

Dissertation

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Chapter One: Introduction

Abstract Lung cancer remains a threat to human health throughout the world. Due to the accumulation of mutations from tobacco or otherwise, lung cancers have a vast mutational spectrum that affects many genes. These characteristics make lung cancer a challenge in understanding how the disease forms. Furthermore, mutational tumor heterogeneity is linked to drug resistance and a poor prognosis for patients. Interestingly, MAPK pathway activation appears to be a prerequisite for lung adenocarcinoma formation. In chapter 1, using genetically engineered mouse models of *BRAF^{V600E}* lung tumorigenesis, we analyzed the contribution of commonly mutated genes to tumor progression. It was found that, while *BRAF^{V600E}* drives benign tumor formation, loss of SETD2 function drives tumor progression to larger tumors. Thus, we identified SETD2 as a common alteration in *BRAF^{V600E}* lung cancer and is a bone fide tumor suppressor.

Genetic sequencing of human lung adenocarcinomas has also revealed the widespread inactivation of p53 as a common alteration in late-stage lung adenocarcinoma. Normal p53 function restricts tumor plasticity, and thus losing p53 function enables tumor cells to adopt more malignant cell identities. However, not all p53 mutations are created equal. In chapter 2, we characterized the effect of three p53 mutant alleles (P53-null, P53-R172H, and p53-R245W) on *BRAF^{V600E}* lung tumorigenesis. It was found that p53-R172H and p53-R245W induce more widespread tumor dedifferentiation, thus leading to worst tumor outcomes compared to p53-loss. Cell lines derived from the tumors allowed for the dissection of signaling pathway dynamics, which led us to observe higher MAPK pathway activation in p53-R172H and p53-R245W cell lines. We also found that p53-R172H and p53-R245W cell lines were more refractory to MAPK pathway inhibition with small molecule inhibitors. These altered phenotypes seen in p53-R172H and p53-R245W are likely a reflection of their altered transcriptomic profiles.

Although genetically-engineered mouse models are foundational in understanding how lung adenocarcinomas develop, progress, and metastasize, little is known about the transcriptional heterogeneity of genetically-engineered mice. In Chapter 3, we profiled the single-cell transcriptomes of *BRAF^{V600E}* lung tumors. We identified distinct clusters of tumor cell types and determined the transcriptional response to MAPK pathway-targeted therapy.

By quantifying the tumor-suppressive landscape and the heterogeneous transcriptional landscape of *BRAF^{V600E}* cells, our work has contributed valuable knowledge to lung adenocarcinoma biology.

NSCLC Lung cancer is a global hazard to human health. The historical consumption of cigarette smoke has caused lung cancer to be the most common and deadly form of cancer. Fortunately, smoking rates have decreased in the past 30 years, with lung cancer rates also declining. However, even if cigarette consumption is eradicated, lung cancer would still significantly affect human health. Indeed, approximately 25% of all lung cancer cases arise in never smokers (Warren and Cummings, 2013). Secondary causes of lung cancer include Radon exposure, high-heat cooking, family history, and air pollution (Malhotra et al., 2016). An individual's risk of lung cancer also increases with age due to decreases in the efficiency of tumor-suppressive mechanisms such as tumor immunity and apoptosis(de Groot and Munden, 2012).

Lung cancer can be broadly classified into two major histological categories based on how the malignant cells look under a microscope: Non-small Cell Lung Cancer (NSCLC) and Small Cell Lung Cancer (SCLC). Within NSCLC, tumors are further characterized based on histological structures, anatomical location, cell-of-origin, and genetic drivers(Herbst et al., 2008). For example, lung adenocarcinoma is the most common subtype and is characterized by glandular and papillary structures, while squamous cell lung carcinoma is characterized by keratin-pearls(Nikitin et al., 2004). Moreover, adenocarcinomas arise from alveolar-type-2 (AT2) cells in the distal lung and alveoli and are typified by genomic alterations throughout the MAPK pathway, such as EGFR, KRAS, BRAF, PI3K

(Ferone et al., 2020; Jackson et al., 2001). On the other hand, squamous cell carcinomas likely arise from basal cells and typically harbor SOX2 gene amplification (Mollaoglu et al., 2018; Mukhopadhyay and Oliver, 2014).

The 5-year survival rate for patients with lung cancer was 25% in 2020 but varied depending on the stage of disease at the time of diagnosis (Siegel et al., 2021). Like other solid cancers, lung cancers are most deadly after primary tumors have metastasized to other organ systems like the liver, kidney, bone, or nervous system (Matsunaga and Shida, 1971). Lung cancer patients with no local or distant metastasis typically have a 5-year survival percentage of approximately 40% compared to approximately 5% with distant metastasis [Woodard et al.; 27740970]. Ultimately, the cause of death of many lung cancer patients includes primary lung tumor burden, infection, pulmonary embolism or hemorrhage, or complications of metastatic dissemination (Nichols et al., 2012). Although patient survival has improved due to improved targeted and immune therapies, the still poor prognosis reflects a need to understand better the molecular mechanisms underlying lung adenocarcinoma initiation, progression, maintenance, and response to targeted- or immune therapy.

The current standard of care for Lung adenocarcinoma patients is often a combination of several conventional chemotherapeutics, including Cisplatin, Carboplatin, Paclitaxel, or Pemetrexed (Ghimessy et al., 2020). Fortunately, patients who have been identified to have EGFR, ALK, ROS, TRK, or BRAF genetic alterations are often offered pathway-targeted therapies (Arteaga and Engelman, 2014; Planchard et al., 2016). As of late, many patients receive immunotherapy first, followed by pathway-targeted therapy after progression on immunotherapy. Furthermore, in the last seven years, exciting therapeutic options for KRAS mutant lung cancer have been developed and tested, such as a combination of MEK-inhibition and autophagy-inhibition and direct inhibitors of KRAS G12c and KRAS G12D (Kinsey et al., 2019; Reita et al., 2022; Zheng et al., 2022)

MAPK signaling Standard activation from the top of the MAPK pathway begins with extracellular ligands (EGF, NGF) binding to Receptor-Tyrosine Kinases (RTKs), leading to loading of RAS with GTP, which activates RAS. Many RAS mutations lead to either more efficient loading of GTP or less efficient GTP hydrolysis, which both serve to activate RAS. GTP-loaded RAS recruits and activates RAF kinases (ARAF, BRAF, CRAF) which then phosphorylates and activates MEK1/2 kinases, which then phosphorylates and activates ERK1/2. Ultimately, this leads to the activation of a suite of cellular processes required for proliferation, differentiation, migration, and cell survival. Therefore, activating mutation in this pathway, from RTKs to kinases, keeps the pathway in a constitutively active state. For example, the most common mutation in *BRAF* is V600E. This valine to glutamic acid substitution is located in the kinase domain and mimics phosphorylation, allowing BRAF(V600E) to become independent of upstream regulations like RAS. This active state drives many of the cancer hallmarks required for tumor formation. It is important to note that oncogenic activation of the MAPK pathway differs from normal ligand-mediated pathway activation. For example, mitogens, such as EGF only transiently activate the pathway even with constant ligand stimulation. Various biological processes integrate this immediate and intense spike in pathway activation in the cell, resulting in specific target genes' transcription. In contrast, oncogenic signaling thru mutational activation of BRAF(V600E) results in constant kinase signaling that is integrated differently than normal pathway stimulation. Thus, an oncogene-specific transcriptional program is activated.

The importance of this pathway is reflected in the strong anti-tumor responses that are seen when small molecule inhibitors are used to target various molecules involved in the MAPK pathway. For example, Dabrafenib and Trametinib, inhibitors of BRAF and MEK kinases, respectfully, are used in patients that harbor BRAF(V600E) melanomas and lung cancers (Planchard et al., 2021; Robert et al., 2014). Although there is strong preclinical and clinical evidence that BRAF+MEK blockade can lead to tumor regression, not all patients respond to treatment (Tabbò et al., 2021). Therefore, multiple pathways need to be targeted to see a complete response, such as PI3K or autophagy (Deuker et al., 2014; Kinsey et al., 2019; Truong et al., 2020).

Great effort has elucidated the proto-oncogenes that drive cancer formation when mutated, such as *MYC*, *RAS*, *PI3K*, and others. Nevertheless, we still do not fully understand the mechanistic details that determine how genes cooperate to drive cancer.

Genetically engineered mouse models Genetically-Engineered mouse models (GEMMs) of human cancer have allowed for the reproducible analysis of cancer biology driven by specific mutations frequently found in cancer patients. These models are helpful in understanding the molecular mechanisms that drive tumor formation and testing tumor responses to targeted or immune therapies. Early mouse models were not engineered but instead selected inbred mouse strains that were natural hosts of tumor transplantation(Strong, 1978). However, these models suffer from long latencies, incomplete penetrance, and are often not genetically similar to human cancers. Therefore, carcinogen-induced models such as UV, Urethane, radiation, cigarette consumption, or MNU treatment models were developed to resemble carcinogen-induced human cancers(Kemp et al., 1994; Westcott et al., 2014). After techniques were developed to genetically manipulate embryonic stem cells came the ability to engineer mice with specific mutations that could drive tumor formation. Since genes involved in cancer are also crucial for normal mouse development, conditional gene manipulation was developed to allow for temporal control.

The most commonly used conditional gene expression system relies on inducible transcription factors, such as the tetracycline-dependent regulatory system, where the treatment of mice with doxycycline toggles gene transcription levels of transgenes. More recently, Cre-lox recombination strategies have been utilized to get more cell-type-specific control. In this system, genes of interest are constructed to contain LoxP-sites that flank a particular genetic element. Upon delivery of Cre-recombinase, the DNA sequence in between LoxP sites is excised. With this approach, one can knock out a gene by engineering LoxP sites around critical exons or induce gene transcription by flanking a strong stop signal such as a polyadenylation sequence upstream of a gene-of-interest sequence such as KRAS(G12D)(Jackson et al., 2001).

Genetically engineered mouse models of human cancer are important preclinical models because they resemble the physiological environment of tumor growth in which tumors arise as progeny from a single initiating cell. These tumors can approximate the genetic alterations, transcriptional landscape, histology, and responsive disposition or lack, therefore, seen in human cancers(cite). However, these models often take months to develop tumors, and compound genetic alterations take time to develop. Later in this thesis, we will utilize TUBA-SEQ to both quantify tumor burden and cooperation of BRAF(V600E) with other common alterations in lung cancer.

Due to the robust connection between lung adenocarcinomas and mutational activation of genes involved in the MAPK-pathway, such as *EGFR*, *KRAS*, and *BRAF*, many labs have generated genetically-engineered mouse models (GEMMs) that harbor conditionally-activated cancer-specific mutant alleles of either EGFR (L858R), KRAS (G12D), or BRAF(V600E).

Our lab has previously developed a conditional mouse model of BRAF (V600E) human lung cancers(Dankort et al., 2007; Shai et al., 2015; van Veen et al., 2019). In the *Braf(CAT)* model, normal BRAF is expressed from a conditional allele prior to Cre-mediated recombination. LoxP sites flank human *BRAF* cDNA encoding normal *BRAF* exons 15-18. Downstream of the LoxP sites is the mutant exon 15 that encodes the murine equivalent of the T1799A mutation that gives rise to the BRAF (V600E) oncoprotein. Downstream of the mutant exon, there is a P2A element and a CAAX-tagged *TdTomato*. Therefore, after Cre-mediated recombination, the BRAF (V600E) oncoprotein and fluorophore *TdTomato* are expressed at normal levels(van Veen et al., 2019).

BRAF (V600E) Tumorigenesis and Cooperating Events for Progression Mutational activation of the MAPK pathway forms lung tumors in mice. Specifically, expression of BRAF (V600E) in the Surfactant-Protein C expression cells (alveolar type 2 pneumocytes) elicits benign tumor formation that fails to

progress to lung adenocarcinoma. It is hypothesized that the cell cycle arrest is dependent on the tumor suppressors P53 and CDKN2A. Loss of either P53 or CDKN2A allows tumors to progress to malignant and deadly lung adenocarcinoma. Also, activating WNT signaling and PI3K signaling also allows BRAF (V600E) adenomas to progress to adenocarcinoma (Green et al., 2015; Juan et al., 2014; van Veen et al., 2019). These observations are in concordance with the "multiple-hit" hypothesis that states that cancers do not arise from single mutagenic events but are the consequence of sequential assaults on the genome that activate more than one oncogenic pathway. These multiple genetic hits allow cells to adopt traits to form cancer, which Bob Weinberg and Douglas Hanahan coined as the "Hallmarks of Cancer" (Hanahan, 2022; Hanahan and Weinberg, 2011). The traits include suppression of apoptosis, active proliferation, altered metabolism, changes in cell identity, and forming blood vessels to supply tumor cells with adequate resources for growth(Hanahan, 2022). BRAF(V600E) expression alone is sufficient to activate many of the hallmarks of cancer, but additional pathways need to be activated or suppressed to develop into malignant tumors. Great effort has elucidated the proto-oncogenes that drive cancer formation when mutated, such as *MYC*, *RAS*, *PI3K*, and others. Yet we still do not fully understand the mechanistic details that determine how genes cooperate to override tumor suppressive mechanisms and drive cancer progression.

Talk here about the prevalence of mutations in the egfr-ras-RAF-mek-erk pathway

Identification of this pathway as central to luad tumorigenesis

Development of gemms to study BRAF^{V600E}

Proliferation arrest by this model

How does p53 loss lead to sustained wnt-signalling? Do dominant-negative recapitulate?

P53- mediated tumor suppression Cancers pose an evolutionary challenge for organisms regardless of their life span. Therefore, suppression of the outgrowth of neoplastic cells is built into many organisms. One such mechanism is the engagement of the p53 pathway. At the time of writing this dissertation, there are approximately 105,000 manuscripts on PUBMED that mention P53. P53 is well studied because of its prominent role in tumor suppression and because it is the most frequently altered gene in human cancer.

The p53 protein was initially discovered as a protein that bound large T antigen of the SV40 virus in infected cells (Linzer and Levine, 1979).P53 is a transcriptional factor with a DNA-binding domain that has a DNA-binding domain recognizes explicitly two decameric half-sites. P53 also contains two N-terminal transactivation domains and a C-terminal oligomerization domain critical for P53-target gene activation(Laptenco and Prives, 2006). Central to the ability of P53 to induce transcriptional activation is tetramerization. Thus, if one or more P53 protein in the complex is compromised in transactivation or DNA binding, transcriptional activity is compromised. Therefore, if one copy of P53 is mutated, the organism's ability to suppress tumor formation is compromised.

The importance of p53 is best demonstrated in people and in mice that have just one mutated copy of P53. Humans with the familial- inherited Li-Fraumeni syndrome, in which they have one or more mutations in p53, are almost certain to develop cancer early on and throughout their life(Li et al., 1988). Similarly, mice lacking two function P53 genes are prone to leukemias and lymphomas and have a shortened lifespan due to cancer incidence(Donehower et al., 1992). On the other hand, organisms such as the African and Asian elephants, which have an abnormally low level of cancer incidence, have been shown to have higher multiple copies of *TP53* in their genomes(Abegglen et al., 2015).

The most well-known cellular functions of P53 include its ability to induce cell-cycle arrest in response to DNA damage(el-Deiry et al., 1993). Many stress signals, including oncogene activation, have been shown to stimulate a reversible or irreversible cell cycle arrest(McNeal et al., 2021; Zhu et al., 1998). However, the strength of P53 induced cell cycle arrest or apoptosis is likely cell type- and cellular stress-specific and is an active area of p53 research.

Once a cell encounters a stressor such as DNA damage or oncogene activation, P53 is stabilized and accumulates in the nucleus to activate a suite of target genes that can lead to several cellular phenotypes, such as cell-cycle arrest or apoptosis.

P53 was initially thought to be an oncogene as P53 is accumulated in lung cancer, which is not common in normal tissues(Bartek et al., 1990; Iggo et al., 1990). Moreover, ectopic expression of a P53 cDNA was found to aid in transforming primary cells induced by RAS. However, early studies erroneously used mutated P53 instead of wild-type, leading to the misclassification of P53 as an oncogene. We now know P53 suppresses cell growth and transformation(Levine and Oren, 2009).

Oftentimes, P53 is mutated in the DNA-binding domain at sites commonly referred to as "hotspots" due to their extraordinary frequency. These hotspot mutations poison the ability of p53 to bind DNA, therefore, blocking its function. Interestingly, since one mutated P53 can disrupt the entire P53 tetramer complex, a single mutation can exert dominant-negative effects by inhibiting the normal tumor-suppressive functions of P53. Although one mutant P53 allele is enough to compromise transcriptional activity, there is still selective pressure to lose the other wild-type copy. Loss of hemizygosity implies there is still residual tumor suppression in the presence of a wild-type P53 allele(Muzumdar et al., 2016).

Early on in P53 studies, there were reports of specific P53 mutations having a "gain-of-function" (GOF) effect. For example, Li-Fraumeni patients with specific missense mutations in P53 would develop tumors earlier than Li-Fraumeni patients with loss-of-function (LOF) mutations (Bougeard et al., 2008). Furthermore, experimental cell biology studies would express mutant-p53 in P53-null cells and demonstrate enhanced tumorigenic potential (Levine, 2020). Further evidence for GOF mutant-P53 in mice indicated that missense mutant P53 induced different cancer types and enhanced metastasis than LOF P53 (Olive et al., 2004). Furthermore, many groups have shown that mutant-p53 can alter signal transduction affecting chemoresistance and metabolism.

The cellular mechanisms that GOF P53 is involved in are also well-characterized but context-dependent. For example, in Pancreatic cancers driven by KRAS(G12D), Mutant P3 interacts with CREB to induce FOXA1 transcription, which enhances Beta-Catenin signaling to augment liver metastasis (Kim et al., 2021).

Chapter Two: Quantitative tumor burden and small library screen using TUBA-SEQ

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##Abstract Large-scale sequencing of cancers has revealed that lung adenocarcinomas have a high mutation rate. Although it is clear that the evolution of lung adenocarcinoma is centered around mutational activation of the EGFR-RAS-RAF-MEK-ERK pathway, little is known about secondary mutations that drive disease progression. Since lung cancers are riddled with mutations, it is challenging to glean which mutations support cancer growth and which are merely bystanders. Therefore, pre-clinical mouse models that are syngenic are essential to determining the oncogenicity of a particular gene. Here, we use Tuba-Sequencing combined with CRISPR/Cas9 gene editing to simultaneously determine the contribution of the loss of one of eleven tumor suppressor

genes in BRAF^{V600E} mouse lung tumorigenesis. We show that loss of either *Apc* or *Setd2* leads to enhanced tumor growth. We validated SETD2 loss as a cooperating mutation with single-guide RNAs. Functionality, SETD2 is a histone lysine methyltransferase that marks active gene bodies by depositing me³ marks on H3K36. In BRAF^{V600E} lung tumors, *Setd2* loss decreases H3K36me³ and H3K36me³ was correlated with larger tumors. SETD2 RNA is expressed less in human lung cancers than in normal lungs and can indicate worse overall survival. Moreover, SETD2 loss is common in human BRAF^{V600E} lung adenocarcinomas. Together, our results highlight the potential importance of SETD2 in the progression of BRAF^{V600E} lung adenocarcinoma.

Introduction

Next-generation sequencing has allowed for the unbiased identification of mutated genes in human cancer (2014; Hoadley et al., 2014; Kandoth et al., 2013). The wealth of knowledge gained from such large-scale sequencing projects has revealed actionable mutations that can be targeted with small molecule inhibitors, such as dabrafenib and trametinib. However, identifying possible targets is not enough, as many cancers with similarly mutated oncogenes, such as BRAF^{V600E}, respond differentially to BRAF^{V600E}-targeted therapy. For example, melanoma patients with BRAF^{V600E} mutations respond well to BRAF^{V600E}-targeted therapy, whereas colorectal cancer patients with BRAF^{V600E} mutations do not respond(Corcoran et al., 2012). The disconnect between putative driver oncogenes identified in human cancers and response to targeted therapy is a problem that can be solved with rigorous pre-clinical experimentation, such as in GEMMs.

