SIRT2 deletion enhances KRAS-induced tumorigenesis in vivo by regulating K147 acetylation status

Supplementary Material

Mice

Sirt2 knockout mice (1) were crossed with the well-established LSL-Kras<sup>G12D</sup> knock-in mouse model (2) (provided by Dr. Harold L. Moses, Vanderbilt University) to generate Sirt2<sup>+/+</sup>:LSL-Kras<sup>G12D</sup> and Sirt2<sup>−/−</sup>:LSL-Kras<sup>G12D</sup> mice. These mice were next crossed with the Ptf1<sup>Cre</sup> driver line (3) (provided by Dr. Harold L. Moses, Vanderbilt University) to direct recombination in pancreas and generate Sirt2<sup>+/+</sup>:LSL-Kras<sup>G12D</sup>,Ptf1<sup>Cre</sup> and Sirt2<sup>−/−</sup>:LSL-Kras<sup>G12D</sup>,Ptf1<sup>Cre</sup> mice. Alternatively, the Sirt2<sup>+/+</sup>:LSL-Kras<sup>G12D</sup> and Sirt2<sup>−/−</sup>:LSL-Kras<sup>G12D</sup> mice were infected intranasally with an adenoCRE virus to direct recombination of the LSL-Kras<sup>G12D</sup> allele and initiate expression of Kras<sup>G12D</sup> in the lungs (4). To generate Sirt2 conditional knockout mice, embryonic stem (ES) cells were transfected with a targeting vector (containing floxed exons 5-8 of the Sirt2 gene as well as a floxed neo gene) by electroporation and selected using G418. ES clones were analyzed by both Southern blotting and PCR for cells with the targeted mutation. Next, cells were microinjected into blastocysts from C57BL/6 mice, and chimeric males were bred to C57BL/6 females to obtain germ-line transmission following standard procedures. Offspring carrying a loxP-neo-loxP cassette were crossed with Ella-Cre transgenic mice to delete the neo gene and generate Sirt2 floxed (Sirt2<sup>fl/fl</sup>) mice with loxP sites flanking exons 5-8 of the Sirt2 gene. Mice were genotyped using tail DNA with the following primers: (a) 5'-GCC TTA GCT ACA TAG AAG GC-3', (b) 5'-GAA TGA CCT ACA ATG GGC CA-3', and (c) 5'-GTG TAG CCC TGG CTC TTC TA-3'. These mice were crossed with the LSL-Kras<sup>G12D</sup> and Ptf1<sup>Cre</sup> mice in order to generate Sirt2<sup>fl/fl</sup>,LSL-Kras<sup>G12D</sup>,Ptf1<sup>Cre</sup> mice where exons 5-8 of the Sirt2 gene
are deleted and \textit{Kras}^{G12D} is expressed in the pancreas simultaneously. Mice were housed, fed, and treated in accordance with the guidelines approved by the Northwestern University IACUC.

**Immunohistochemistry**

For immunohistochemistry (IHC), sections were stained using the protocol described below. Briefly, slides were heated via pressure cooker in DAKO retrieval buffer, and endogenous peroxidases quenched in 3% hydrogen peroxide in methanol for 30 min. Tissues were blocked with 5% BSA in PBS for 30 min and exposed to primary antibodies against BrdU (Millipore), CK19 (DSHB, University of Iowa), Ac-KRAS-K147 (Eurogentec), pERK (Abcam and Cell Signaling), Ki67 (Abcam), thyroid transcription factor 1 (Abcam), and SIRT2 (Proteintech) at 1:100-1:200 overnight at 4 °C. Slides were developed using DAKO secondary antibodies and DAB substrate/buffer. Trichrome staining was performed per the manufacturer’s instructions (Sigma Aldrich).

