locus was targeted with a dual guide approach (Fulcher et al., 2019) (using the sense guide (pBabeD vector, DU54976): GCGAATATGATCCAACAATAG; antisense guide (pX335 vector, DU54980): GCTGAATTAGCTGTATCGTCA; and the GFP-KRAS donor (pMK-RQ vector, DU57406). Briefly, 1 μg of each of the guideRNA plasmids and 3 μg of the donor plasmid were co-transfected into A549 cells. Plasmids were mixed with 1 μl of Opti-MEM (Gibco) and 20 μl of 1 mg/ml polyethyleneimine (Polysciences), vortexed vigorously for 15 s and added to 70% confluent cells in a 10-cm dish. The next day, cells were selected in puromycin (2.5 μg/ml) for 48 h and re-transfected with the same plasmids once they reached 70% confluence. Next generation positive cells were obtained through FACS sorting and surviving single cell clones were screened by genomic DNA based PCR and Western blot to validate homozygous knock-in of the GFP-tag on the endogenous KRAS gene. For PCR based screening the following primers were used: Fw: ATCCAGAGACTACTGCTGACAGC; Rv: CATGACCTCAAGTGTCCTACAGGTG. PCR products of positive clones were cloned with the StrataClone PCR Cloning Kit (Agilent) into the supplied vector system, according to the manufacturer’s protocol. Sequencing of positive clones was carried out by the MRC-PPU DNA Sequencing and Services with a custom primer close to the RAS mutation site (Rv: CAAAGAATGGTCCTGCACCAG).

SDS PAGE and Western Blotting

Cell lysates were adjusted to uniform protein concentration and mixed with 6x reducing Laemmli SDS sample buffer (Fisher Scientific). 10-20 μg of total lysate protein, or immunoprecipitates were resolved by SDS polyacrylamide gel electrophoresis (PAGE). After PAGE, proteins were transferred onto methanol activated PVDF membrane (Immobilon-P or Immobilon-FL, Merck) in Tris/glycine buffer containing 20% methanol in a tank blotting system for 85 min at a constant voltage of 85 V. The membranes were then re-incubated with methanol for 2 minutes and stained with Ponceau S solution to gauge uniform protein transfer (Sigma). After de-staining membranes in TBS-T (50 mM Tris–HCl pH 7.5, 150 mM NaCl, 0.1% Tween-20), they were blocked for 1 h in 5% non-

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fat milk (Marvel) in TBS-T. Primary antibody incubation was done overnight at 4°C in 5% milk/TBS-T. Following 3x10 min washes in TBS-T, membranes were incubated with respective HRP-conjugated (CST) or fluorescently labelled (Biorad) secondary antibodies for 1 h, washed again 3x10 min in TBS-T and developed on a ChemiDoc gel imaging system (Biorad) using the respective channels. HRP-conjugated blots were incubated with Immobilon Western Chemiluminescent HRP Substrate (Millipore).

**Immunoprecipitation**

Cell lysates were adjusted to 1 µg/µl in lysis buffer. Either GFP-trap beads (ChromoTek) or Anti-FLAG-M2-Affinity agarose resin (SigmaAldrich) was equilibrated with lysis buffer. 300-500 µg of total protein was added to 10-15 µl of beads (50% slurry) and incubated for an hour at 4°C under agitation. Centrifugation steps at 200xg were done at 4°C for 2 minutes. Supernatant (flowthrough) was separated from beads, and beads were washed 3-5 times in lysis buffer. Proteins were eluted in lysis buffer containing Laemmli SDS sample buffer by boiling at 95°C for 5 minutes.

**Antibodies**

Antibodies were purchased from Thermo Fisher (Alpha tubulin, MA1-80189; rat-HRP, 31470; B-RAF, 702187), Abcam (panRAS, ab206969; HIF1α, ab1), Sigma (K-RAS4B, WH0003845M1; Flag-HRP, A8592-.2MG; GFP, 11814460001), CST (GAPDH, 2118S; ERK1/2, 9102S; phospho ERK1/2 (T202/Y204), 9106S; MEK1/2, 4694S; phospho MEK1/2 (S221), 2338S; AKT, 9272S; phosphor AKT (S473), 12694S; phospho EGFR receptor (Y1068), 3777; c-myc, 5605; rabbit-HRP, 7074S; mouse-HRP, 7076S), SantaCruz (EGF receptor, sc-03-G) and Bio-Rad (rabbit starbright 700, 12004161). Primary antibodies were generally used in 1:1,000 dilutions in 5% milk TBS-T, except from RAS (1:500), and GAPDH & alpha-tubulin (1:5,000). Secondary antibodies were used in a 1:5,000 dilution in 5% milk TBS-T. Other primary antibodies to detect RAS species were obtained from Proteintech (N-RAS, 10724-1-AP; H-RAS, 18295-1-AP; K-RAS2B, 16155-1-AP; K-RAS2A, 16156-1-AP) and Invitrogen (H-RAS, PA5-22392; K-RAS, 415700). Antibodies for immunofluorescence were purchased from MBL/Caltag Medsystems (GFP, 598), Abcam (ATPB, ab14730), BD Biosciences (P120 Catenin, 610133), Sigma (Flag-M2, F1804) and Thermo Fisher (AlexaFluor488 [donkey anti-rabbit], A21206; AlexaFluor594 [goat anti-mouse], A11005).