Genomic sequencing of human tumors has also been a crucial insight into how cancers form and what signaling nodes might be necessary for progressive stages of cancer evolution. For example, pancreatic cancers have a stereotypical histological progression brought about by known mutational activation. Specifically, mutations in KRAS initiate tumorigenesis, and loss of TP53 function and SMAD3 are associated with cancer progression(Orth et al., 2019). However, this is more difficult to do in lung adenocarcinoma because of the high mutational burden often due to the consumption of cigarettes(Alexandrov et al., 2016; Yoshida et al., 2020).

Indeed, in highly mutated cancers such as melanoma and lung cancer, recurrent mutations are not necessarily predictive of biological importance. To better understand the importance of candidate tumor suppressors or oncogenes, researchers exploit tumor cell lines and GEMMs to identify bona-fide genetic drivers of cancer. However, these systems are suboptimal, as they are either not physiologically relevant (tissue culture) or are not scalable to determine the effect of many mutations (GEMMs). Moreover, techniques to measure the cooperation of genes to drive lung cancer are qualitative and are not sufficiently quantitative to allow for large-scale studies. Typically, lung adenocarcinoma is quantified by lung weight or histological analyses such as tumor burden (Percentage of the lung that is tumor), tumor size, and immunohistochemistry (IHC) of tumor markers. Although essential and informative, these analyses are not high-throughput and rely on many mice to achieve statistical confidence. Furthermore, there is intrinsic bias when calculating tumor size and tumor burden, as tissue-sectioning results in tumor sections that do not accurately portray actual tumor size. Therefore, more advanced techniques are required for large-scale analysis of mouse tumors that are both quantitative and high throughput.

Recently, Monte Winslow and his lab at Stanford University have developed **Tumor- Barcode Sequencing** (Tuba-Seq), a rapid, multiplexed, quantitative assay for analyzing the contribution of tumor-suppressor loss *in vivo*. This technique delivers a uniquely barcoded lentiviral encoding Cre recombinase to activate Cre-mediated oncogenes and a guide RNA to inactivate a single tumor suppressor gene. Therefore, each tumor will be driven by the same oncogene but with different tumor suppressors inactivated. After some time (6-24 weeks), tissue is harvested for whole-lung DNA extraction, and targeted DNA-sequencing is performed to quantify relative tumor size to spike-in

controls(Rogers et al., 2017). Therefore, the contribution of a single tumor suppressor can be quantified in a single animal.

$\text{BRAF}^{\text{V600E}}$ has previously been shown to be a potent driver of lung tumorigenesis(Dankort et al., 2007). However, tumors fail to progress past adenoma-like tumors. Given the theory that $\text{BRAF}^{-\text{V600E}}$ induces growth arrest after an initial burst of neoplastic growth, it would be of interest to know if all tumors grow to a predetermined and absolute size or if there is a spectrum of tumor sizes that are determined by extrinsic factors such as proximity to blood vessels or specific stromal cells. Therefore, we used Tuba-Seq combined with CRISPR/CAS9 genome editing to explore the contribution of 11 tumor suppressor genes and five controls in our $\text{BRAF}^{\text{V600E}-}$ - driven lung tumor models.

Results

Adenoviral vectors and lentiviral vectors introduced in the lungs of GEMMs offer temporal control and can be titrated to achieve an optimal level of tumor numerosity. However, only lentiviral vectors can be used to barcode tumors because they integrate into the genome while adenoviruses do not(DuPage et al., 2009; Rogers et al., 2017). With CRISPR/CAS9 gene editing, lentiviruses have been used to analyze many genetic interactions of oncogenic *Kras* driven lung adenocarcinoma(Rogers et al., 2017; Winters et al., 2017). To enable simultaneous activation of $\text{BRAF}^{\text{V600E}}$ and loss of tumor suppressor genes, we bred *Braf*^{CAT}; *H11b*^{LSL-CAS9/+} mice. To determine the size of $\text{BRAF}^{\text{V600E}}$ tumors and the contribution of 11 tumor suppressor genes (*Trp53*, *Rb1*, *Cdkn2a*, *Apc*, *Rbm10*, *Stk11*, *Setd2*, *Arid1a*, *Atm*, *Keap1*, *Smad4*) on $\text{BRAF}^{\text{V600E}}$ lung tumorigenesis, we obtained tittered barcoded-lentiviral plasmids that encode Cre-recombinase as well as a sgRNA that inactivates one of eleven tumor suppressors common in lung adenocarcinoma. All genes tested are putative tumor suppressor genes commonly lost in human lung cancer patients. Each lentiviral solution contained ~1000 individually barcoded lentiviruses. We infected both *LSL-Cas9* and *CAS9* deficient mice in order to determine the contribution of genetic manipulation on tumor size. Mice meant to be analyzed after eight weeks received a viral titer of 1×10^5 , while mice meant to be analyzed after 16 weeks received a viral titer of 1×10^4 . After euthanasia, lungs were collected for genomic DNA extraction and barcode amplification, followed by the preparation of sequencing libraries. Before genomic extraction and library preparation, a known quantity of known-barcoded cells was added to benchmark sequencing reads to a known cell number(figure).

We determined the mean size of a $\text{BRAF}^{\text{V600E}}$ driven tumor to be 21,248 thousand cells (figure). Interestingly, there is a widespread distribution of tumor sizes, with the most giant $\text{BRAF}^{\text{V600E}}$ tumor to be ~300,000 cells, along with many small tumors (figure).

Gross, histologic images of experimental lungs show an increase in tumor burden with both time (Figure xxx) and CAS9 status (figure xxx). Therefore, based on macroscopic inspection, at least one tumor suppressor appeared to cooperate with $\text{BRAF}^{\text{V600E}}$ to increase tumor burden. In order to determine the fitness landscape of any one particular tumor suppressor, we calculated the relative number of tumor cells in a given tumor using the Tuba-Seq genomics pipeline. After eight weeks post tumor initiation, there is no statistically significant contribution of tumor suppressor loss (Figure). SETD2 loss was trending towards statistical significance. However, 16 weeks post-initiation revealed that both SETD2 loss and APC loss cooperated with $\text{BRAF}^{\text{V600E}}$ to form larger tumors (Figure x). APC loss was a suspected hit as we have previously shown that activation of Wnt-beta-catenin signaling promoted $\text{BRAF}^{\text{V600E}}$ tumorigenesis. As APC loss is predicted to increase Beta-catenin signaling activity, this result was unsurprising to us. However, loss of SETD2 was a novel finding in the context of $\text{BRAF}^{\text{V600E}}$ lung tumorigenesis.

Given our findings of SETD2 loss increased tumor burden concurrent with $\text{BRAF}^{\text{V600E}}$, we next validated SETD2 in a new cohort of mice with single-guide RNAs targeting SETD2. Indeed, loss of

SETD2 increased tumor burden from 5% to over 30% in some animals. Since SETD2 antibodies are unreliable in both western blots and immunohistochemistry, we used H3K36me3 as a surrogate marker for SETD2 activity by immunohistochemistry. Whereas 90% of nuclei in BRAF^{V600E} only tumors were H3K36me3, more than 60% of tumors of mice infected with SETD2 CRISPR virus were negative for H3K36me3. Furthermore, when classified by H3K36me3-positive or -negative, there is even more substantial evidence that SETD2 loss leads to larger tumor formation.

Discussion

The mutational landscape of lung adenocarcinoma is complex due to the lifelong bombardment of DNA by carcinogens ingested into the distal lung epithelium[23770567]. While understanding which altered genes play causal roles in tumor formation and progression is required for personalized medicine, the noisy mutation profiles in lung adenocarcinoma make it hard to do so. However, there is a central role for the MAPK pathway during the initiation and progression of lung adenocarcinoma (Cicchini et al., 2017). Oncogenic BRAF^{V600E} is a significant driver of lung adenocarcinoma. However, mouse models with activation of BRAF^{V600E} only lead to benign tumorigenesis(Dankort et al., 2007; Shai et al., 2015). Although it is known that activation of WNT and PI3k signaling and loss of *Trp53* and *Cdkn2a* genes allow BRAF^{V600E} benign lung tumors to progress to malignant disease, it is a time-consuming and laborious process to effects individual tumor suppressors, in GEMMs. Therefore, we sought to systematically test a variety of known tumor suppressors in lung adenocarcinoma. Furthermore, until now, we did not know the absolute number of cells in a BRAF^{V600E} adenoma, only its area, which is confounded by tissue sectioning and stromal cell contamination. Prior to Tuba-Seq, it was challenging to accurately determine the size of BRAF^{V600E} adenomas. Based on population doublings and cross-sections of BRAF^{V600E} tumors in histological sections, it is estimated that BRAF^{V600E}-driven lung adenomas grow about 15-20 populations doublings and reach a size of 1mm³(Juan et al., 2014). With Tuba-Seq, we accurately discerned the absolute size of all BRAF^{V600E} lung adenomas, regardless of size. Interestingly, our results have revealed that BRAF^{V600E}-driven tumors widely differ in absolute tumor cell number. Our results support the hypothesis that each tumor-initiating cell has unique barriers to tumor outgrowth. This heterogeneous response to BRAF^{V600E} activation likely reflects intrinsic differences between individual cell states. Indeed, tumorigenic potential can vary significantly due to different cell types (AT2 vs. Club cell vs. AT1) or be influenced by subtle differences within a cell type, such as proximity to supporting stromal cells such as fibroblast or immune cells. Since we used a ubiquitous promoter to drive BRAF^{V600E} and CAS9 expression, not all sequence likely reads come from adenomas or tumors but from average cell turnover during tissue homeostasis. Therefore, in our quantification and analysis, we focused on tumors greater than 5,000 cells.

We combined Tuba-Seq with CRISPR/CAS9 gene editing of eleven putative tumor suppressor genes with four inert control sgRNAs. We identified APC and SETD2 loss as a cooperating genetic event that allows BRAF^{V600E} tumors to grow larger. Therefore, APC and SETD2 are bona fide tumor suppressors in lung adenocarcinoma. APC loss is known to activate the Wnt-signaling pathway and is a well-established tumor suppressor in colon cancer(Powell et al., 1992). Furthermore, our lab has previously published a requirement for WNT-signaling for the progression of BRAF^{V600E} lung tumors(Juan et al., 2014). Specifically, inhibition of WNT signaling by a Porcupine inhibitor prevents tumor onset, while constitutive activation of the pathway drives tumor progression to high grade. Therefore, it is not surprising that APC loss cooperates with BRAF^{V600E} to induce tumor growth.

We have also identified SETD2 loss as a potent tumor suppressor(Figure 2). SETD2 is mutated in other solid malignancies on cBioPortal, such as Kidney and bladder cancers(Cerami et al., 2012). SETD2 is a histone methyltransferase that deposits trimethylation marks on H3K36(Bhattacharya and Workman, 2020). Chromatin modifiers use H3K36me³ to mark actively transcribed genes(Edmunds et al., 2007). Mining of TCGA mRNA expression and survival analysis further revealed that SETD2 mRNA is lower in

lung adenocarcinoma and is associated with poorer overall survival(Figure 2). Interestingly, *SETD2* alterations are also significantly associated with *BRAF^{V600E}* mutations (Figure 2). The unique association of *BRAF^{V600E}* mutation and *SETD2* loss may reflect an interesting synergistic effect between MAPK signaling and loss of chromatin regulation. One hypothesis is that *SETD2* loss drives spurious transcriptional activation.

Similarly, recent work by Monte Winslow and David Felder's labs have identified *SETD2* loss as a cooperating event in KRAS-G12D lung tumorigenesis(Walter et al., 2017). However, the mechanism by which *SETD2* loss drives tumor progression in either *BRAF^{V600E}* or KRAS-G12D remains elusive.

Although we have identified novel tumor suppressors in *BRAF^{V600E}* lung tumorigenesis using Tuba-Seq, we have failed to confirm previously identified tumor suppressors *CdkN2a* and *Trp53* as cooperating events. One hypothesis is that the sgRNAs targeting *Trp53* and *Cdkn2a* are inefficient. However, previous Tuba-Seq experiments did confirm *Trp53* editing and even demonstrated tumorigenic cooperation with KRAS-G12D(Rogers et al., 2018). Another hypothesis is that the effect size of *trp53* loss is so tiny that many mice and tumors are needed to detect robust cooperation. Indeed, p53 loss is not that to suddenly make tumors immediately proliferate. Instead, p53 loss merely allows for tumor cells to sample various transcriptional programs that can be co-opted for increased fitness in the face of stress. Along similar lines, perhaps 16-weeks is too little time to allow for *Trp53* or *Cdkn2a* deficient tumors to proliferate. Likely, it is a combination of small effect size, inefficient gene editing, and insufficient time that prevented us from identifying *Trp53* as a tumor suppressor in *BRAF6V600E*^ driven lung tumorigenesis using Tuba-Seq combined with CRISPR/CAS9.

Since GEMMs lack the mutational burden exhibited by human lung adenocarcinomas, they are good models to study gene-gene interactions in tumorigenesis. Our data provide further insight into the essential biological mechanisms for *BRAF^{V600E}* lung tumorigenesis. Additional studies of *SETD2*'s role in lung adenocarcinoma will further reveal how it alters the evolution of lung adenocarcinomas. Furthermore, given its striking association with *BRAF^{V600E}* mutations, it will be essential to know how *SETD2* alters the response to *BRAF^{V600E}*-targeted therapy. A complete understanding of the consequences and outcomes of genetic cooperation between *BRAF^{V600E}* and tumor suppressors or oncogenes will help clinicians better understand tumor evolution and paths to drug resistance.

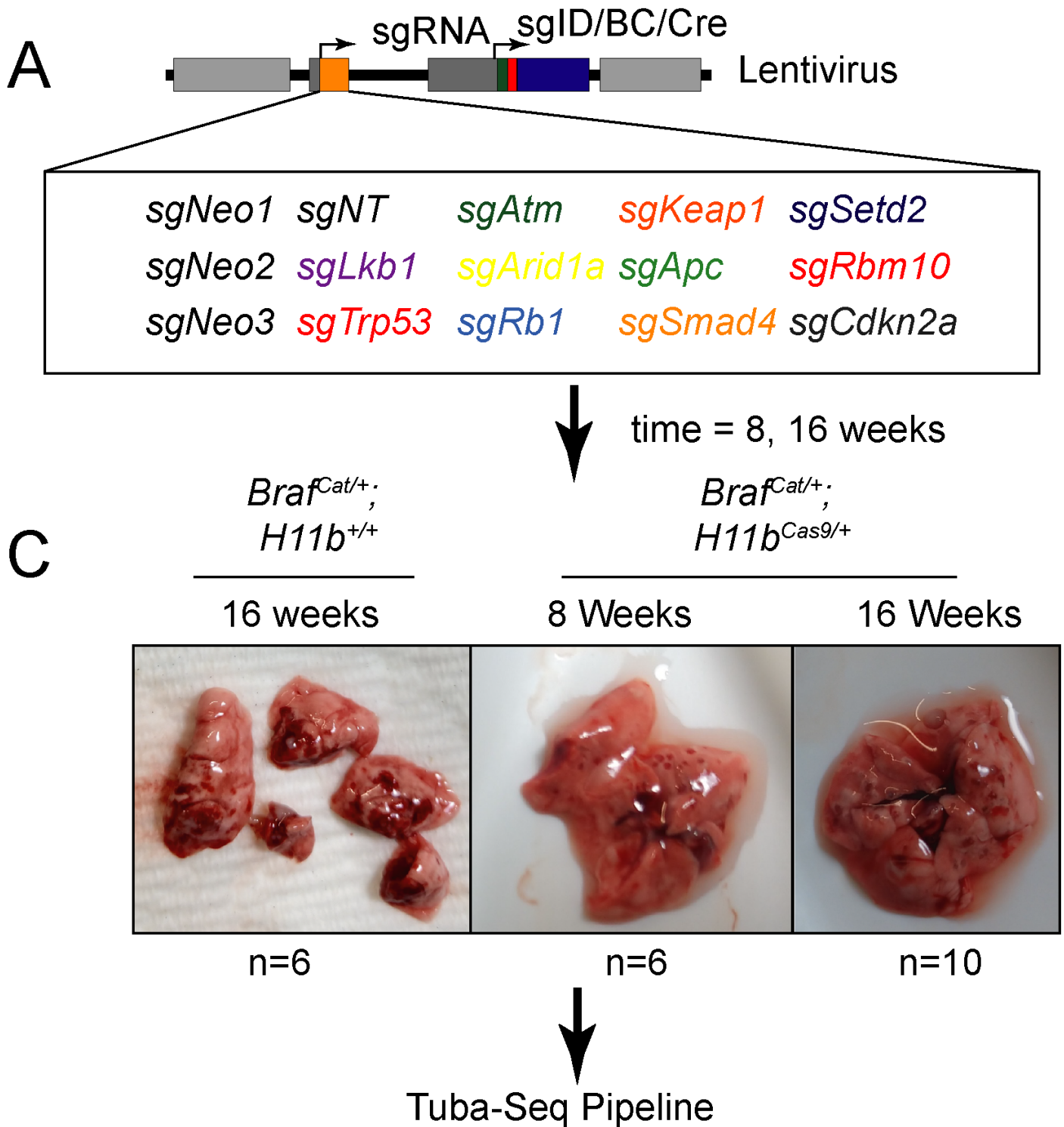
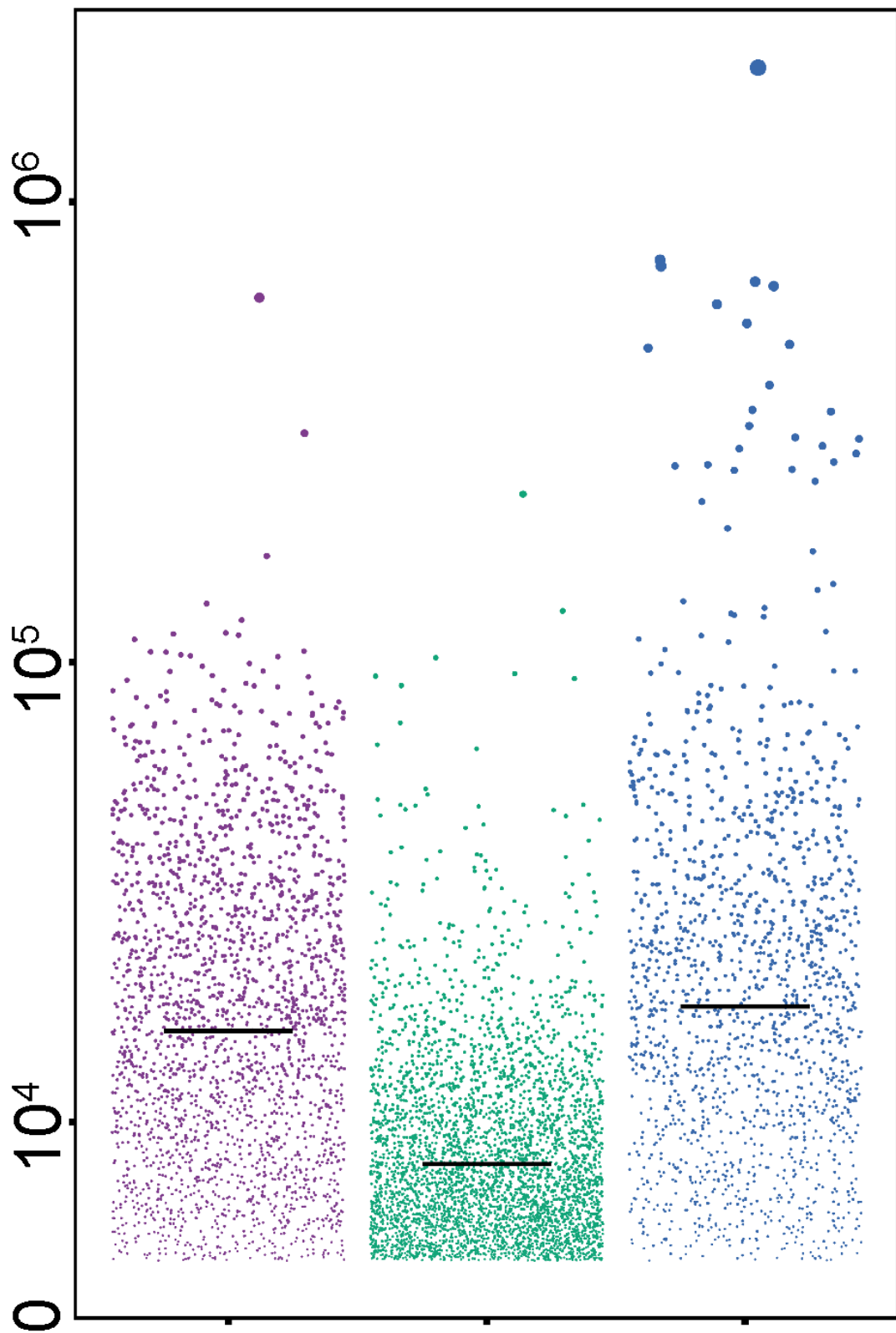


Figure 1: Figure 2.1. Overview of Tuba-Seq experimentation (A) Lenti-sgRNA-Cre plasmid library with collection of targeted tumor suppressors. (B) *Braf*^{CAT/+} and *Braf*^{CAT/+}; *H11b*^{LSL-Cas9/+} were infected with either 10^4 (16 week *Braf*^{CAT/+}; *H11b*^{LSL-Cas9/+}) or 10^5 (8 weeks *Braf*^{CAT/+}; *H11b*^{LSL-Cas9/+} and 16 weeks *Braf*^{CAT/+}; *H11b*^{LSL-Cas9/+}) (B) Representative gross images of experimental mouse lungs.