**Plasmids/transfection/infection**

All plasmids for overexpressing SIRT2 have been described previously (1). Flag-tagged \textit{KRAS} and \textit{KRAS}^{G12V} constructs in pCMV2 vector were kindly provided by Lewis C. Cantley (Weill Cornell Medical College). HA-tagged \textit{KRAS} and \textit{KRAS}^{G12V} constructs in pGCN vector were provided by Mark R. Philips (New York University). Site-directed mutagenesis to make K147 acetylation mutants (K147R and K147Q) as well as subcloning to a pCDH-GFP-puro vector (SBI, Systems Biosciences) were performed by Bioinnovatise. For transient overexpression, 293T cells were transfected using polyethylenimine (PEI) at a ratio of 3 μL PEI/μg DNA. For stable overexpression, 293T cells were used to produce viral particles to infect cells (5). All cells
were selected after treating with puromycin (Cellgro) at 2 µg/mL or G418 (Cellgro) at 200 µg/mL for 2 weeks. For knocking down mouse Kras, human SIRT2 and mouse Sirt2, target sequences (Kras: 5’- GGATTTGGTGACATGCAGTTGA-3’, human SIRT2: 5’- AAGTAGTGACAGATGGTTGGC-3’ and mouse Sirt2: 5’-TTCCAGCTGTCTATGTCTG-3’) subcloned into a pLKO.1-puro vector (Dharmacon), were used to produce viral particles, followed by cell infection, and selection as described earlier. For siRNA experiments, control siRNA (sc-37007) and mouse siSIRT2 (sc-40989) were bought from Santa Cruz Biotechnology.

**Cell treatments**

To stimulate KRAS signaling, cells were first starved for 24 h in 0.1% FBS medium followed by treatment with EGF (100 ng/mL) (Sigma). To inhibit the activity of endogenous sirtuins, cells were treated with 2 µM nicotinamide (NAM) (Sigma) for 12 h before lysing the cells. To specifically inhibit SIRT2, cells were treated with 5 µM AGK2 (Santa Cruz) for 12 h before cell lysis.

**Intranasal infection**

AdenoCRE (2.5x10^7 particles, Gene Transfer Vector Core, University of Iowa) was first mixed with Minimum Essential Medium (MEM) and 2 M CaCl2 (final concentration 10 mM) in a final volume of 125 µL/mouse infected. Using a protein gel loading tip, half of the virus mixture (62.5 µL) was administered by placing the pipette tip at the opening of one nostril. The virus solution was slowly expelled from the tip while the mouse inhaled the drop that was forming. Mice were left to recover for 10-15 minutes. After breathing returned to normal, the procedure was repeated with the remaining 62.5 µL of the virus mixture. Mice were monitored for up to 6
months. All procedures were performed in accordance with approved Northwestern University IACUC protocol.

**Western blotting**

Cells or tissue samples were lysed using buffer A (25 mM Tris-HCl pH 7.5, 150 mM NaCl, 1 mM EDTA, 0.5% NP-40, and 5% glycerol), followed by sonication and centrifugation at 14,000 rpm for 25 min at 4 °C. After protein quantification using the Bradford assay, equal amounts of protein (20-40 µg) were mixed with 2× Laemmli lysis buffer, boiled for 5 min, separated on a denaturing SDS-polyacrylamide gel, and transferred to a PVDF membrane. The membrane was blocked in 5% milk/PBS/0.05% Tween for 1 h and incubated with antibodies against pERK, ERK (Cell Signaling), Ras (Thermo Scientific), KRAS (Proteintech), tubulin, actin, Flag, HA (Sigma), SIRT2 (Sigma and Proteintech), Ac-K (Millipore), and Ac-KRAS-K147 (Eurogentec). Membranes were washed with PBS/0.05% Tween and then incubated with HRP-conjugated secondary antibody (Santa Cruz Biotech, Santa Cruz, CA) at room temperature for 1 h. Protein banding was analyzed by an enhanced chemiluminiscence method (Amersham Biosciences, Piscataway, NJ) and resolved on x-ray film (Eastman-Kodak, Rochester, NY) per the manufacturer’s specifications. Quantification of western blot bands was performed using Gel Analyzer software.