**Immunofluorescence**

Cells were seeded in a 12-well dish onto coverslips and grown over night. The next day, cells were washed twice in PBS and fixed for 10 minutes in 4% formaldehyde/PBS (Sigma). Coverslips were washed in DMEM (Gibco) containing 10 mM HEPES followed by a 10 min incubation. Coverslips were washed in PBS and permeabilised for 3 min in either 0.2% NP-40/PBS or 0.2% Triton X-100/PBS. Coverslips were washed twice in PBS and blocked for 15 min in 3% BSA (Sigma) in PBS. Primary antibody incubation was done for 1-2 h at room temperature. After appropriate antibody dilutions in blocking solution, residual antibody was washed away in 0.2% Tween/PBS (3x10 min). Secondary antibody incubation was done for 30 min at 1:300 antibody dilution in the dark. The same wash steps were repeated, but the first wash contained DAPI (0.5–1 µg in 10 ml, SigmaAldrich). Finally, coverslips were dipped in water, air dried and mounted on slides with Vectashield (Vector Laboratories). Fluorescence signals were analysed on a Deltavision Widefield microscope (GE). Images were deconvolved using the default settings of softWoRx Imaging software and further analysed using OMERo (Alian et al., 2012).

**Cell Proliferation Assays**

After trypsinization, live cell numbers were determined in a Neubauer haemocytometer in the presence of trypan blue. Cell numbers were adjusted to 5000 cells per ml in the respective growth medium. 5000 cells were added per well of a 12-well dish, and each line was grown in triplicates. After 7 days, relative cell numbers were determined by crystal violet staining. In short, cells were washed in PBS, fixed for 5 min in fixing buffer (10% methanol, 10% acetic acid), washed in PBS again and incubated for 30–60 min in crystal violet solution (0.5% crystal violet in 20% methanol). Plates were then incubated in a Licor Odyssey using the 700 nm channel. Subsequently, 1 ml methanol was added to each well and plates were incubated shaking for 30 min. Depending on the colour of 1 set of cells, 100-200 µl of supernatant was loaded in triplicate on a 96-well plate and absorbance at 570 nm was measured in an Epoch microplate spectrophotometer (BioTek). Values were normalized to the untreated sample and a one-way ANOVA analysis with Dunnett’s multiple comparisons test was done.

**Flow Cytometric Analysis**

Cells were trypsinized, washed and resuspended in PBS containing 1% FBS. Cells were then analysed on a FACS Canto II flow cytometer. Cells were analysed with the following gating strategy: (i) cells: in a plot of FSC-A vs. SSC-A, a gate was drawn surrounding the major population of cells, removing debris and dead cells. (ii) single cells: in a plot of FSC-A vs. FSC-W, a gate was drawn around an area corresponding to single cells. (iii) in the ‘single cells’ population on a GFP-A vs. PE-A plot a gate was drawn around GFP-positive cells in A549GFPKRAS sample, using WT A549 cells as a negative control. Gates (i) and (ii) were adjusted to the individual cell lines. Gate (iii) was kept unchanged within an experiment.

**Sample preparation for Tandem Mass Tag (TMT) Labelling**

Transduced and selected A549 cells were processed for TMT labelling as described previously (Tovell et al., 2019). In short, samples were lysed in 8 M Urea and 50 mM Ammonium bicarbonate containing buffer, cleared after benzonase treatment, reduced with 5 mM DTT at 45°C for 30 min, alkylated with 10 mM iodoacetamide at room temperature in the dark for 20 min), quenched by addition of 5 mM DTT, digested with Lys-C (1:200 (w/w), Lys-C/protein) for 4 h at 30°C, diluted with 50 mM Ammonium bicarbonate to 1.5 M final
Urea concentration, followed by trypsin digestion (1:50 (w/w), trypsin/protein) at room temperature overnight. 1% TFA was added to stop the digestion. The acidified digest samples were desalted on 200 mg Sep-Pak tC18 cartridges, and the eluents were dried by using speed vacuum centrifugation (Thermo). Tandem Mass Tag labelling was performed according to the manufacturer’s protocol using the TMT Labelling Kit (Thermo, 90110). The complete labelled samples were then mixed andfractionated with high pH reverse phase C18 chromatography using an Ultimate 3000 high-pressure liquid chromatography system (Dionex) at a flow rate of 569 µl/min using two buffers: A (10 mM ammonium formate, pH 10) and B (80% ACN, 10 mM ammonium formate, pH 10). Briefly, the desalted TMT labelled peptides were resuspended in 200 µL of buffer A (10 mM ammonium formate, pH10) and fractionated on a C18 reverse phase column (4.6 × 250 mm, 3.5 µm, Waters) with a gradient as follows: 3% B to 12.5 % B in 10 min, 12.5% to 40% buffer B in 45 min, 40% B to 60% B in 25 min, 60% B to 80% B in 10 min, 80% B to 100% B in 2.5 min, 100% B for 5 min, ramping to 3% B in 2.5 min and then 3% for 10 min. A total of 90 fractions were collected and then concatenated into 30 fractions, which were further desalted over C18 StageTips and speed vacuum dried prior to LC-MS/MS analysis.