Number of tumor cells



16 Weeks

n= 2,211

mean= 21,248

$Braf^{Cat/+};$

8 Weeks

n= 4,106

mean= 9,095

$Braf^{Cat/+};$

16 Weeks

n= 1,514

mean= 29,867

$Braf^{+/+};$

$H11b^{+/+}$ $H11b^{\text{Cas9}/+}$

Figure 2: Figure 2.2. Jitter plot of calculated tumor cell number based on Tuba-Seq pipeline. Tumor size corresponds to dot size. Mean tumor number for 16 weeks $Braf^{\text{CAT}/+}; H11b^{+/+}$ is 21,248 cells. Mean tumor number for 8 weeks $Braf^{\text{CAT}/+}; H11b^{\text{LSL-Cas9}/+}$ is 9,095 cells. Mean tumor number for 16 week $Braf^{\text{CAT}/+}; H11b^{\text{LSL-Cas9}/+}$ is 29,867 cells.

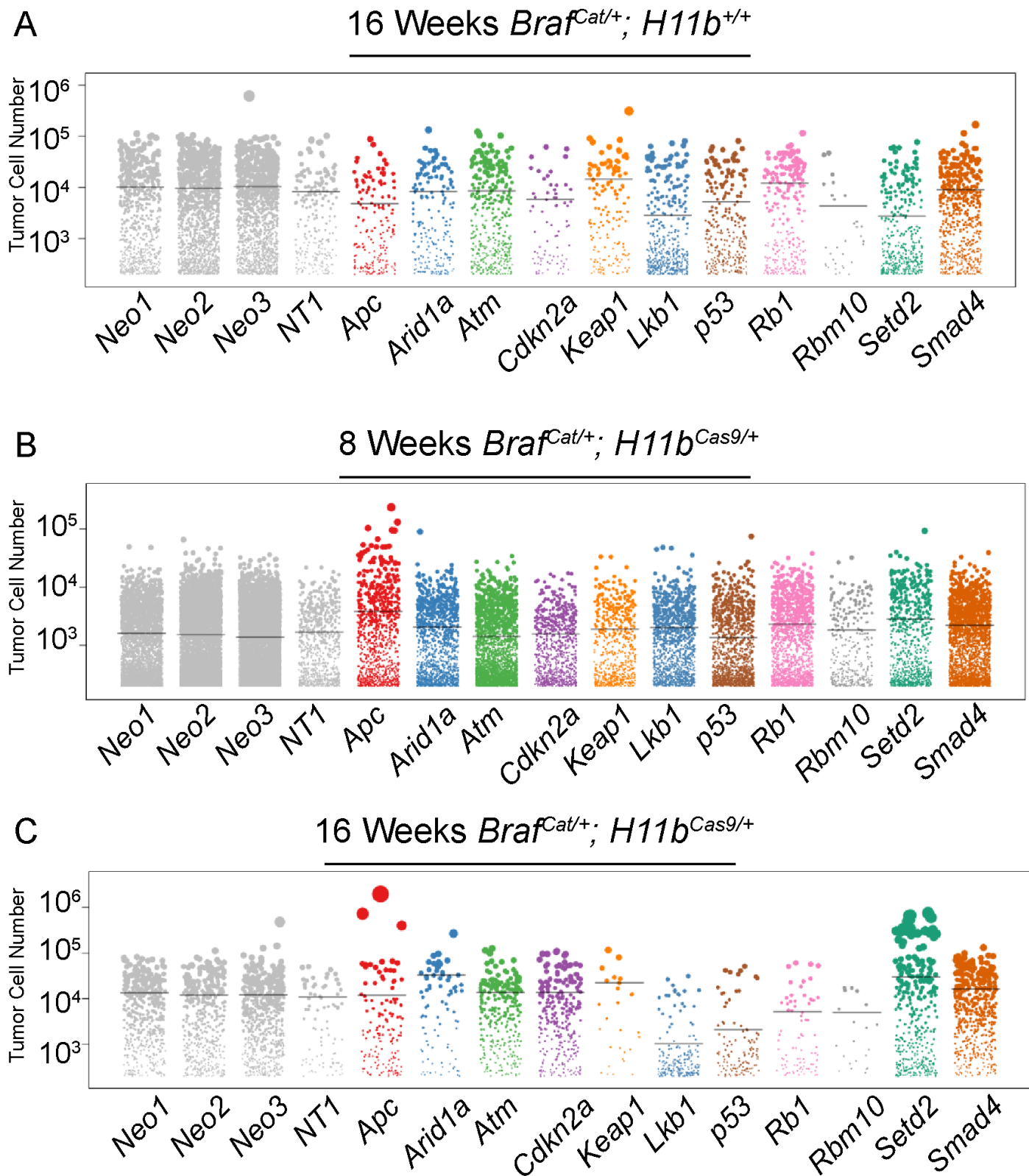


Figure 3: Figure 2.3. Jitter plot of calculated tumor cell number based on Tuba-Seq pipeline seperated by sgID. (A) 16 weeks $Braf^{\text{CAT}/+}; H11b^{+/+}$ (negative control) has no apparent cooperative genetic events. (B) 8 weeks $Braf^{\text{CAT}/+};$

H11b^{LSL-Cas9/+} identified Apc loss as a cooperating genetic event based on increase in tumor cell number (C) 16 week *Braf*^{CAT/+}; *H11b*^{LSL-Cas9/+} identified Apc and Setd2 loss as cooperating events.

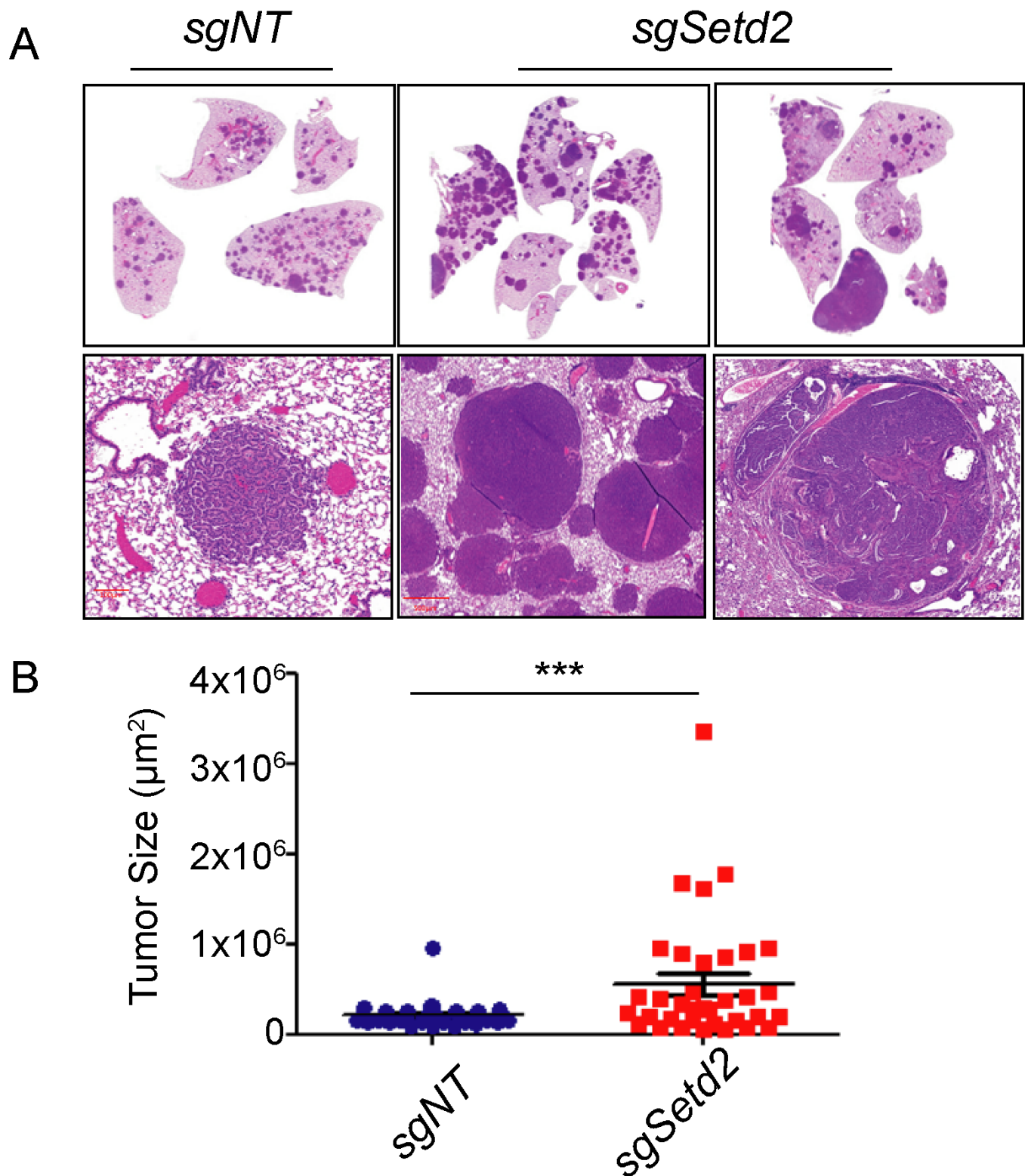


Figure 4: Figure 2.4. Validation of *Setd2* loss as cooperating genetic event in *BRAF*^{V600E} lung cancer. (A) Representative images of lung tumor histology of 16 weeks *Braf*^{CAT/+}; *H11b*^{LSL-Cas9/+} mice infected with 2.5×10^4 PFU Lenti-sgSetd2-Cre or Lenti-sgNT1-Cre. (B) Quantification of tumor size based on histology, n= 30 tumors. * = p-value<.05.

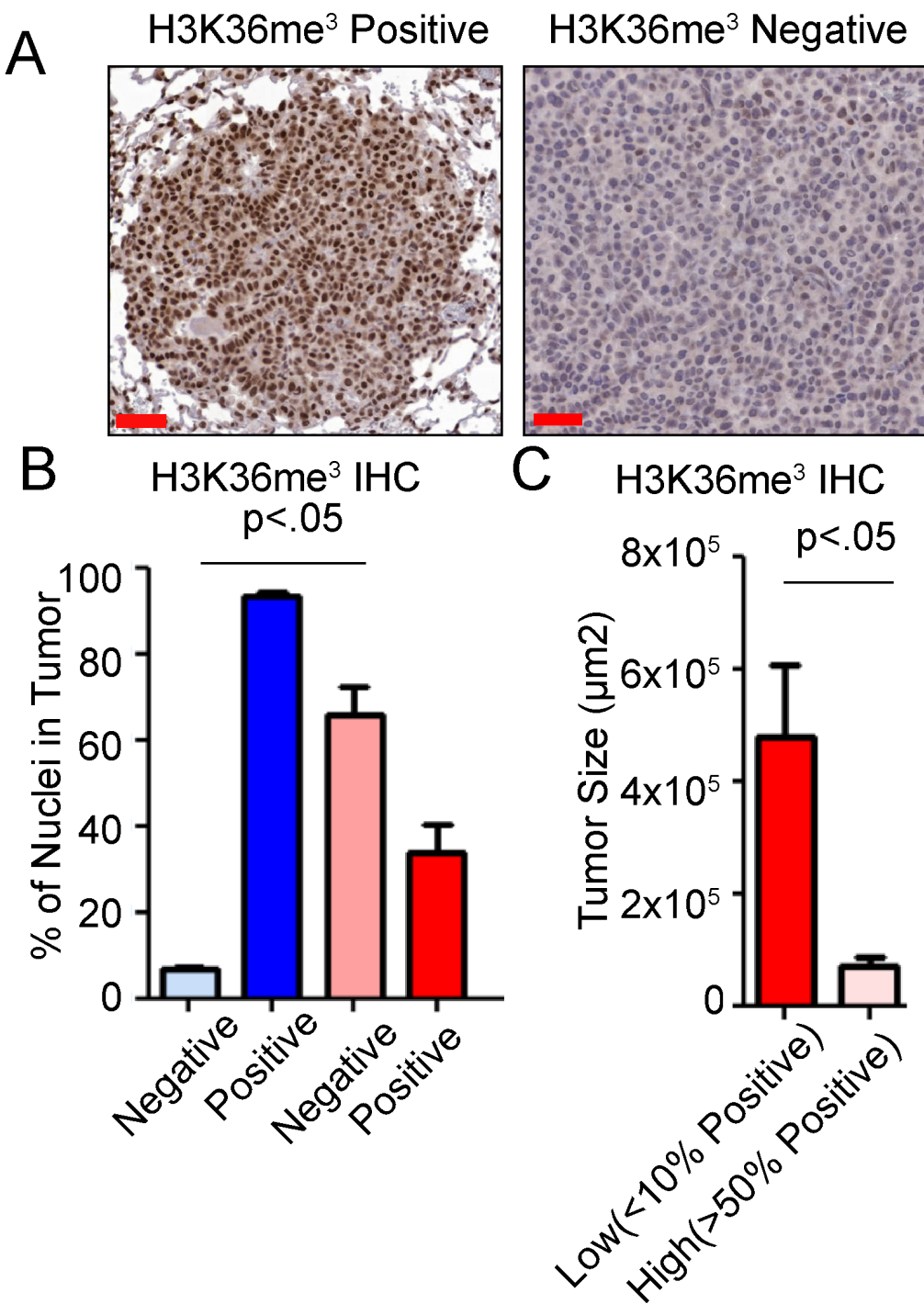


Figure 5: Figure 2.5. H3K36me³ is lost in larger tumors in *Set2d*-edited lung tumors.(A) Representative images of H3K36me3 (surrogate for *setd2* loss) depicting either “high” or “low” staining indicative of protein expression. (B) Quantification of H3K36me³ positive tumors in either *sgSetd2* or *sgNT1* targeted mice. = p -value $<.05$ for negative tumors. (C) Quantification of tumor size paired with H3K36me³ negative tumors. Tumor size and H3K36me³ positivity were simultaneously measured using QuantCenter and binned in either “low” or “High” and then average tumor size was determined for each group. = p -value $<.05$.

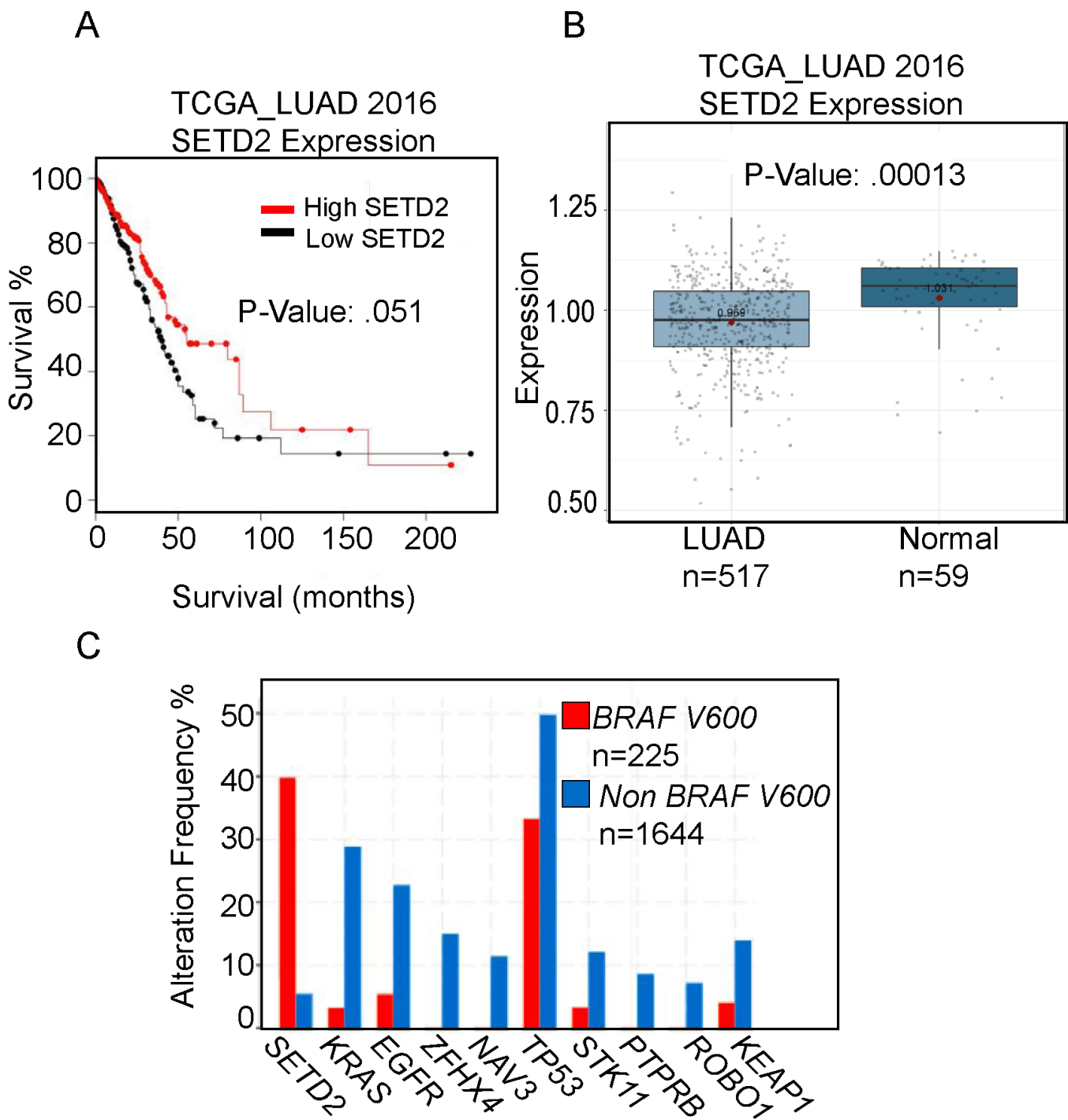


Figure 6: Figure 2.6. *SETD2* loss is common in human cancers, an indicator of poor prognosis and frequently occurs with *BRAF^{V600E}* mutations.(A) Survival analysis with "Survival" R package indicates a trend towards poorer overall survival with low *SETD2* mRNA in TCGA Lung adenocarcinoma dataset. (B) *SETD2* mRNA is attenuated in Lung adenocarcinoma compared to normal lung tissue (C) *SETD2* mutations significantly co-occur with *BRAF^{V600E}* mutations and are present in 40% of *BRAF^{V600E}* mutant Lung Adenocarcinoma.