**Cell proliferation**

For measuring proliferation of cells, three different methods were used. For the MTT- [3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide] proliferation assay, 5x10^3 cells/well were plated in 24-well plates. At different time points (0-6 days), MTT solution (BosterBio) was
added to the cells and after incubation for another 2 h, the resultant formazan crystals were dissolved in dimethyl sulfoxide (100 µL) and the absorbance intensity measured using an **xMark™ Microplate Absorbance Spectrophotometer** (Bio-RAD, USA) at 540 nm. All experiments were performed in triplicate, and the proliferation rate was expressed as a percentage relative to the cell absorbance at day 0. For counting the actual cell number, 5x10^4 cells/well were plated in 24-well plates. At different time points (0-6 days), cell proliferation was estimated after counting cell numbers using the **TC20™ Automated Cell Counter** (Bio-RAD, USA) following the manufacturer’s instructions. For colony forming ability, 100, 250, 400, 500 or 1000 cells per well were plated onto 6-well tissue culture plates in appropriate medium for each cell line used in this assay. After 2 weeks, colonies were stained with crystal violet and counted.

**Cellular transformation**

Transformation ability of NIH3T3 cells was assessed by checking colonies formed by confluent cells. A total of 5 x 10^5 cells/well were seeded in 6-well tissue culture plates in complete medium which was replaced every 3-4 days. Fourteen days post-confluence, the cells were fixed and stained with crystal violet (Sigma), and the number of colonies composed of >50 cells was determined. All assays were performed three times. The mean ± SD was calculated from three independent experiments. Cellular transformation was also assessed by observing anchorage-independent growth using the soft agar assay. Briefly, 10^4 cells/well were seeded into 0.35% low melting point agarose (Lonza) on top of a 0.5% agarose layer in 6-well plates. After 3 weeks, colonies were fixed in methanol and stained with 0.1% crystal violet. For quantification, crystal
violet-positive colonies were counted. All assays were performed three times. The mean ± SD was calculated from three independent experiments.

**Deacetylation assays**

Both in vitro and in vivo deacetylation assays were performed as described previously (5).

**Dot blot assay**

Samples (2 µL containing 10 ng – 0.5 µg protein) were spotted onto a nitrocellulose membrane at the center of the grid using a narrow-mouth pipette tip. Membrane was left to dry followed by blocking of non-specific sites after incubation with 5% milk in Tris-Buffered Saline/Tween 20 (TBST) for 0.5-1 h at room temperature. Next, the membrane was incubated with the primary antibody diluted in 1% milk in TBST for 30 min at room temperature. After washing three times with TBST (3 x 5 min), the membrane was incubated for 30 min at room temperature with a secondary antibody conjugated to HRP. Finally, membranes were exposed to x-ray films following incubation with a chemiluminiscence reagent.

**Custom anti-Ac-K147 antibody development**

The anti-Ac-K147 antibody was generated by Eurogentec following the Speedy 28-day protocol. For immunization of rabbits, a peptide carrying acetylated K147 (N-IETSAK(Ac)TRQGVD-C) as well as a control non-acetylated peptide were synthesized and attached to keyhole limpet hemocyanin and (KLH) carrier. After purification, the raised antibody, together with the synthesized peptides used for immunization, was shipped to our laboratory where they were used in additional validation experiments.
**GDP/GTP exchange**

Cells were lysed following the same procedure as described for the Ras activity assay. 2 mg of cell lysate for each condition was used. For 2 mL total lysate, 40 µL 0.5 M EDTA pH 8.0 was added (for a final concentration of 10 mM) followed by vortexing. Then 40 µL 100 mM GDP was added (for a final concentration of 2 mM). Sample was vortexed again and the mixture was incubated at 30°C for 15 minutes with constant agitation. Each 2 mL sample was split in 4 followed by addition of different amounts of GTPγS (10 mM) to the mixture to reach a final concentration of 0, 0.1, 0.5 and 1 mM. After incubating at 30°C for 15 minutes, the exchange reaction was terminated by placing the samples on ice, adding 32 µL of 1 M MgCl₂ (for a final concentration of 60 mM) and vortexing. GTP-bound “active” KRAS was determined by using the Thermo Scientific Active Ras Pull-Down and Detection Kit (Thermo Scientific) according to the manufacturer’s instructions.