**LC-MS/MS Mass Spectrometry**

LC-MS/MS analysis was done as described previously (Tovell et al., 2019), with a Thermo Dionex Ultimate 3000RSLC Nano liquid chromatography instrument. Peptides were quantitated by Nanodrop and the sample was dissolved in 0.1% formic acid. 1 µg of each fraction was loaded on C18 trap column with 3% ACN/0.1% TFA at 5 ul/min flow rate. Peptides were separated over an EASY-Spray column (C18, 2µm, 75µm x 50cm) with an integrated nano electrospray emitter (flow rate 300nl/min). Peptide separation was done over 180 min with a uniform gradient: the first 10 fractions starting from 5%~30% buffer B in 125 min (Note: the middle 10 fractions starting from 7% and the last 10 fractions starting from 10%), 30%~45% buffer B in 30 min, 45%~95% buffer B for 5 min, followed by a 5 min 95% B. Eluted peptides were analysed on an Orbitrap Fusion Lumos (Thermo Fisher Scientific, San Jose, CA) mass spectrometer. Spray voltage was set to 2 kV, RF lens level was set at 30%, and ion transfer tube temperature was set to 275 °C. The Orbitrap Fusion Lumos was operated in positive ion data-dependent mode with synchronous precursor selection (SPS)-MS3 analysis for reporter ion quantitation. The mass spectrometer was operated in data-dependent Top speed mode with 3 seconds per cycle. The full scan was performed in the range of 350–1500 m/z at nominal resolution of 120 000 at 200 m/z and AGC set to 4x105 with maximal injection quantitation. The Orbitrap Fusion Lumos was operated in positive ion data-dependent mode with synchronous precursor selection (SPS)-MS3 analysis for reporter ion quantitation. The mass spectrometer was operated in data-dependent Top speed mode with 3 seconds per cycle. The full scan was performed in the range of 350–1500 m/z at nominal resolution of 120 000 at 200 m/z and AGC set to 4x105 with maximal injection quantitation. The isolation width was set to 0.7 m/z with no offset. Dynamic exclusion was set to 60 seconds. Monoisotopic precursor selection was set to peptide, maximum injection time was set to 50 msec. Charge states between 2 to 7 were included for MS2 fragmentation. The top 5 fragment ions from each MS2 scan was notched out for MS3. The MS3 scan were performed with an isolation width of 2 m/z in the quadrupole, normalised HCD collision energy of 65% and analysis of fragment ions in the orbitrap using 50 000 resolving power with auto normal range scan from m/z 100 to 500 and AGC target of 5x105. The maximal injection time for MS3 scan was set to 86 ms.

**LC-MS/MS Data Analysis**

LC-MS/MS data analysis was done as described previously (Tovell et al., 2019). All acquired LC-MS data were analysed using Proteome Discoverer software v.2.2 (Thermo Fisher Scientific) with Mascot search engine. Maximum missed cleavages for trypsin digestion was set to 2. Precursor ion tolerance was set to 20 ppm. Fragment ion tolerance was set to 0.6 Da. Carbamidomethylation on cysteine (+57.021 Da) and TMT-10plex tags on N termini as well as lysine (+229.163 Da) were set as static modifications. Variable modifications were set as oxidation on methionine (+15.995 Da). Data were searched against a complete UniProt Human Proteome (Reviewed 20,143 entry downloaded at Nov 2018). Peptide spectral match (PSM) error rates with a 1% FDR were determined by target-decoy strategy coupled to Percolator modelling of true and false matches. Both unique and razor peptides were used for quantitation. Reporter ion abundances were corrected for isotopic impurities based on the manufacturer’s data sheets. Reporter ions were quantified from MS3 scans using an integration tolerance of 20 ppm with the most confident centroid setting. Signal-to-noise (S/N) values were used to represent the reporter ion abundance with a co-isolation threshold of 50% and an average reporter S/N threshold of 10 and above required for quantitation from each MS3 spectra to be used. The S/N value of each reporter ions from each PSM were used to represent the abundance of the localised phosphorylation sites. The precursor spectra with higher than 25% co-isolation were further manually checked. The total peptide amount was used for the normalisation. Protein ratios were calculated from medians of summed sample abundances of replicate groups. Standard deviations were calculated from three biological replicates. The standard deviation of three biological replicates lower than 25% were used for further analyses. To determine the significant differences between different treatments, ANOVA model was used for statistical significance analysis.

**QUANTIFICATION AND STATISTICAL ANALYSIS**

Statistical analysis was done using GraphPad Prism. For comparison of two groups, unpaired, two-tailed t-tests were performed. For comparison of more than two groups an Ordinary ANOVA with post-hoc Dunnett’s multiple comparisons test was performed. A p-value < 0.05 was considered statistically significant. Sample sizes are indicated in the respective figure legend.