Chapter Three

Abstract

TP53 missense BRAF^{V600E} mutations both frequently occur in lung adenocarcinoma. While genetically engineered mouse models (GEMMs) of lung adenocarcinoma are invaluable for understanding the initiation, progression, and response to therapeutics, little is known about how different hotspot point mutations in p53 affect disease progression and therapeutic responses. To this end, we use GEMMs, tumor-derived cell lines, and human lung cancer cell lines to test how murine homologs of p53 hotspot mutants R172H and R245W compare to p53-deletion. We find that conformation mutation p53^{R172H} drives more aggressive lung adenocarcinomas and dedifferentiation. We also observe p53^{R172H} and p53^{R245W} accumulation in only high-grade tumors that have activated ARF expression. In vitro and in vivo studies demonstrate enhanced resistance to BRAF+MEK blockade with p53^{R172H} and p53^{R245W}, a standard treatment regimen for BRAF^{V600E} mutant patients. Mechanistically, mutant-p53 alters the transcriptome of cancer cells and enhances entry into the cell cycle by a yet to be determine mechanism.

Introduction

Lung adenocarcinoma (LUAD) has remained the deadliest form of cancer since 1987 (Siegel, Miller et al. 2021). It is characterized by alterations in the MAP kinase (MAPK) pathway, including activating mutations in the oncoproteins EGFR, KRAS (G12C, G12D, G12V) or BRAF^{V600E} (2014; Dankort et al., 2007; Jackson et al., 2001; Politi et al., 2006). The recognized predominant cell of origin for LUAD, lung alveolar type-2 pneumocyte cells (AT2), are exquisitely responsive to the transformative properties of constitutive activation of the MAPK pathway (Sainz de Aja and Kim, 2020; Sutherland et al., 2014). Indeed, genetically engineered mouse models (GEMMs) of mutationally activated EGFR, KRAS, or BRAF form lung tumors that strongly resemble key characteristics of human LUAD. However, tumors formed in all three models (KRAS:G12D, EGFR: Deletion 19 or L858R, or BRAF^{V600E}) develop only histologically low- to mid-grade cancer and are growth-arrested (Dankort et al., 2007; Jackson et al., 2001; Politi et al., 2006). Prior research has determined that oncogene-induced tumor growth arrest occurs through activation of several tumor suppression mechanisms, most notably, p53-mediated tumor suppression (Shai et al., 2015).

Alterations in the *TP53* tumor suppressor gene are common in all cancer types, including LUAD (Bouaoun et al., 2016). P53 encodes a transcription factor that responds to various cell stresses, such as DNA damage and oncogene activation. In response to stress, p53 is stabilized and induces anti-proliferative and genome-mending effects through trans-activation of a diverse set of P53-target genes (Boutelle and Attardi, 2021; Vogelstein et al., 2000). p53 mutations in cancer frequently occur in the DNA binding domain, where a single amino acid change results in dysfunctional DNA-binding and fails transcriptional activation (Olivier et al., 2010). Therefore, mutated p53 defunct in p53 transcriptional regulation results in failed tumor suppression and cancer outgrowth. Furthermore, defective p53 signaling will permit future genomic insults to remain unchecked, thus fostering an environment primed for malignant potential. Since mutant-p53 lacks transcriptional activity through its inability to bind DNA, it cannot induce expression of its negative regulators, ARF, MDM2, and MDMX (Perry et al., 1993; Wu et al., 1993). Lack of negative feedback results in mutant-p53 stabilization. Accumulation of mutant p53 has been reported to elicit gain-of-function p53 activity. Thus, any gain-of-function activity in mutant-p53 cannot be studied with p53-genetic deletion. Many research groups have characterized gain-of-function phenotypes of mutant-p53 in cancer. For instance, mice expressing mutant-p53 have altered disease phenotype compared to p53-null models (Lang et al., 2004; Olive et al., 2004). Specifically in lung cancer model systems, mutant-p53 has

modulated SMAD3, NF κ B, p63, TGF β , and others to drive various hallmarks of cancer invasion, chemoresistance, metabolism, and propagation of a stem cell fate (Adorno et al., 2009; Gaiddon et al., 2001; Klemke et al., 2021). Although the mechanisms that lead to these altered processes are controversial, it is thought that the main effect is through mutant-p53 binding and sequestering known p53-associated proteins. This often leads to the activation of transcription factors such as NF-Y (Kim et al., 2021).

In BRAF^{V600E} GEMMs, when p53 function is abolished by *Trp53* gene deletion or by *Cdkn2a* deletion, lung tumors can escape growth arrest and undergo malignant transformation (Dankort et al., 2007; Shai et al., 2015). Furthermore, p53 pathway dysregulation allows a cell to sample diverse phenotypic landscapes that allow for enhanced cell fitness and tumor outgrowth (Marjanovic et al., 2020). However, BRAF^{V600E} lung tumorigenesis and progression to malignant lung adenocarcinoma have only been studied in the context of p53 genetic deletion, not p53 mutation, which is more frequently seen in human lung cancer. Specifically, p53-R172H and p53-R245W are murine homologs of typical hotspots in human cancer R175H and R245W, respectively.

Here, we characterize the differential impact of p53(null), as well as p53 hotspot mutations p53^{R712H} (R175H in humans) and p53^{R245W} (R248W in humans) on BRAF^{V600E} lung tumorigenesis. We observed significantly differential effects of p53-R172H on BRAF^{V600E}-driven lung tumorigenesis that were not observed when p53 is deleted, such as enhanced tumor size and decreased survival. Our findings highlight functional differences in mutant-p53 biology, as well as the importance and the utility of clinically relevant mutant-p53 mouse models.

Results

P53-mutant Alleles Differentially Alter BRAF (V600E) Lung Tumorigenesis

Expression of BRAF^{V600E} in the distal epithelium of the mouse lung elicits clonal tumorigenic outgrowths of alveolar type 2 (AT2) pneumocytes (Dankort et al., 2007). Prior work has shown that *Trp53*^{f/f} cooperates with BRAF(V600E) to form malignant lung tumors when Cre recombinase is activated (Dankort et al., 2007; Shai et al., 2015). Therefore, we compared the tumor burden of *Braf*^{Cat/+} mice with either p53 loss (*Trp53*^{f/f}), p53^{R172H} (*Trp53*^{R172H/R172H}), or p53^{R245W} (*Trp53*^{R245W/R245W}) after infection with 106 pfu of adenovirus expressing surfactant protein C specific Cre (ad5-Spc-Cre) (Figure 2). As expected, mice developed lung cancer, consistent with previous findings (Figure 8 B-C). However, there was a striking, highly reproducible increase in the tumor size of tumors harboring *Trp53*^{R172H/R172H} and to a lesser extent *Trp53*^{R245W/R245W}, compared to *Trp53*^{f/f} (Figure 8 A,B). These data led us to hypothesize that p53^{R172H} and p53^{R245W} expression alters tumor progression uniquely from p53 deletion, and these differences may be potentially evident of gain-of-function effects imparted by mutant-p53. We next looked at early time points to determine if p53 mutation affects the growth and proliferation of early stage lung tumors. However, we did not find any differences in tumor size between the three different p53 alleles (Figure (fig3.3?) A, B).

Since lung cancer patients with mutations in *TP53* often start only have one mutated *p53* allele unless the loss of heterozygosity (LOH) occurs. Therefore, we generated mice heterozygous for both *Braf*^{CAT} and either *Trp53*^f, *Trp53*^{R172H}, or *Trp53*^{R245W} and observed their survival over time (Figure 10 A). We noted that *Trp53*^{R172H/+} mice had diminished median overall survival compared to *Trp53*^{f/+} mice (183 days and 216 days), with *Trp53*^{R245W/+} mice falling in between the two groups (median survival 212 days) (Figure 10 A).

We observed a distinct separation of survival between mice harboring lung tumors that are either *Trp53*^{f/+}, *Trp53*^{R172H/+}, and *Trp53*^{R245W/+}. This separation is diminished when altered p53 alleles are homozygous (Figure 10 B). *Trp53*^{R172H/R172H} mice have a median survival of 120 days, while *Trp53*^{f/f}

mice have a median survival of 136 days. Importantly, this observed difference in survival is statistically significant (Log-rank test p< 0.001) and is likely to reflect the worse diseased, characterized by enhanced tumor burden and size.

Mutant-p53 Accumulation is Associated with Dedifferentiation and p19 Induction Increased malignancy in lung adenocarcinoma is often associated with altered states of tumor cell identity and dedifferentiation (Marjanovic et al., 2020; van Veen et al., 2019). Since LUADs often arise from AT2 cells, we surveyed the expression of AT2 cell identifiers, such as NKX2-1, a master regulator of AT2 transcriptional programs, and SPC, a marker of differentiated AT2 cells (Marjanovic et al., 2020). Quantification of over 50 tumors from *Braf^{V600E/+}* mice with either *Trp53^{f/f}*, *Trp53^{R172H/R172H}*, and *Trp53^{R245W/R245W}* revealed that *Trp53^{R172H/R172H}* mice had significantly more tumors that stained negative for NKX2-1 and SPC (Figure 11 A, B). This finding strongly indicates that mutant-p53 tumors are dedifferentiated. Since HMGA2 expression is associated with LUAD dedifferentiation and worse patient prognosis, we performed IHC for HMGA2(Di Cello et al., 2008; Gao et al., 2017; Winslow et al., 2011). We noted an increase in HMGA2-positive tumors in *Trp53^{R172H/R172H}*, and *Trp53^{R245W/R245W}* tumors compared to *Trp53^{f/f}* (Figure 11 A, B). Thus, this data reveals that *Trp53^{R172H/R172H}* mice exhibit a higher tumor burden and worse overall survival than *Trp53^{f/f}*. However, the tumors developed in these mice are more likely to exhibit malignant progression and dedifferentiation.

One striking feature of lung tumors in mice, either heterozygous or homozygous for *Trp53^{R172H}* or *Trp53^{R245W}* is how few tumors and tumor cells have detectable levels of mutant-p53 protein as measured by Immunohistochemistry. One critical aspect of normal p53 function is self-regulation through transcriptional activation of its negative regulator MDM2 (Wu et al., 1993). Thus, we predicted that homozygous p53 mutant cells would exhibit increased mutant-p53 protein. In our models, we observed that mutant p53 protein accumulation is restricted to malignant cells exhibited by a lack of papillary architecture and nuclear atypia (Figure 12 A). Indeed, p53 immunostaining is correlated with increased phosphorylated-ERK1/2 detection (Figure 12 B). Moreover, immunofluorescence of p53 and its positive regulator, p19/ARF, reveal that cells with high p19/ARF are also high in p53 accumulation (Figure 12 C). This result suggests that even in homozygous mutant-Trp53_ mice, there is still a barrier to mutant-p53 accumulation. Our data suggest that elevated levels of p19/ARF are necessary for mutant-p53 accumulation, even with homozygous p53 mutations. Collectively, these data indicate that *Trp53^{R172H}* drives more aggressive lung adenocarcinoma formation in BRAF(V600E) driven mouse models of lung cancer.

P53^{R172H} and P53^{R245W} cell lines are refractory to BRAF/MEK inhibition Given that previous research has determined that p53 mutations can confer resistance to both chemotherapies and pathway-targeted therapies (Huang et al., 2011; Rusch et al., 1995), we next questioned whether our p53 alterations would be sufficient to drive therapeutic resistance in BRAF^{V600E}-driven genetically engineered mice(Planchard et al., 2016). Interestingly, *Trp53^{R172H/R172H}* mice retain a more considerable tumor burden after a week four treatment of MAPK blockade *in vivo* (Figure 13 A).

To further investigate the effects of p53-mutants on *BRAF^{V600E}* driven lung cancers, we derived cell lines from *Braf^{CAT/+}* murine tumors expressing either *Trp53^{f/f}*, *Trp53^{R172H/R172H}*, or *Trp53^{R245W/R245W}* 14 weeks after infection with ad5-Spc-Cre. Cell lines expressing either *Trp53^{R172H/R172H}* or *Trp53^{R245W/R245W}* were resistant to single-agent dabrafenib (Figure 13 B). Of note, *Trp53^{R172H/R172H}* and *Trp53^{R245W/R245W}* cell lines express increased ERK1/2 activity, as evidenced by increased phosphorylated ERK1/2, both with and without MAPK-blockade (Figure 13 C). Furthermore, *Trp53^{R172H/R172H}* and *Trp53^{R245W/R245W}* were more resistant to the combination of dabrafenib plus trametinib, than *Trp53^{f/f}* cell lines(Figure 14 A-C). Not only are resistant cells still proliferative in the face of MAPK-blockade, they are resistant to cell death imparted by MAPK-inhibition[Figure 14 C].

Next, we reasoned if mutant-p53 expression is the cause differential cancer phenotypes such as increase proliferation and drug resistance, then eliminating mutant-p53 expression would have a growth inhibitory effect and sensitize cells to MAPK-pathway inhibition. We cloned a short-hairpin RNA plasmid that induces GFP expression and knockdown of mutant-p53 with supplementation of doxycycline(Figure 15 A). Despite sucessful p53 knockdown, observed no change in *in vitro* proliferation or *in vivo* tumor outgrowth(Figure 15 B and C). Therefore, mutant-p53 cells are not addicted to mutant-p53 expression.

To gain relevance to human lung cancer, we infected BRAF(V600E) mutant HCC364 cells with lentivirus expressing human cancer hotspot mutant p53^{R175H}. p53^{R175H} expressing cells were more proliferative and refractory than parental HCC364 cells (Figure 16 A and B). Collectively, these *in vitro* and *in vivo* data suggest that missense mutant p53^{R175H} and its murine homolog p53^{R172H}, as well as p53^{R245W} can support cancer cell viability in the face of BRAF+MEK blockade that p53-null mutations cannot achieve.

P53^{R172H} and P53^{R245W} alter the transcriptome of BRAF(V600E) lung tumor cell lines Due to our observations that mutant p53^{R172H} drives more aggressive cancer phenotypes such as increased growth, dedifferentiation, and resistance to FDA-approved therapeutics, we reasoned that this would be reflected in a change in the transcriptional landscape. This is due in part to the transcriptional trans-activation potential by mutant p53. RNA sequencing of cell lines generated from p53-mutant mouse lung tumors does reveal transcriptomic differences between both p53^{R172H} and p53^{R245W} cells compared to p53-null cell lines (Figure 17 4 A and B). Interestingly, genes involved in cell cycle regulation and MYC pathway are upregulated in both p53^{R172H} and p53^{R245W} cell lines (Figure 4 17 B). These results suggest that mutant-p53 affects the transcription of genes involved in pro-growth signaling pathways. We also observed transcriptional differences between *Trp53*^{R172H/R172H} and *Trp53*^{R245W/R245W} cell lines(Figure 18 A and B). Thus, we hypothesize that *Trp53*^{R172H/R172H} and *Trp53*^{R245W/R245W} have similarities in their function, but they also have some clear differences.

PDGFR expression is increased in mutant-p53 cells Our *in vitro* RNA analysis revealed an upregulation of the platelet-derived growth factor receptor(PDGFR) in *Trp53*^{R172H/R172H} cell lines(Figure 19 A). Since PDGFR is a known regulator of the MAPK pathway, we sought to validate its expression and determine its important in *Trp53*^{R172H/R172H} and *Trp53*^{R245W/R245W} cell lines. Indeed PDGFRB is more highly expressed and can be activated by recombinant PDGF stimulation(Figure 20 B). Although *Trp53*^{f/f} cells do not have any detectable levels of PDGFR or Phosphohorylated-PDFRB, they still upregulated MAPK-pathway activation in response to ligand stimulation(Figure 20 B). Furthermore, expression of PDGFRB is evident in malignant *Trp53*^{R172H/R172H} and *Trp53*^{R245W/R245W} tumors, but not *Trp53*^{f/f} tumors(Figure 20 C). We next sought to determine the sensitivity to PDGFR-signaling blockade with Imatinib, a known PDGFRB inhibitor. Neither PDGFR stimulation or PDGFR inhibition had any effect on the proliferation of mutant-p53 cell lines(Figure ?? A an B). Therefore, PDGFR is a validated differentially expressed gene in *Trp53*^{R172H/R172H} and *Trp53*^{R245W/R245W} cell lines, but is not important for its growth.

Discussion

While lung adenocarcinomas primarily contain missense mutations in p53, most studies using mouse models of lung adenocarcinoma utilize a p53-null (*Trp53*^{f/f}) allele instead of alleles encoding hotspot missense mutations (Gao et al., 2013). This is most likely because prior mutant-p53 alleles were null for *Trp53* prior to Cre-mediated recombination, leaving every cell in the mouse sensitive to leukemias, lymphomas, and sarcomas due to being partially p53-deficient (Jackson et al., 2005; Olive et al., 2004; Wijnhoven et al., 2005). Our work has been made possible by newly described “wild-type to mutant” p53 alleles that allow for mice to be fully wild-type for p53 in all cells prior to Cre-mediated recombination(Zhang et al., 2018). This strategy is crucial as it allows for temporally extended

experiments not confounded by separate malignancies. Here, we utilize " wild-type to mutant" $Trp53^{R172H}$ and $Trp53^{R245W}$ alleles R245W to determine how they alter tumor progression in a BRAF(V600E) model of lung cancer. We show that $p53^{R172H}$ and $p53^{R245W}$ accelerate lung tumor progression and increase the rate of dedifferentiation beyond the complete loss of p53 expression.

Although point mutations in the *TP53* gene can poison the natural tumor suppressive mechanisms of p53 protein, not all point mutations are equivalent in executing these functions. Mutations in the DNA-binding domain are most common and potent at ablating normal p53 function. Taking advantage of a conditional BRAF^{V600E} lung tumor model where BRAF^{V600E} expression in AT2 cells drives benign tumor formation, we show that either $p53^{R172H}$ or $p53^{R245W}$ can bypass growth arrest and influence tumor progression and therapeutic sensitivity beyond how p53 loss can. Specifically, $P53^{R172H}$ reduces survival and sensitivity to dabrafenib plus trametinib *in vivo*. Interestingly, a single allele of $p53^{R172H}$ has a more robust tumor-promoting phenotype than $P53^{R245W}$. These findings further support a hierarchy model of p53 missense mutations, where conformation mutants such as $p53^{R175H}$ have a more substantial dominant-negative effect on wild-type p53, while mice with a single $trp53^{fl}$ allele can still suppress tumorigenesis with one copy of wild-type but are prone to loss of heterozygosity (Muzumdar et al., 2016; Olive et al., 2004; Zhang et al., 2018).

In BRAF-driven lung adenocarcinoma, the repressive role of wild-type p53 is initiated by oncogenic stress brought about by increasing levels of MAPK-pathway activation(Dankort et al., 2007). Hyperactivation of the MAPK pathway triggers p19/ARF to block MDM2, leading to p53 accumulation(Christophorou et al., 2006; Feldser et al., 2010; Junntila et al., 2010). Therefore, p53 acts late in lung adenocarcinoma, unlike pancreatic adenocarcinoma, where p53 is triggered early, likely through PDAC-specific increases in p19/ARF through increased PI3K, STAT3, or DMP1 (Muzumdar et al., 2016). Our studies in BRAF(V600E) mutant lung adenocarcinomas reinforce this concept, as mutant-p53 does not accelerate growth in early tumors (supplemental figure). Furthermore, p53 is not detected until later stage tumor development and correlates with p19/ARF expression and increased phosphorylated-ERK1/2 levels. Our studies indicate that mutant-p53 levels are associated with increased expression of p19/ARF. Therefore, any gain-of-function phenotype manifested by mutant-p53 must depend on accumulated p19/ARF. Thus, ablating the ARF tumor suppressor in human cancers with GOF *TP53* mutants may paradoxically lead to tumor regression.

While most *TP53* mutations in cancer result in an inability to bind DNA, they are often stable and preserve functional oligomerization and transcriptional transactivation domains(Garcia and Attardi, 2014). Thus, mutant-p53 can still interact with and affect the function of cellular processes. Supporting this rationale, we observe altered transcriptomes with both $p53^{R172H}$ and $p53^{R245W}$ cell lines compared to p53-null. Consistent with previous mutant-p53 GOF studies, we observed enriched genes associated with EMT and cell-cycle entry with $p53^{R172H}$ and $p53^{R245W}$.