**Mass spectrometry**

KRAS<sup>G12V</sup> was purified from 293T cells expressing either Flag-KRAS<sup>G12V</sup> alone or Flag-KRAS<sup>G12V</sup> and SIRT2. For purification, an anti-Flag antibody covalently attached to agarose (ANTI-FLAG<sup>®</sup> M2 affinity gel, Sigma) was used to immunoprecipitate KRAS<sup>G12V</sup>, followed by elution with excess free Flag peptide according to the procedure described in (6). Eluted samples were run on a 4-20% polyacrylamide gel, followed by silver staining (ThermoFisher Scientific). Protein bands were cut and sent to the MSRC Proteomics Laboratory at Vanderbilt University for mass spectrometry analysis as described before (6).
Cloning, expression, and purification of KRAS proteins

KRAS, KRAS\textsuperscript{K147Q}, KRAS\textsuperscript{G12V}, and KRAS\textsuperscript{G12V/K147Q} gBlocks\textsuperscript{®} gene fragments (IDT) were cloned into pMCSG7 vector. Each plasmid was digested with SspI restriction enzymes and the linearized vector was gel-purified. Ten µL of Gibson assembly\textsuperscript{®} master mix (New England Biolabs), 1 pmol pMCSG7 linearized vector, and 3 pmol of each gBlock product were incubated for 1 h at 50 °C. One µL of the Gibson assembly\textsuperscript{®} reaction mix was transformed into TOP10 electro-competent cells (Sigma). All the sequences were validated by DNA sequencing. To improve expression and solubility of the KRAS proteins, the catalytic domain of each KRAS construct (residues 1-169) was amplified from purified pMSCG7, using the following primers: FWD 5’-CTTTAAGGAGGCTCTCCTCCCATGACTGAATATAAACTTGTGGTTGG-3’ and REV 5’-CCGCTAATGCTGCCCTTTTCTTTATGTCTTTTCGAATTTTCTCGAATCTGAAAAATG-3’. All PCR products were gel-purified and inserted into pMCSG58-CPD\textsubscript{Vc} vector. pMCSG58-CPD\textsubscript{Vc} vector comprises the C-terminal domain of the Vibrio cholerae MARTX toxin cysteine protease domain (CPD) followed by a 6x His-tag. The addition of inositol hexakisphosphate (InsP6) triggers the CPD autoprocessing, releasing the target protein in solution without the 6His-tag. pMCSG58-CPD\textsubscript{Vc} was digested with SmaI and gel-purified. The Gibson assembly\textsuperscript{®} cloning method was used to insert each PCR product into pMCSG58-CPD\textsubscript{Vc} linearized vector as described above. Plasmid insert sequences were confirmed to be accurate by DNA sequencing, and each plasmid was transformed into E. coli BL21(DE3)/MAGIC cells. Cultures were grown in Terrific Broth supplemented with 100 µg/mL ampicillin and 30 µg/mL kanamycin at 37 °C until OD\textsubscript{600} = 0.8 and then induced with 1 mM isopropyl-b-D-thiogalactoside at 25 °C for ≈18 h. Bacteria were
harvested by centrifugation, re-suspended in buffer A1 (50 mM Tris pH 8.3, 500 mM NaCl, 10 mM MgCl₂, 0.1% Triton X-100, 5 mM β-mercaptoethanol), and lysed by sonication. After centrifugation at 30,000 x g for 30 min, the soluble lysates were loaded onto a 5-mL Ni-NTA HisTrap column using the ÄKTA protein purification system (GE Healthcare). The column was washed with buffer B1 (10 mM Tris pH 7.5, 500 mM NaCl, 10 mM MgCl₂, 50 mM imidazole) followed by elution in the same buffer with 500 mM imidazole (buffer C1). Proteins were further purified by size-exclusion chromatography (Superdex 200 (26/60), GE Healthcare) in buffer D1 (10 mM Tris-HCl pH 7.5, 500 mM NaCl, 10 mM MgCl₂, 5 mM β-mercaptoethanol). KRAS₁₋₁₆₉-CPD fusion proteins were incubated with 100 µM InsP₆ (Sigma) for 1 h at 25 °C under gentle agitation. Protein samples treated with InsP₆ were loaded again onto a Ni-NTA HisTrap (GE Healthcare), and the flow-through fractions containing KRAS₁₋₁₆₉ proteins were collected. Purified proteins were analyzed by SDS-PAGE.
References

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**A**

Comparison of Kras^G12D-Ptf1 and Sirt2^-/-Kras^G12D-Ptf1.