Many GOF phenotypes exhibited by mutant-p53 have centered around drug-resistant mechanisms. Previous studies have shown that mutant-p53 can confer resistance to pathway-targeted therapy and chemotherapies. Our studies highlight that $p53^{R172H}$ and $p53^{R245W}$ can confer resistance to BRAF-targeted therapy. Here, we show p53-R172H and p53-R245W cell lines are resistant to BRAF(V600E) inhibition and have less sensitivity to the combination of BRAF+MEK inhibition. In pancreatic cancer models with LSL-p53-R172H expression, mutant-p53 regulates PDGFR-signaling to promote an invasive phenotype sensitive to Imatinib(Weissmueller et al., 2014). Consistent with these findings, our transcriptional analysis and immunoblotting indicate p53-R172H, and p53-R245W cells express PDGF-ligands and exhibit PDGFR-pathway activation.

In summary, we have compared the tumorigenic potential of three *Trp53* alleles ($Trp53^{fl/fl}$, $Trp53^{R172H}$, and $Trp53^{R245W}$) concurrently with BRAF^{V600E}. $Trp53^{R172H}$ and $Trp53^{R245W}$ promoted more aggressive lung cancers exhibited by larger tumors and tumor dedifferentiation. However, only $Trp53^{R172H}$ was observed to shorten survival compared to $Trp53^{fl/fl}$. Moreover, $Trp53^{R172H}$ and

Trp53^{R245W} cell lines were refractory to dabrafenib and trametinib and were more proliferative. *Trp53^{R172H}* and *Trp53^{R245W}* cell lines had altered transcriptomes compared to p53-null cells and were highly enriched in genes involved in cell cycle proliferation. Our work highlights the importance of studying disease-relevant *TP53* alleles in pre-clinical modeling systems.

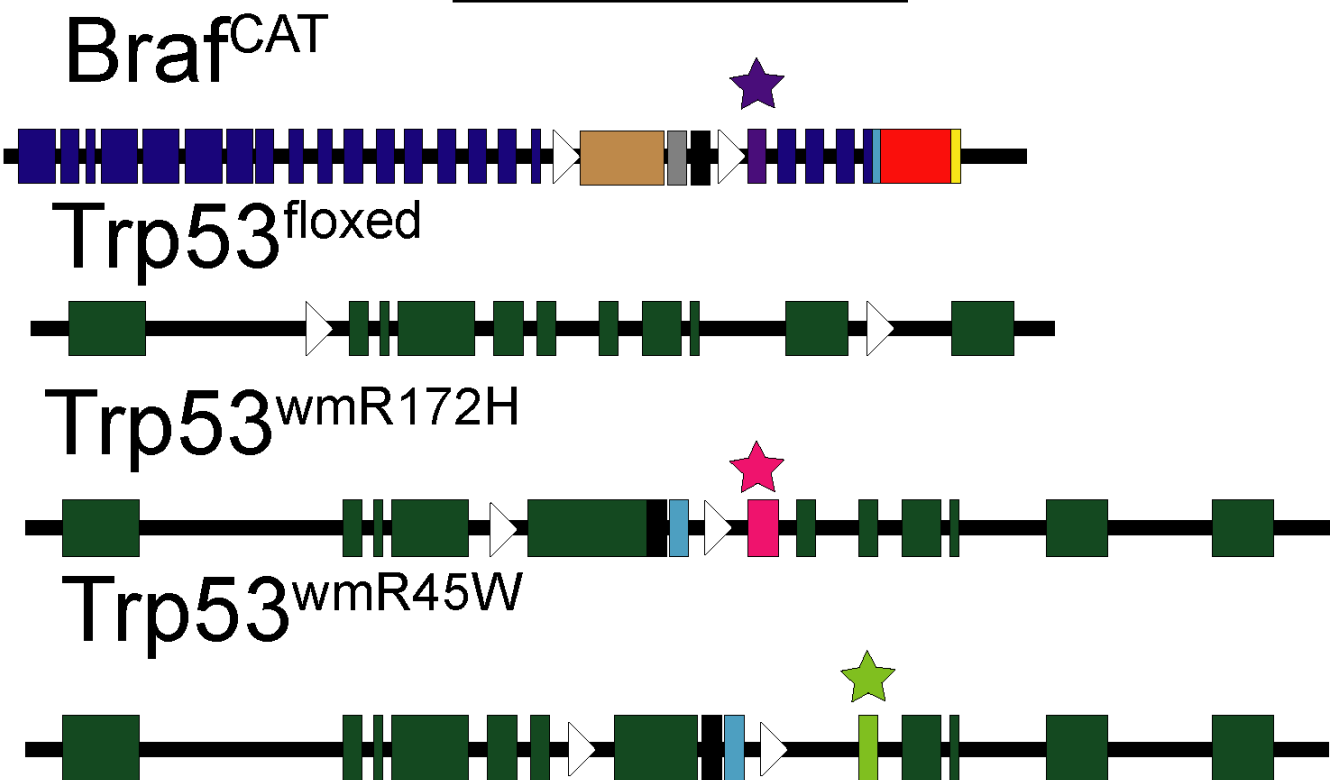
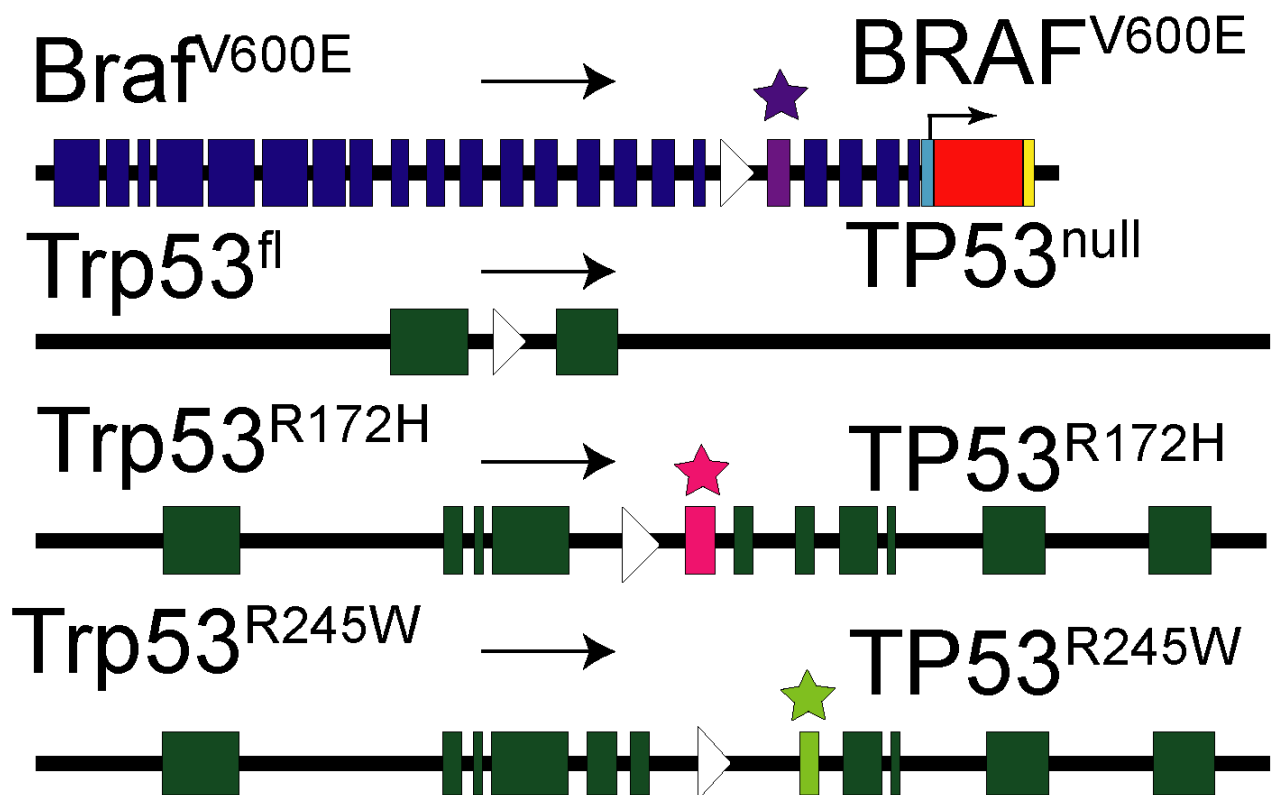
A**Before Cre****B****After Cre**

Figure 7: Cre-mediated wildtype-to-mutant alleles used in study. (A) Prior to Cre-mediated recombination, normal *Braf* and *Trp53* are expressed from their endogenous loci. (B) After Cre-mediated recombination, the indicated mutant proteins are expressed, in addition to a TdTomato fluorescent protein to mark expression of *BRAF^{V600E}*.

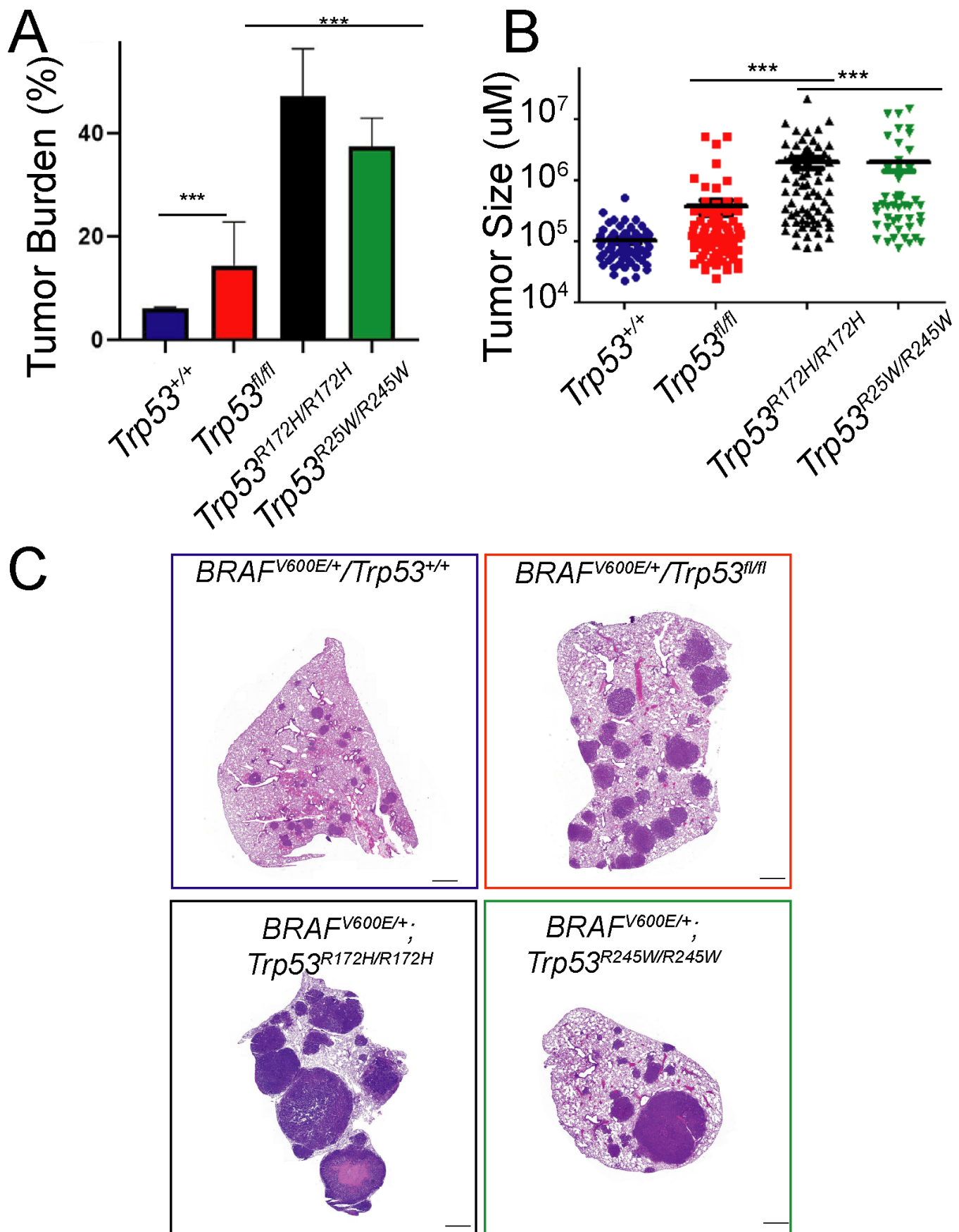


Figure 8: Mutant-p53 alleles differentially alter $\text{BRAF}^{\text{V600E}}$ driven lung tumorigenesis. (A) Tumor burden analysis 14-weeks post Cre-recombinase. (B) Tumor size 14 weeks post Cre-Recombinase. (C) Representative histological images 14 weeks post Cre-Recombinase. *** = $p\text{-value}<.01$

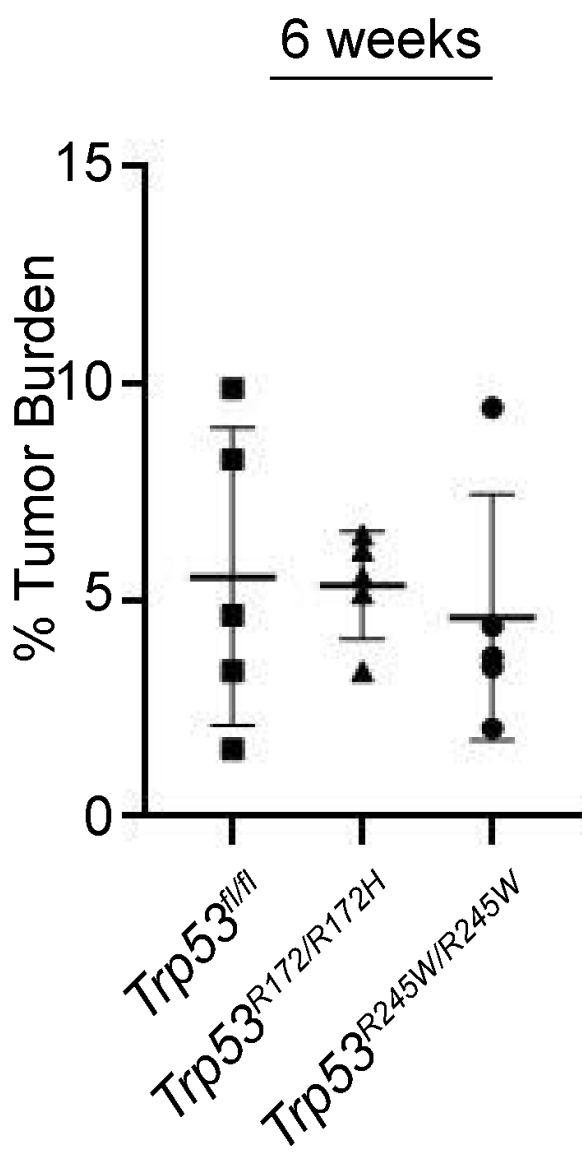
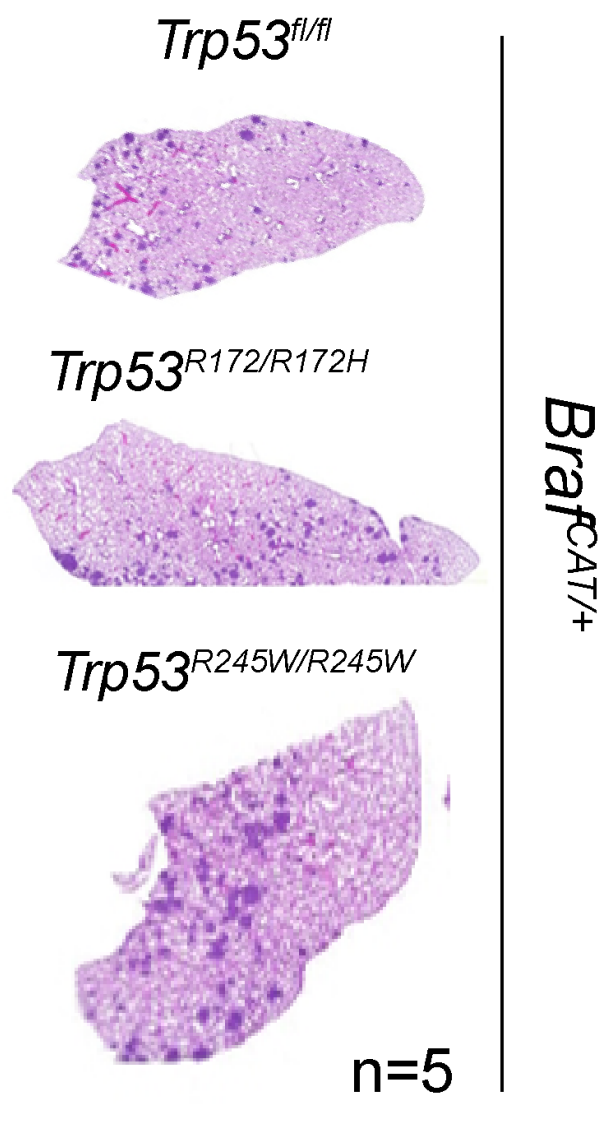
A**B**

Figure 9: Mutant-p53 does not alter the growth of early stage tumors (A) Tumor burden calculation after 6 weeks of tumor growth (B) Representative images of H&E stained lung sections.

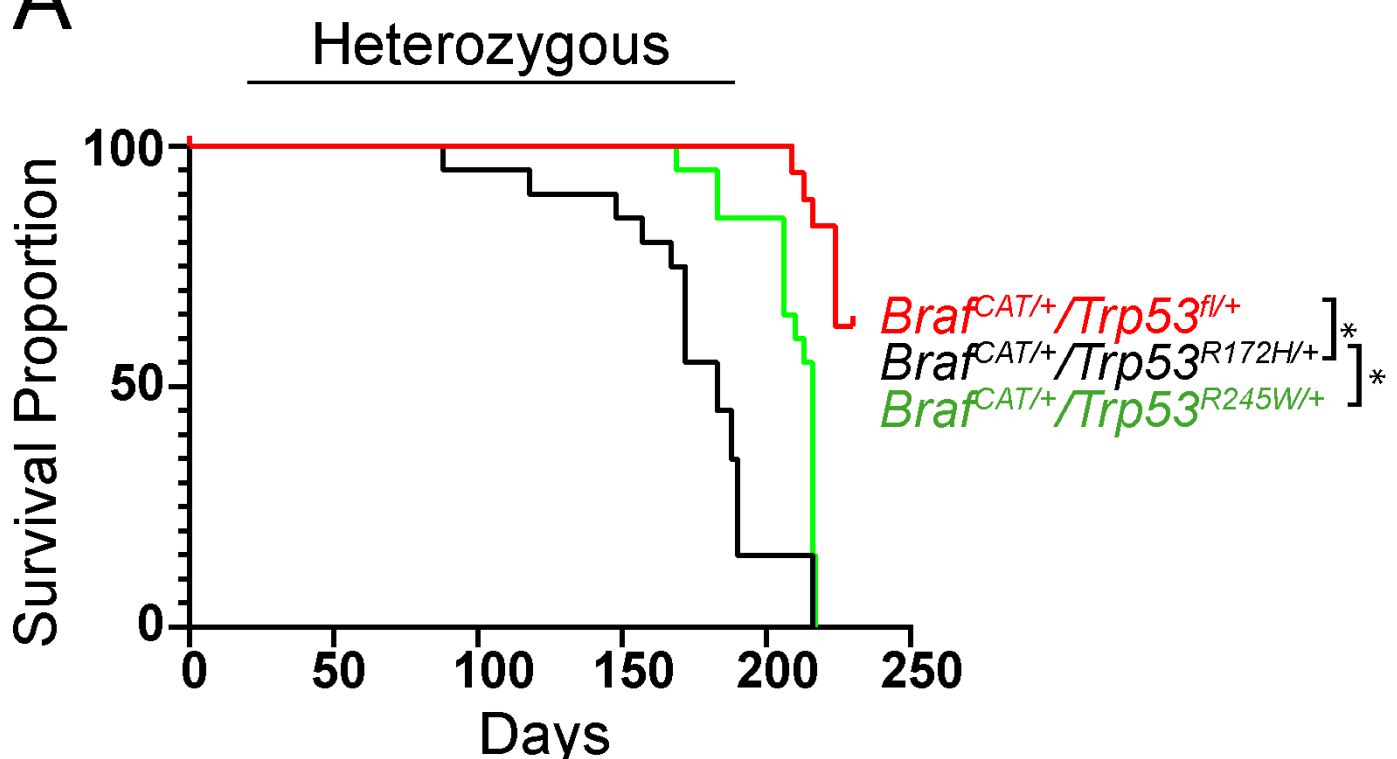
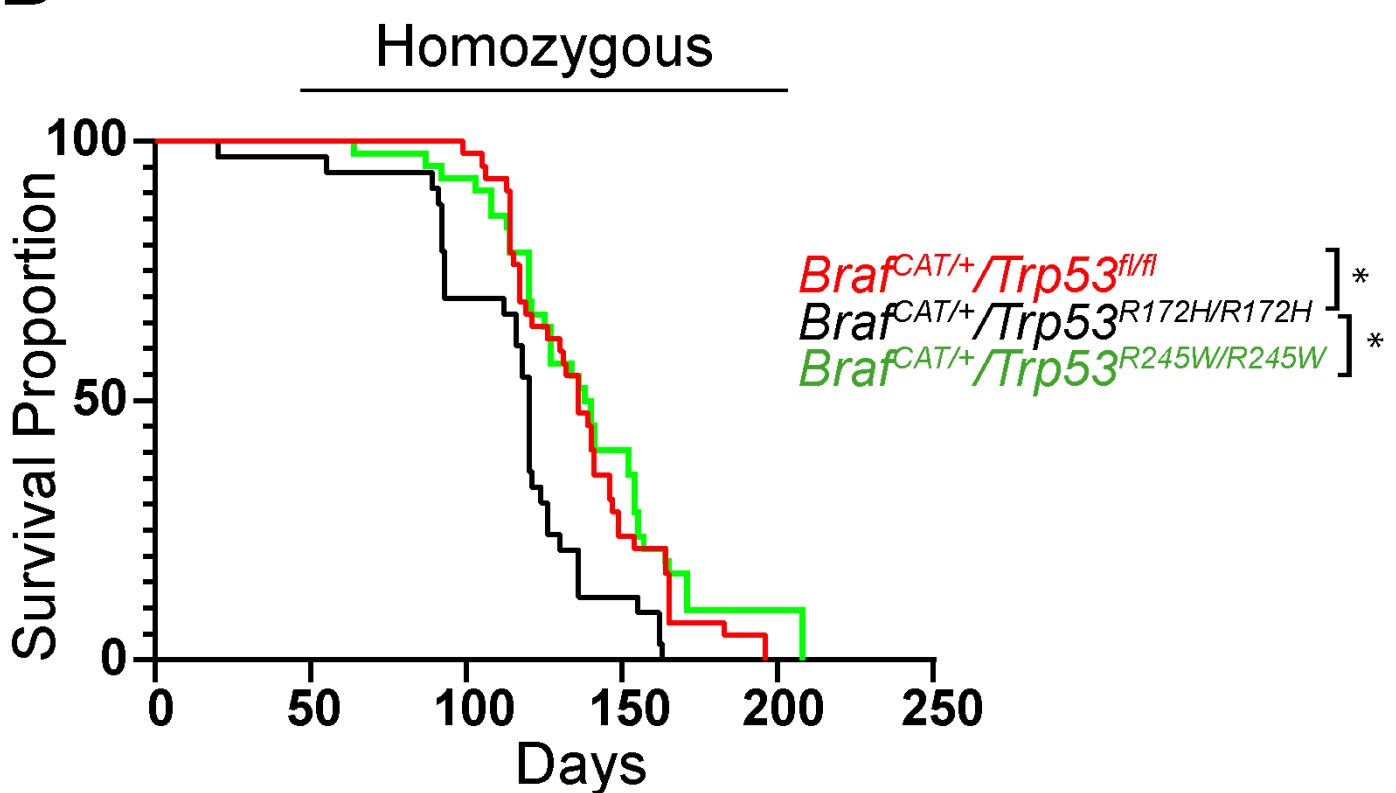
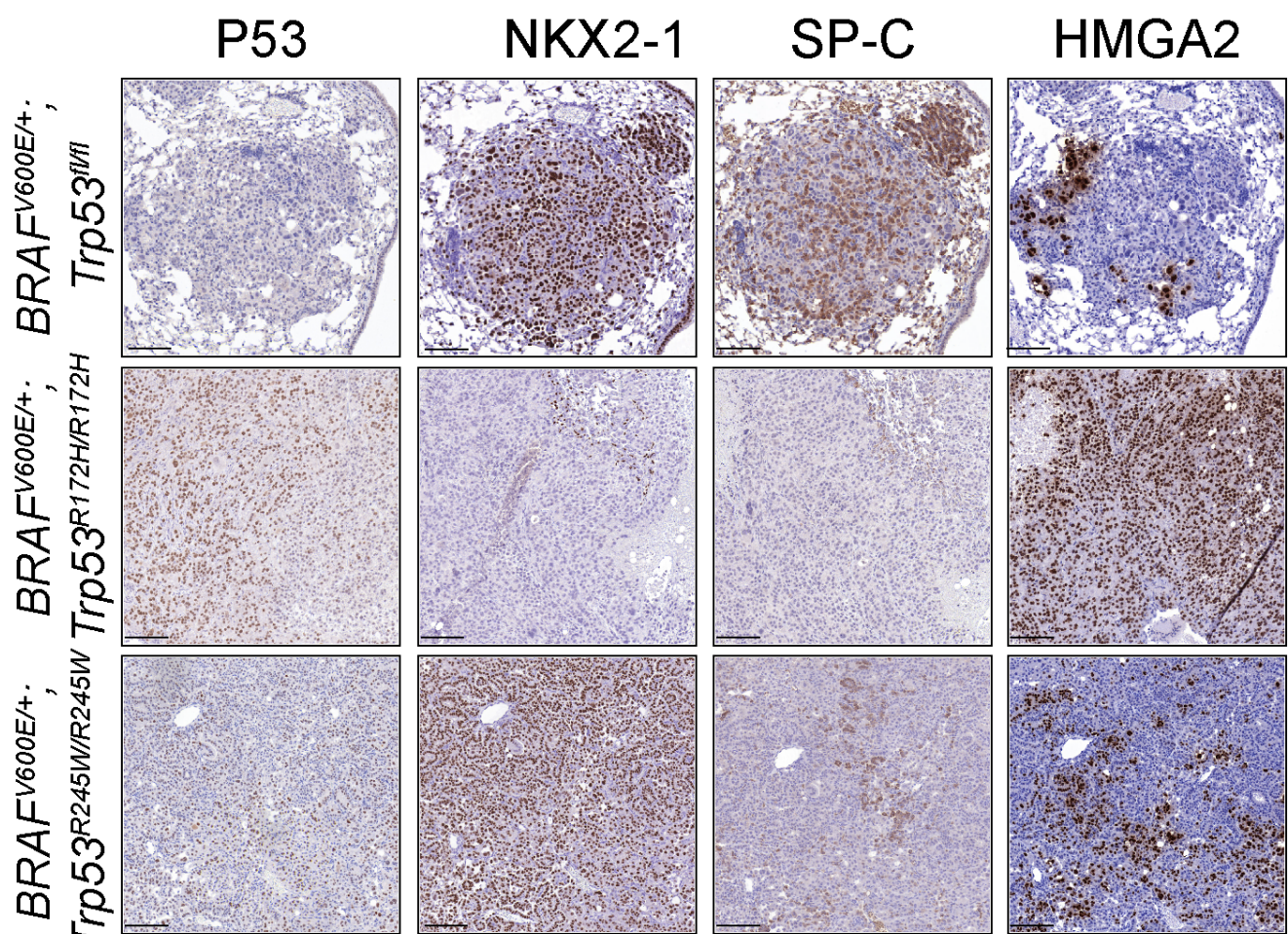
A**B**

Figure 10: Mutant-p53 alleles differentially affect the survival of mice due to progressive lung adenocarcinoma. (A) Kaplan-Meier plot of $Braf^{CAT/+}$ mice with p53-mutant alleles. (B) Kaplan-Meier plot of $Braf^{CAT/+}$ mice with homozygous p53-mutant alleles.* = p -value < .05

A



B

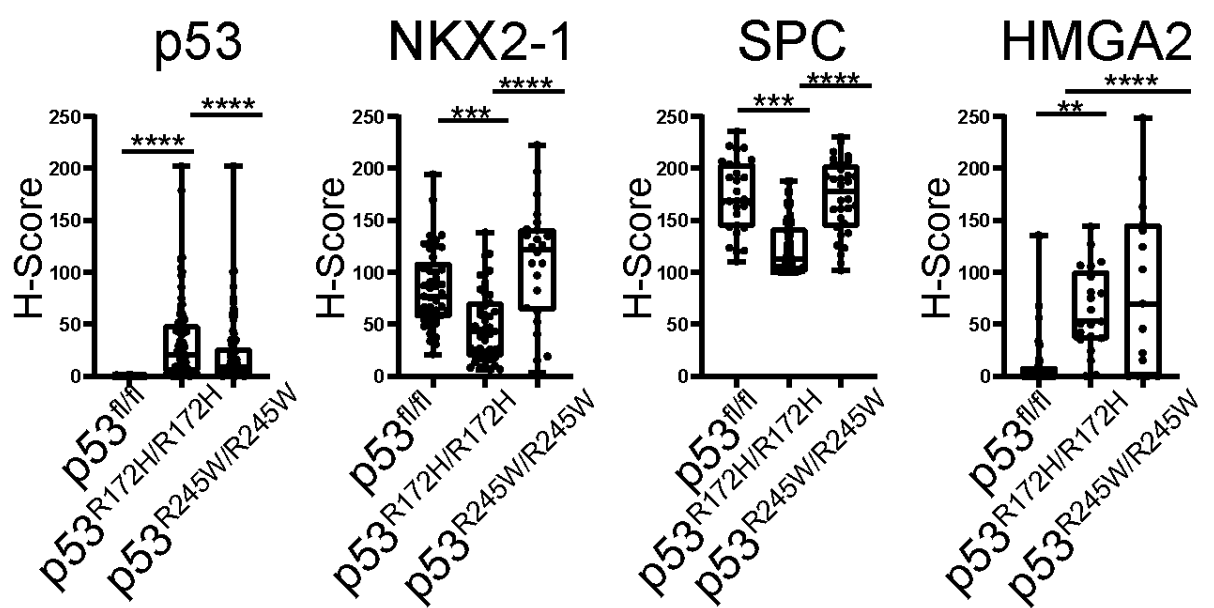
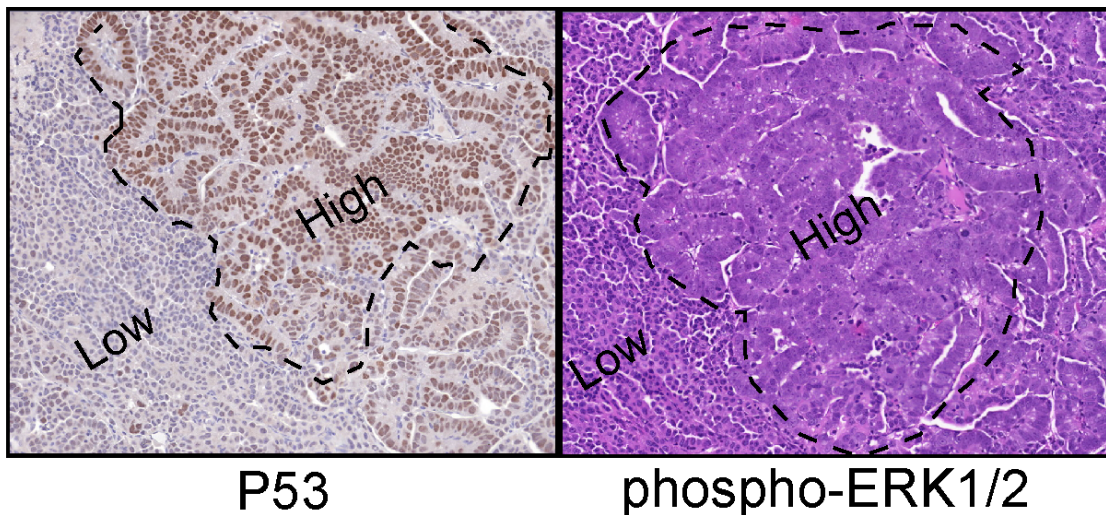


Figure 11: Missense mutant-p53 alters lung adenocarcinoma differentiation status. (A) Representative IHC images of tumors stained with antibodies targeting p53, NKX2-1, SPC, and HMGA2. (B) Quantification of immunostaining in (A) based on H-Score. *** = p -value < .01 ; **** = p -value < .001.

A

P53

phospho-ERK1/2

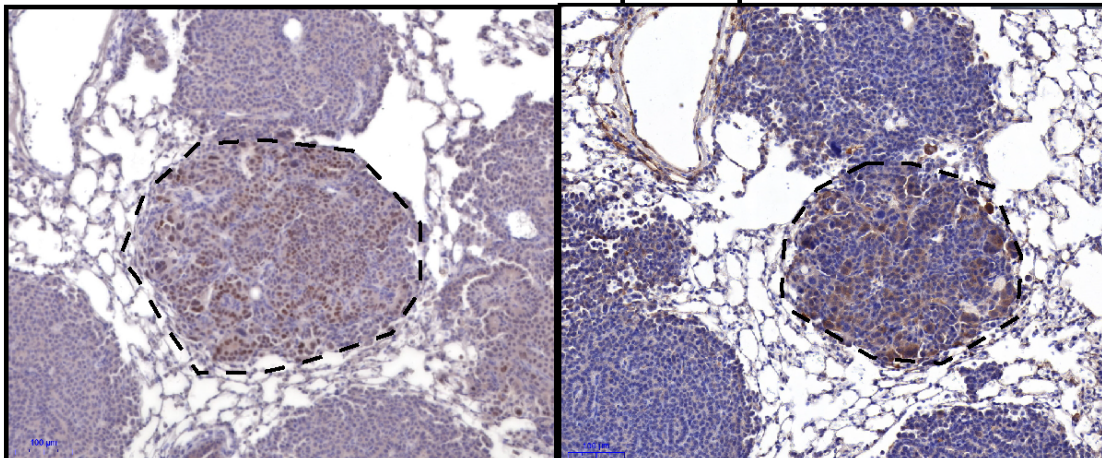
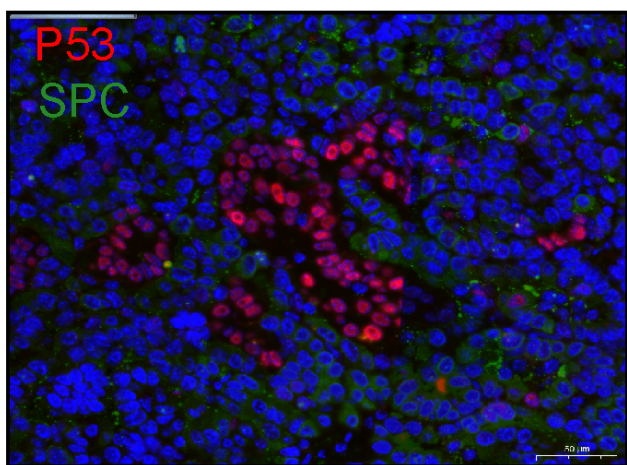
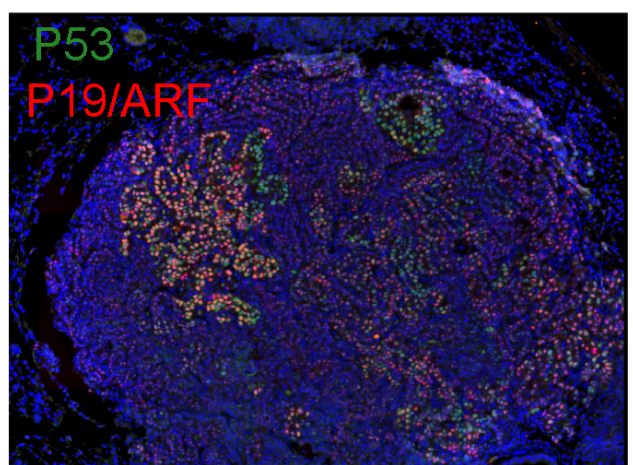
B**C****D**

Figure 12: p53-accumulation occurs in high grade tumors and correlates with p19/ARF expression. (A) Representative images of mutant-p53 accumulation in high grade tumors. the tumor area encased by the dashed line exhibit nuclear atypia and enlarged nuclei.(B) Mutant-p53 immunostaining correlates with phospho-ERK1/2 accumulation. (C) p53 immunostaining correlates with loss of SPC expression. (D) mutant-p53 is co-expressed with p19/ARF in p53-mutant tumors.

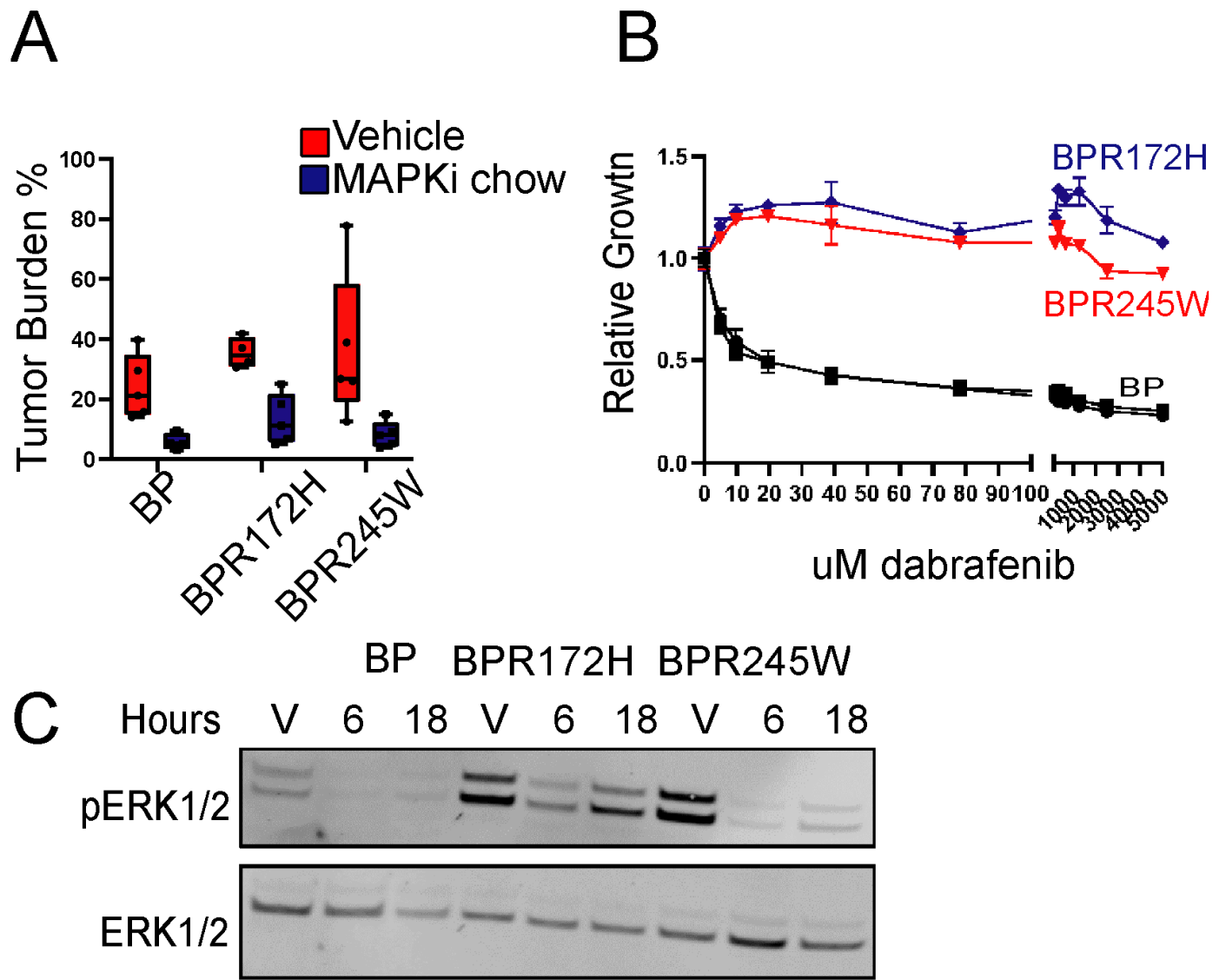


Figure 13: Mouse tumors and cell lines that express P53-R172H and p53-R245W are refractory to MAPK blockade. (A) Tumor burden analysis of GEMMs treated with MAPKi chow for four weeks. (B) *n* vitro dose response to dabrafenib for 48 hours. (C) Western blots of timecourse of cell lines treated with dabrafenib plus trametinib.