**B**

Histograms showing normalized intensity with P-values.

**C**

Genetic model for Eila-Cre and Ptf1-Cre.

**D**

Western blot analysis of SIRT2 expression.

**E**

**F**

Graph showing tumor nests per field.

**G**

Graph showing larger tumor size.

**H**

Immunohistochemical images of Kras^G12D and Sirt2^-/-Kras^G12D.

**I**

Immunohistochemical images of H&E, pERK, and Ki67.
Figure S1. Sirt2 loss enhances KRAS-induced tumorigenesis. (A) The pancreata from Kras<sup>G12D</sup>-Ptf1 and Sirt2<sup>−/−</sup>-Kras<sup>G12D</sup>-Ptf1 mice were stained with a Mallory trichrome stain. Blue staining indicates fibrosis. Representative images are shown (10x). Scale bar 200 μM. (B) SIRT2 expression is decreased in human PDAC samples. The Oncomine genomic microarray database (http://www.oncomine.org) was used to determine SIRT2 expression in normal pancreas versus PDAC. P values are shown (C) Generation of mice carrying pancreas-specific deletion of Sirt2. To generate Sirt2 conditional knockout mice, mice carrying a loxP-neo-loxP cassette (upper) were crossed with Ella-Cre transgenic mice to delete the neo gene and generate Sirt2 floxed mice with loxP sites flanking exons 5-8 of the Sirt2 gene (middle). These mice were crossed with Ptf1-Cre mice in order to delete exons 5-8 specifically in the pancreas (lower). (D) PCR-based genotyping of tail DNA using primers b/c and a/c to identify wild type (+/+), heterozygous (fl/+), and homozygous (fl/fl) Sirt2 floxed mice. No Sirt2 deletion can be detected in tail DNA using primers a and c. (E) Western blot using a SIRT2 antibody in pancreata of Sirt2 floxed mice after crossing with Ptf1-Cre mice showing Sirt2 deletion in pancreas. (F, G) The lungs from Kras<sup>G12D</sup> and Sirt2<sup>−/−</sup>-Kras<sup>G12D</sup> mice (n=5-8), two months after intranasal administration of adenoCRE, were harvested, fixed, sectioned, and H&E stained. Lung sections from both genotypes were analyzed for: (F) nests of tumor-like cells per 10x field and (G) size of tumors. Data represent mean ± SEM, *p<0.05. (H) The lungs from Kras<sup>G12D</sup> mice develop mostly atypical adenomatous hyperplasia (AAH) (2 left panels, arrows indicate AAH, 2<sup>nd</sup> panel shows higher magnification, 20x vs 10x) whereas Sirt2<sup>−/−</sup>-Kras<sup>G12D</sup> mice develop mostly adenocarcinoma (2 right panels, 2<sup>nd</sup> panel shows higher magnification, 20x vs 10x). Scale bar 200 μM. (I) The Sirt2<sup>−/−</sup>-Kras<sup>G12D</sup> mice, but not the Kras<sup>G12D</sup> mice, exhibit increased pERK (middle) and Ki67 (right) staining. The lungs from 3 mice in each group (Sirt2<sup>−/−</sup>-Kras<sup>G12D</sup> and Kras<sup>G12D</sup>) were harvested and analyzed by IHC staining with anti-pERK and Ki67 antibodies. Areas of cells with increased staining display a rust color. Representative images are shown (20x). Scale bar 100 μM.
Figure S2. SIRT2 interacts with KRAS and regulates its activity. (A) Endogenous KRAS and SIRT2 proteins were immunoprecipitated from MEF cell lysates with KRAS (left) and SIRT2 (right) antibodies. Interaction was confirmed by western blotting using anti-SIRT2 and anti-KRAS antibodies, respectively. Endogenous levels of both KRAS and SIRT2 are shown as input. (B) Flag-tagged sirtuins (SIRT1-7) were expressed in 293T cells (upper) followed by immunoprecipitation using an anti-Flag antibody. Interacting proteins were eluted after incubation with a Flag peptide and samples were checked for immunoreactivity against KRAS (lower). (C) Endogenous KRAS was immunoprecipitated in Kraslox MEFs and interaction with different members of the sirtuin family was checked using specific antibodies as indicated. (D) Cell extracts from Sirt2+/+ and Sirt2−/− MEFs were analyzed by western blotting using antibodies against pERK, ERK, SIRT2, and KRAS. The same lysates were used to determine the relative amounts of GTP-bound active KRAS through a specific protein interaction with Raf1-RBD. (E) HCT116 cells were infected with lenti-sh luc (sh ctr) or lenti-shSIRT2 (sh SIRT2), treated with EGF (100 ng/mL) for 2.5 and 5 minutes, and extracts were immunoblotted with antibodies against pERK, ERK, and tubulin. (F, G) Totals of 500 and 1,000 HCT116-sh ctr and HCT116-sh SIRT2 cells were plated in soft agar for 21 days (F) and colonies were counted (G). All experiments were done in triplicate, and representative images are shown. (H) 293T cells were co-transfected with either HA-KRAS (upper) or HA-KRASG12D (lower), as well as different HATs. Acetylated levels of both HA-KRAS and HA-KRASG12D were determined after immunoprecipitation using an anti-Ac-K antibody followed by immunoblotting with an anti-HA antibody. Levels of expressed KRAS and KRASG12V are shown as input after immunoblotting with an anti-HA antibody. (I) 293T cells were co-transfected with either HA-KRAS or HA-KRASG12V HATs, and wild-type Flag-SIRT2. Acetylated levels of both KRAS and KRASG12V were determined after immunoprecipitation using an anti-Ac-K antibody followed by western blotting using an anti-HA antibody. Interaction of both KRAS and KRASG12V with SIRT2 was confirmed after immunoprecipitation using an anti-HA antibody followed by western blotting using an anti-Flag antibody, whereas successful immunoprecipitation of both KRAS and KRASG12V was confirmed by western blotting using an anti-HA antibody. Levels of expressed SIRT2 are shown after immunoblotting using an anti-Flag antibody. (J) 293T cells were transfected with either Flag-KRAS (upper) or Flag-KRASG12V (lower) and cells were subsequently treated with nicotinamide and AGK2. The relative amounts of GTP-bound active Ras were determined through a specific protein interaction with Raf1-RBD. Levels of expressed KRAS and KRASG12V are shown after immunoblotting using an anti-Flag antibody. (K) 293T cells were co-transfected with Flag-KRAS, HATs, and either HA-SIRT2wt (wild-type SIRT2) or HA-SIRT2dn (deacetylation null mutant SIRT2). KRAS activity was determined as described in (J). Levels of expressed KRAS and SIRT2 are shown after immunoblotting using anti-Flag and anti-HA antibodies, respectively. Tubulin is used as loading control.
Figure S3. K147 is a SIRT2 deacetylation target. (A) 293T cells were transfected with Flag-KRAS<sup>G12V</sup> in the absence (sample 1) or presence (sample 2) of exogenously expressed SIRT2. After immunoprecipitation using an anti-Flag antibody, samples were separated on a gel, and bands corresponding to Flag-tagged KRAS<sup>G12V</sup> were excised, in-gel trypsin digested, and analyzed by mass spectrometry. Identified peptides are highlighted in yellow, and acetylated lysines are indicated by red circles. K104 and K147 were found to be acetylated in sample 1, whereas K147 acetylation was not detected in sample 2. (B, C) Spectra for both sites found to be acetylated in sample 1 are shown.
Figure S4. K147 acetylation enhances transformation activity of KRAS and plays a role in regulating guanine nucleotide exchange. (A-C) Cell proliferation in NIH3T3 cells expressing KRAS, KRAS<sup>K147R</sup>, KRAS<sup>K147Q</sup>, KRAS<sup>G12V</sup>, KRAS<sup>G12V-K147R</sup> and KRAS<sup>G12V-K147Q</sup> after knocking down endogenous Kras was checked by measuring the number of cells for 6 consecutive days (A) and by determining the number of colonies formed after 21 days (B, C). For cell proliferation, data represent mean ± SD of three independent experiments, **p<0.01 KRAS<sup>G12V/K147Q</sup> vs KRAS<sup>G12V</sup> cells, ****p<0.0001 KRAS<sup>G12V/K147Q</sup> vs KRAS<sup>G12V</sup> cells. For number of colonies formed, data represent mean ± SD of three independent experiments, *p<0.05 KRAS<sup>G12V/K147Q</sup> vs KRAS<sup>G12V</sup> cells. (D) Representative images of mice bearing the subcutaneous tumors after injecting with Kras<sup>lox</sup> MEFs infected with lenti-KRAS<sup>G12V</sup>, lenti-KRAS<sup>G12V-K147R</sup>, and lenti-KRAS<sup>G12V-K147Q</sup> followed by treatment with 4HT. (E) Representative photos of subcutaneous tumors removed after sacrificing the mice injected with the cells described in (D).
Figure S5. K147 acetylation can be detected in vivo by using a KRAS-Ac-K147 antibody. (A) A dot blot assay was performed using different combinations of antigen/antibody: lane 1, SIRT2/SIRT2; lane 2, BSA/non acetylated KRAS (non Ac); lane 3, BSA/acylated KRAS (Ac); lane 4, non Ac-KRAS peptide/non Ac-KRAS; lane 5, non Ac-KRAS peptide/Ac-KRAS; lane 6, Ac-KRAS peptide/Ac-KRAS. (B) Lysates from 293T cells co-transfected with HA-KRAS<sup>G12V</sup> and either GFP (as control) or p300 were immunoblotted with the anti-KRAS-Ac-K147 antibody.