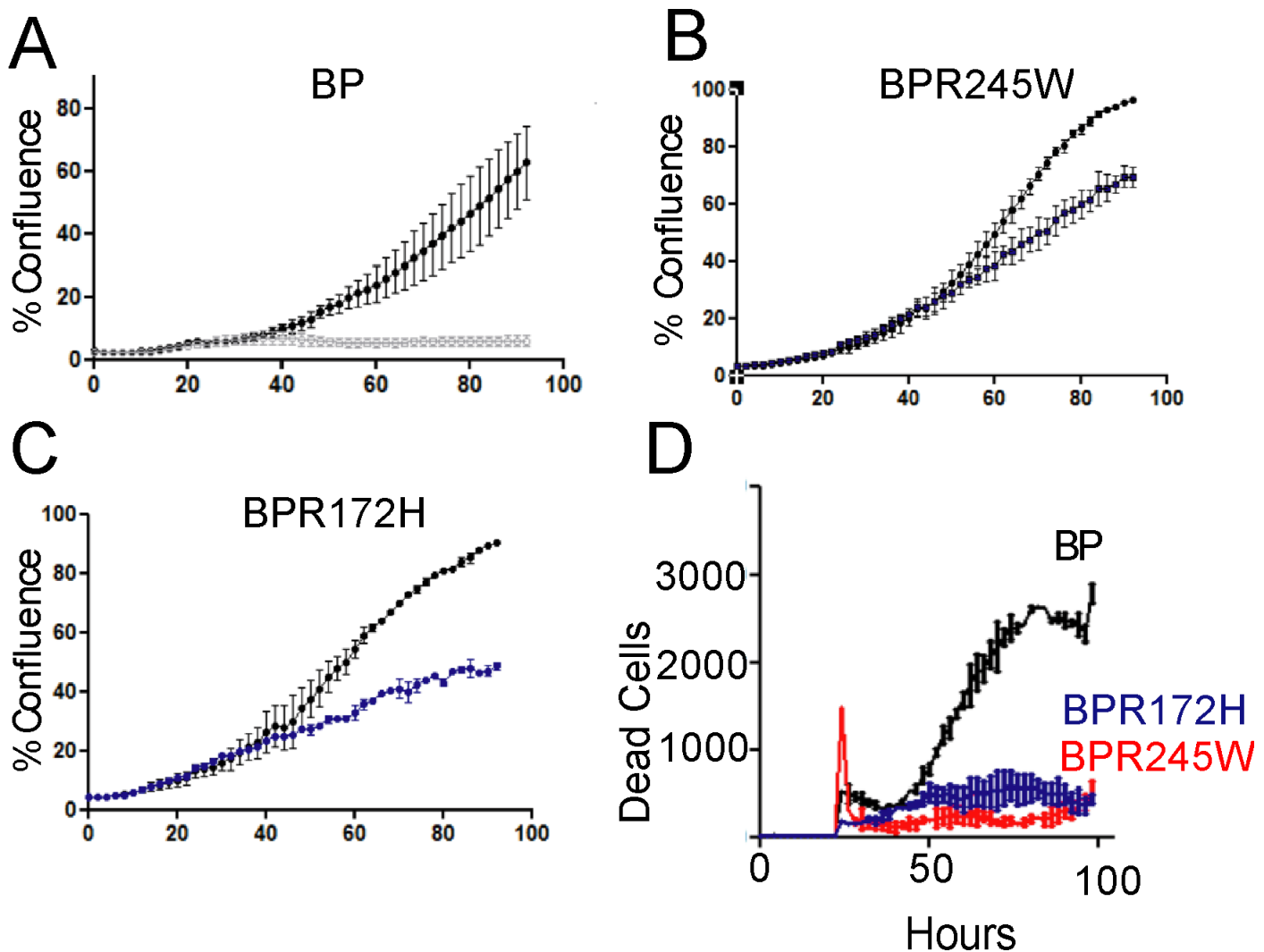


Figure 14: p53-R172H and p53-R245W cell lines are insensitive to MAPKi-induced cell death (A-C) Incucyte analysis of cell lines treated with the combination of dabrafenib plus trametinib for 96 hours. (D) Accumulation of dead cells in response to dabrafenib plus trametinib. Dead cells were quantified by CytoTox Red Dye.

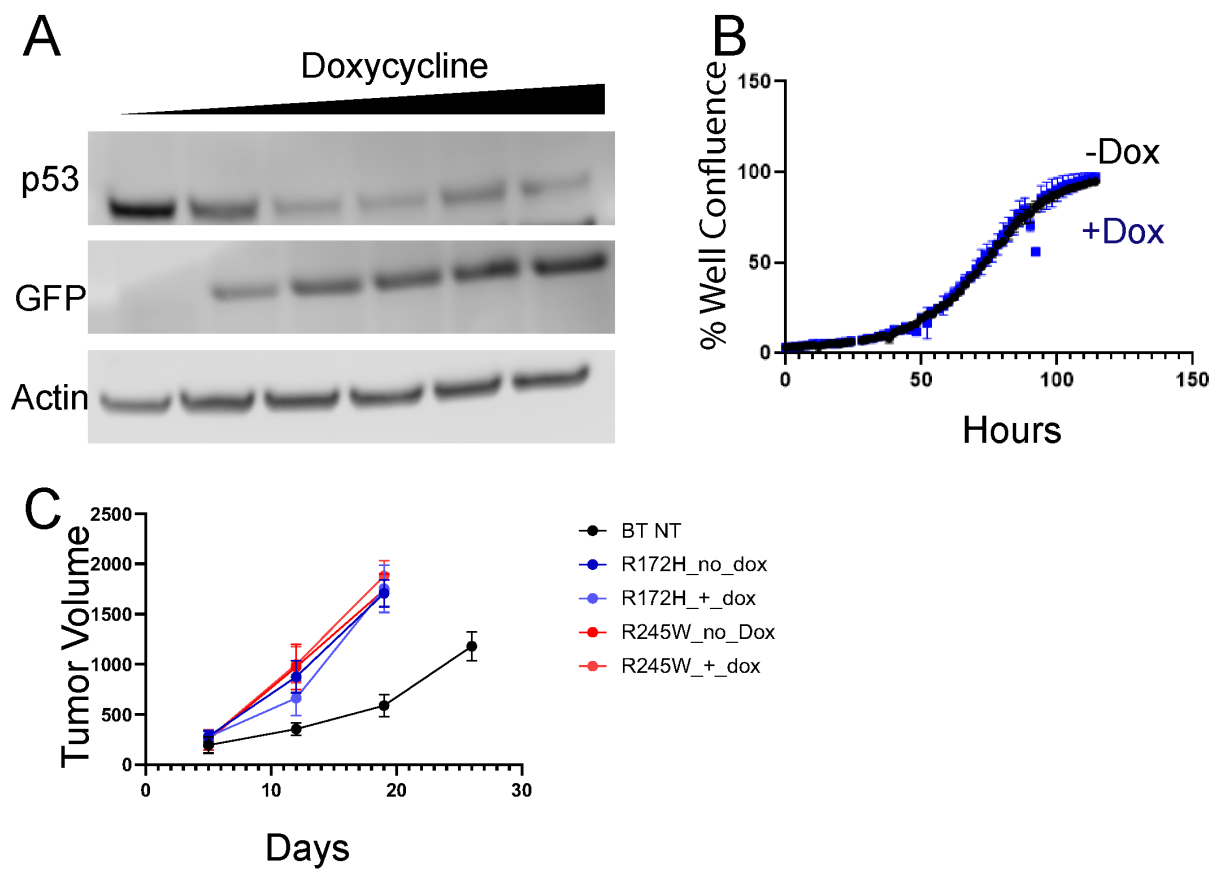


Figure 15: Mutant-p53 knockdown has no effect on cell proliferation or tumor growth. (A) Western blot of increasing Doxycycline concentration to silence mutant-p53 using a short-hairpin RNA (B) Cell proliferation assay with 4ug/mL doxycycline concentration. (C) *In vivo* response to mutant-p53 knockdown.

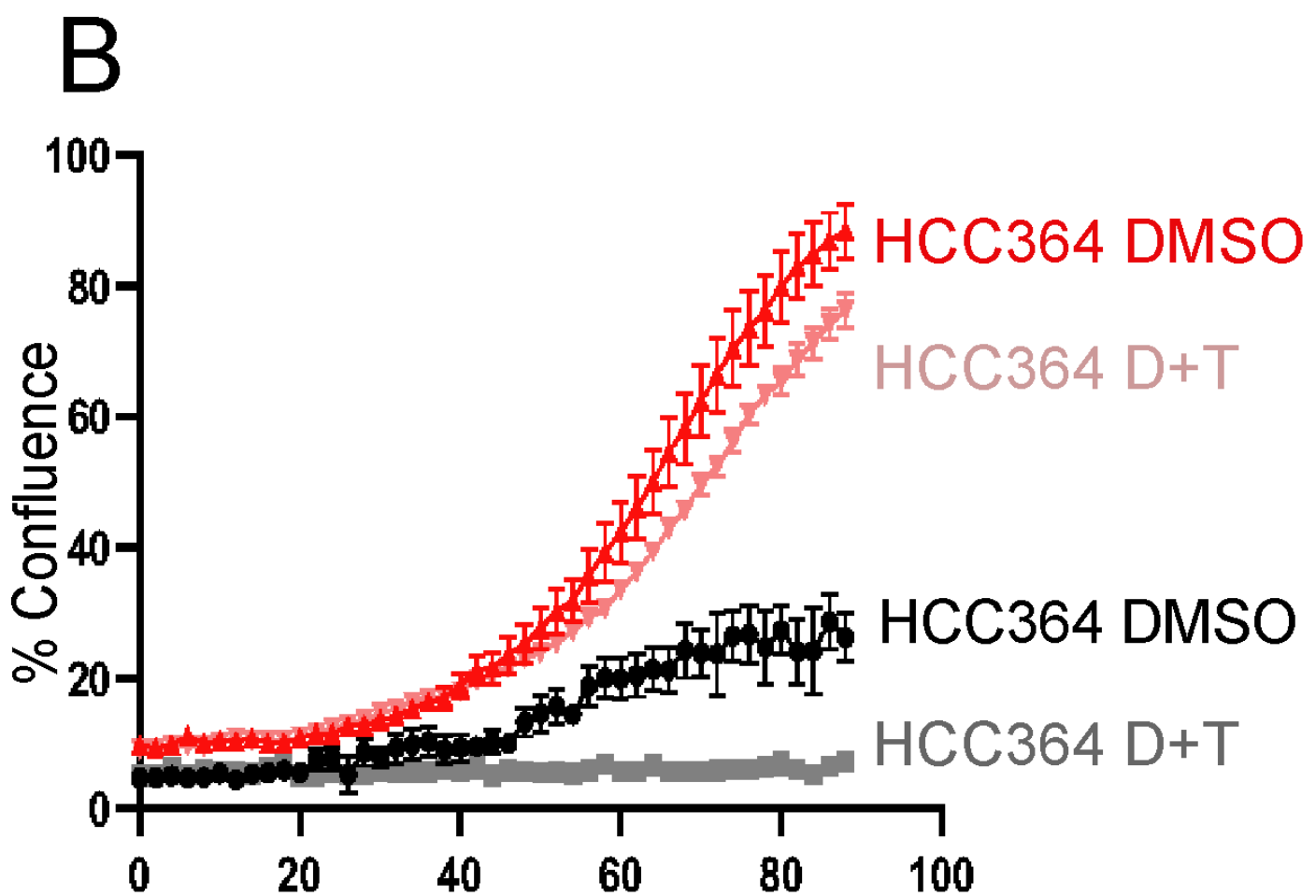
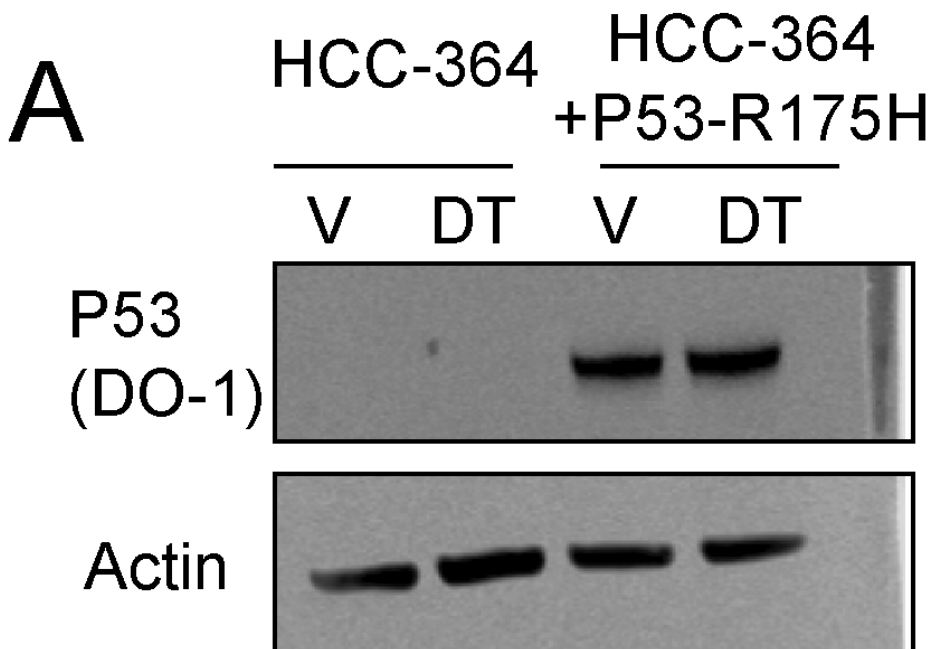


Figure 16: HCC-364 cells that express p53-R175H are refractory to MAPKi. (A) Western blot showing successful ectopic expression of p53-R175H. (B) Incucyte analysis of HCC364 parental and HCC364+p53-R175H in response to dabrafenib plus trametenib.

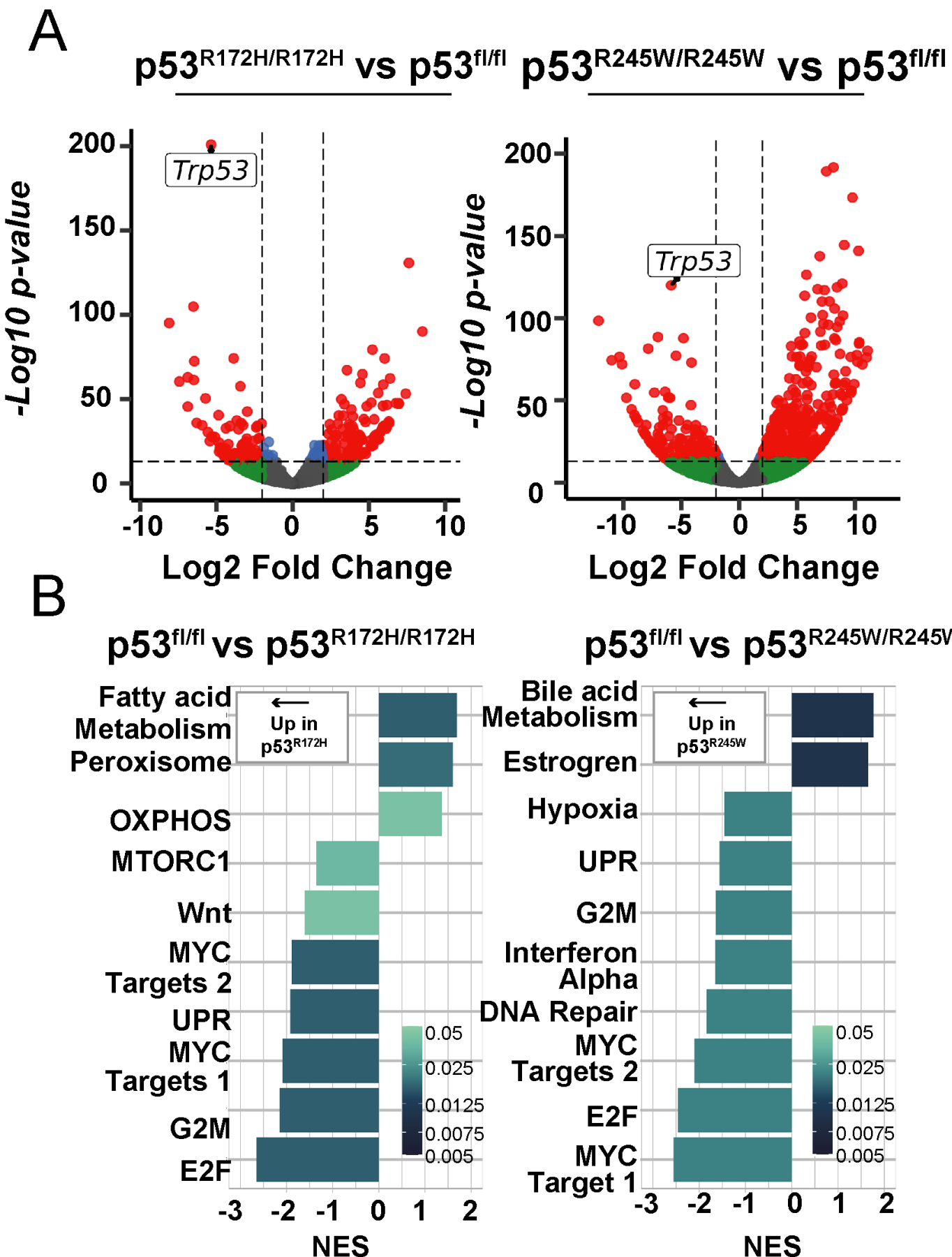
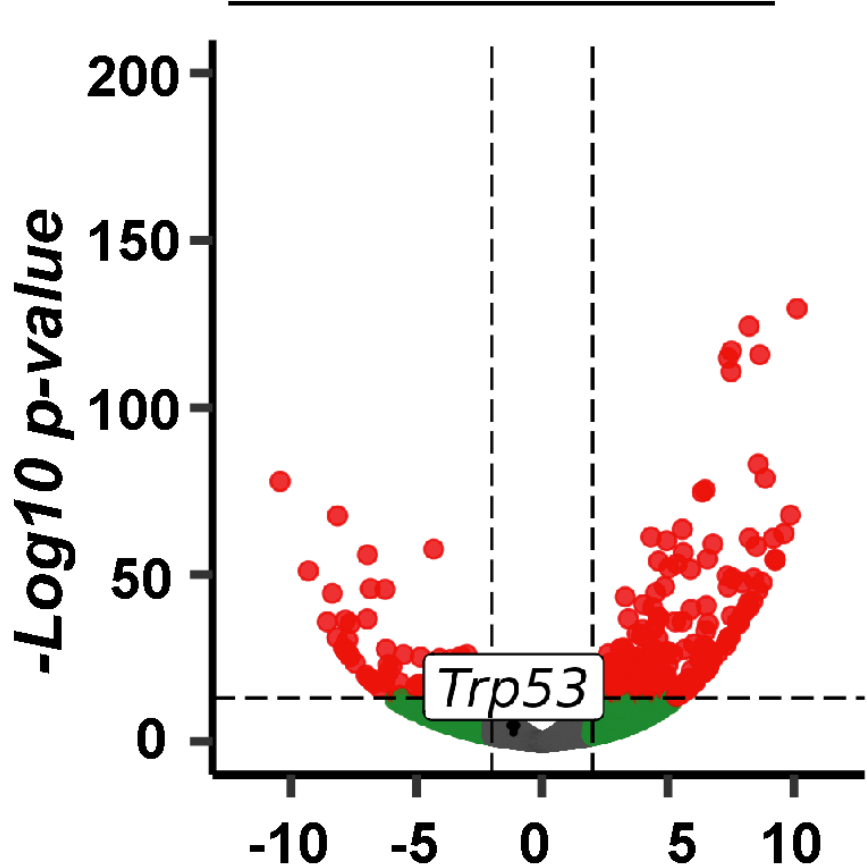
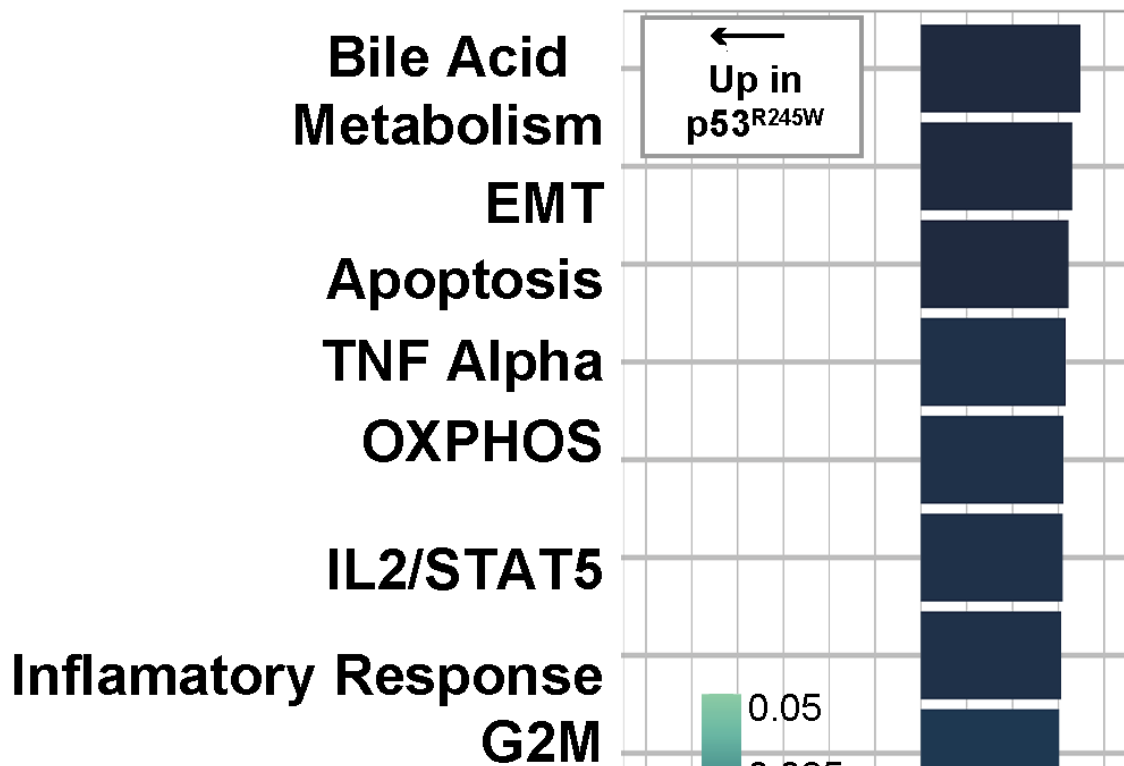