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antibody. Levels of expressed KRAS\textsuperscript{G12V} are shown after immunoblotting using anti-HA antibody. (C) Lysates from 293T cells transfected with either Flag-KRAS or Flag-KRAS\textsuperscript{G12V} in the absence or presence of p300 were immunoblotted with the anti-KRAS-Ac-K147 antibody. (D) Lysates from both Sirt2\textsuperscript{+/+} and Sirt2\textsuperscript{−/−} MEFs were immunoprecipitated using the anti-KRAS-Ac-K147 antibody followed by immunoblotting with a KRAS antibody. Sirt2\textsuperscript{−/−} MEF lysates were also immunoprecipitated using rabbit IgG as a negative control (lane 1). (E) KRAS was immunoprecipitated in Kras\textsuperscript{lox/lox} MEFs followed by western blotting using the anti-Ac-K147 specific antibody. Sirt2 knockdown efficiency (after shSIRT2 lentiviral infection) and Kras knockdown efficiency (upon 4-HT treatment) was confirmed by immunoblotting using SIRT2 and KRAS antibodies. (F) Pancreas tissue sections from Sirt2\textsuperscript{lox/lox}-Kras\textsuperscript{G12D}-Ptf1 (left, 5x, scale bar 200 μM) and Sirt2\textsuperscript{−/−}-Kras\textsuperscript{G12D}-Ptf1 mice were stained by IHC using an anti-KRAS-Ac-K147 antibody. Middle (20x) and right (40x) panels show higher magnification of stained tissues shown in Figure 5G. Scale bar 50 μM. (G) Pancreas tissue sections including areas with normal acinar cells from Kras\textsuperscript{G12D}-Ptf1 (upper) and Sirt2\textsuperscript{−/−}-Kras\textsuperscript{G12D}-Ptf1 (lower) mice were H&E stained (left) or stained by IHC using an anti-KRAS-Ac-K147 (middle) and an anti-pERK (right) antibody. Representative images are shown (5x). Scale bar 200 μM.