Figure 17: P53-R172H and p53-R245W cell lines have altered transcriptome compared to p53-null cell lines. (A) Volcano plot showing up- and down- regulated genes compared to p53-null cells. (B) Pathway analysis of Differentially expressed genes of p53-R172H and p53-R245W cell lines compared to p53-null cell lines.

A**p53^{R245W/R245W} vs p53^{R172H/R172H}****B****Log2 Fold Change**

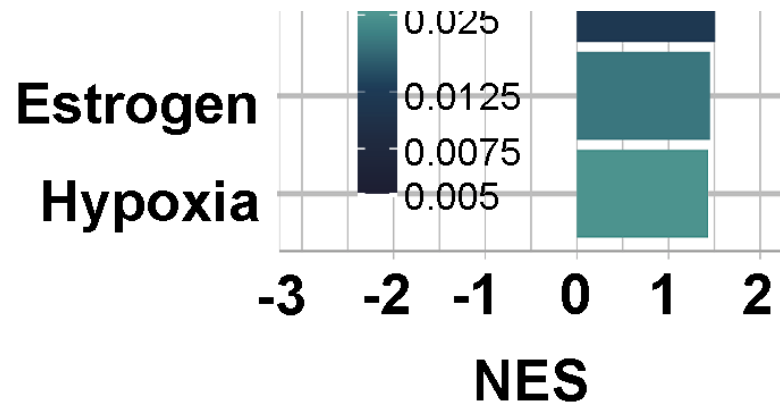


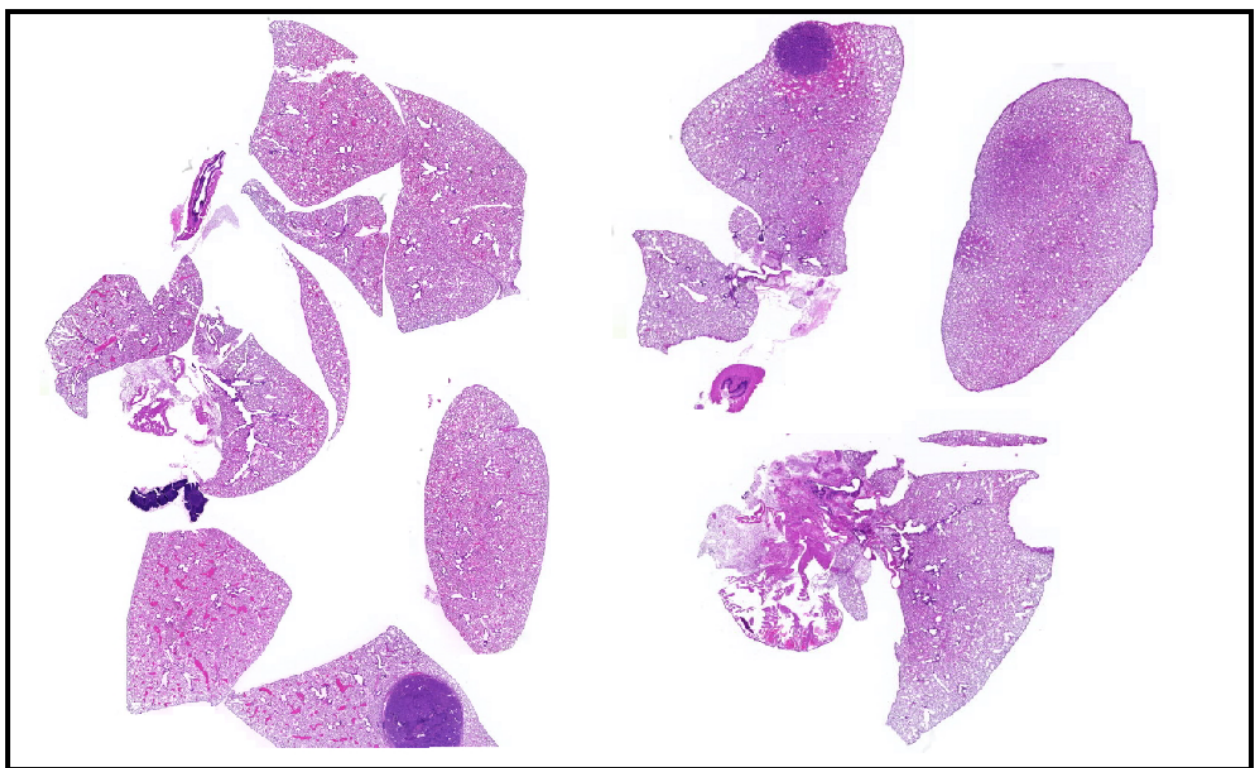
Figure 18: Comparison of differentially-expressed genes between p53-R172H and p53-R245W cell lines. (A) Volcano plot of differentially expressed gene between p53-R172H and p53-R245W tumor-derived cell lines. (B) Pathway analysis of differentially expressed genes between p53-R172H and p53-R245W tumor-derived cell lines.

Figure 19: PDGFRB is upregulated and expressed in mutant-p53 tumors and cell lines. (A) Log-normalized counts of Pdgfrb RNA reads from p53-mutant cell lines. (B) Western blot analysis of basal and PDGF-BB stimulated cells. (C) IHC of PDGFRB in p53-mutant GEMM tumors.

Figure 19: PDGFRB is upregulated and expressed in mutant-p53 tumors and cell lines. (A) Log-normalized counts of *Pdgfrb* RNA reads from p53-mutant cell lines. (B) Western blot analysis of basal and PDGF-BB stimulated cells. (C) IHC of PDGFRB in p53-mutant GEMM tumors.

A

Braf^{+/+} ; Trp53^{R172H/R172H}



B

Braf^{+/+} ; Trp53^{fl/fl}

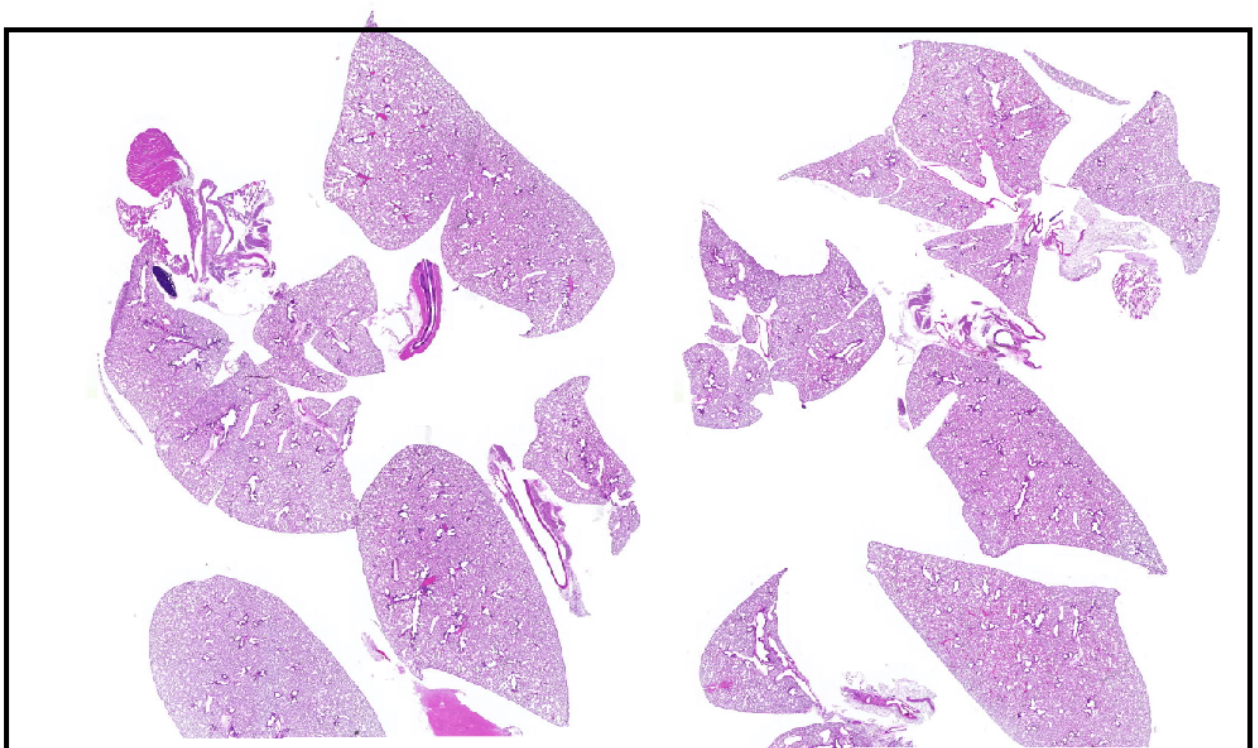


Figure 20: PDGF-stimulation or PDGF patway blockade has no effect on cell proliferation. (A) Growth response to PDGF stimulation. (B) Growth response to Imatinib treatment.

Chapter Four

##Abstract malignant lung adenocarcinomas tumors are initiated from a single neoplastic cell. Yet, there is substantial cellular heterogeneity in a fully malignant tumor. Since human lung adenocarcinomas are heavily mutated, it is difficult to know whether underlying cellular heterogeneity is a consequence of the underlying mutational heterogeneity. Mouse models of lung adenocarcinoma accurately represent genomic and histological aspects of human lung adenocarcinoma. However, genetically-engineered mouse model tumor heterogeneity remains poorly understood. Here, we profile the single cell transcriptomes of genetically-engineered mouse models that are driven by the *BRAF^{V600E}* oncoprotein with 5 separate cooperating genetic events (*Pi3k^{H1047R}*, *CdkN2a^{f1/f1}*, *Trp53^{f1/f1}*, *Trp53^{R172H/R162H}*, and *Trp53^{R245W/R245W}*). There are stereotypically distinct clusters of cells throughout all genotypes of *BRAF^{V600E}*-driven tumors. We also identified known tumor cell identities in both human and mouse lung adenocarcinoma.

Introduction

The mammalian lung is a vast and intricate labyrinth that is responsible for gas exchange between air and body through a meshwork of endothelial cells. The lung organ system begins with a single airway that branches in a geometric and stereotypic pattern(Metzger et al., 2008). These many thousands of branches lead to the business end of gas exchange, the lung alveoli. The alveoli are comprised of two specialized cell types: Alveolar Type 1 (AT1) cells comprise most of the surface area of the lung and are thin squamous cells that mediate oxygen exchange across cell membranes. Alveolar Type 2 (AT2) cells, however, are cuboidal that secrete surfactant proteins that aid in preventing the delicate alveoli from collapsing.

Pathology of the alveoli is a steadfast threat to an organism's life due to the importance of gas exchange. Diseases such as lung adenocarcinoma, pulmonary fibrosis, and emphysema are the most common dysfunctions of the distal lung epithelium(Desai et al., 2014). Lung adenocarcinoma is also the most common type of human lung cancers in both men and women. A large body of literature has suggested that lung adenocarcinomas arise in the alveolar and are descendants of AT2 cells (Ferone et al., 2020; Jackson et al., 2001). Therefore, there is great interest in understanding how AT2 cells develop, maintain cell identity, as well as how they renew themselves during aging or in response to injury (2020; Alanis et al., 2014, 2014; Treutlein et al., 2014).

Recent advances in profiling of single-cell transcriptomes during homeostasis conditions and disease states have greatly improved our understanding of AT2 cell, and lung adenocarcinoma cell biology(LaFave et al., 2020; Marjanovic et al., 2020; Treutlein et al., 2014).Single-cell profiling of human lung cancers have revealed the drastic intra-tumoral heterogeneity that exists in a single tumor(Gerlinger et al., 2012; Xue et al., 2016). Since transcriptional heterogeneity can be considered a source of therapeutic resistance, there is great interest in understanding the evolution of lung adenocarcinoma and how it changes with response to pathway targeted therapy(Maynard et al., 2020).Since GEMMs are an important pre-clinical model for testing out potential therapeutic strategies for lung adenocarcinoma, it is important to understand single-cell heterogeneity in popular GEMM models.

Although GEMMs are a great way to model human lung adenocarcinomas, there are several drawbacks worth mentioning. Human lung adenocarcinomas typically develop from a single tumor-initiating cell and is a consequence of an accumulation of mutations in certain cell types and a weakened immune system across a lifetime. In GEMMs, tumorigenesis is often initiated in many cells at once and is the consequence of an engineered mutation. In contrast to human lung adenocarcinomas that have a high mutational burden, murine lung adenocarcinomas do not have a

high mutational burden and instead display high copy number variation(Westcott et al., 2014). Therefore, we set out to profile the single-cell transcriptomes of various BRAF^{V600E}-driven lung cancer GEMMs in order to determine genotype-specific heterogeneity and response to pathway-targeted therapy.

Results

ScRNA Sequencing of benign and malignant BRAF^{V600E} mouse lung tumors We profiled the single-cell transcriptome of six different BRAF^{V600E} mouse lung tumor models including BRAF^{V600E} on its own, BRAF^{V600E} plus either mutant *Pik3ca*^{H1047R/+}, *Cdkn2a*^{f/f}, *Trp53*^{f/f}, *Trp53*^{R172H/R172H}, or *Trp53*^{R245W/R245W}. We also treated each genotype with dabrafenib (100mg/kg) plus trametinib(2mg/kg) once daily for four days prior to tissue harvesting and 10X genomics library preparation. Importantly, we kept dabrafenib and trametinib throughout all stages of processing to prevent altered transcription in response to drug cessation. Using fluorescently-activated cell sorting (FACS), we purified tumor cells based on Td-Tomato expression 8 weeks after tumor initiation using adenoviral Spc-Cre. In sum, we were able to identify 71,110 cells that passed standard 10X genomics and Seuratbased quality control and filtering (Figure 4.1).

Although we sought to purify tumor cells based on TdTomato-positivity, we did unintentionally collect and analyze stromal cells, such as immune, mesenchymal, and endothelial cell, which are also present in the murine distal lung epithelium (Figure 4.2). Therefore, we removed these cells based on their expression of *Ptprc*(immune), *Pecam1* (endothelial), and *Pdgfra* (mesenchymal) for downstream analysis (Figure 4.3).

Reclustering of only tumor cells(based on their expression of TdTomato and AT2 identity) revealed significant tumor cell heterogeneity(Figure 4.4). Next we sought to determine cluster specific markers using DESEQ2 analysis (Figure 4.5). strong markers of each cluster include

We next sought to determine the effect of dabrafenib plus trametinib on transcriptional heterogeneity (Figure 4.6). Clearly, there is a MAPK specific gene expression program that is decreased in most if not all clusters and genotypes (Figre 4.7).

We were also interested in genotype-specific differences. therefore, we performed DESEQ2 between vehicle-treated mice of each genotype (Figure 4.8).

table of genotypes/treatment/and cells collected Pre-sort pre-batch correction batch correction cell types tumor cells only Feature plots of AT2 markers feature plots of AT1 markers Feature plots of P53 target genes heatmap of P53 target genes heatmap/violin plot of MAPK score

Discussion

Here, we used scRNA-seq to determine transcriptional heterogenity of BRAF^{V600E} driven lung cancer and how it changes in response to concommittant genetic alterations. We modeled lung adenocarcinoma that mimick human oncogenic transformation where MAPK pathway activation drives tumor formation, and inactivation of the p53 pathway allows for tumor progression(Campbell et al., 2016). We also determined how cell state changes in response to MAPK blockade, a common therapeutic option for patients with BRAF^{V600E}-mutant lung adenocarcinomas.

We determined that there is transcriptional heterogeneity and that it is reproducible across genotypes. We also confirmed previousl published cell states in similar models, such as KRAS G12D(Marjanovic et al., 2020). Unfortunely, we did not observe robust dedifferentiation signals such as loss of NKX2-1 and

adoption of a gastric cell state. However, we did collect and identify highly-pastis cell states, such as ITGA2+ and CLDN4+ cells. Our failure to collect and analyze pre-metastatic cell states such as EMT-like cells, is likely because we collected cells only after 8-weeks, which is likely not long enough for these cell states to evolve during lung tumorigenesis. Therefore, adoption of highly malignant cell states, although might require certain genomic alterations like p53 loss, is an outcome of tumor evolution. One hypothesis is that p53 loss causes genomic instability and thus copy number gain, but similar studies failed to detect robust changes in copy number variation(Marjanovic et al., 2020).

We also failed to detect robust differences between genotypes that are known to cause malignant transformation, such as p53 loss, p53 missense mutation, CDKN2A loss, or mutant PI3K^{H1047R} (Dankort et al., 2007; Trejo et al., 2013; van Veen et al., 2019). This is likely a result from a combination of factors, such as time of collection and library capture/sequencing depth. One straightforward hypothesis is that we did not allow for full transcriptomic changes to take effect that is imparted by tumor suppressor or oncogene mutation. Therefore, a timecourse analysis that spans early timepoints (4-6 weeks) and late timepoints (>16 weeks) will likely reveal more informative transcriptional heterogeneity. Indeed, our previous studies have indicated that there is no tumor growth advantage at early time points as late as 6 weeks post tumor initiation. One other possibility that could explain our lack of genotype-specific gene expression programs is that we did not capture enough unique transcripts in order to reveal the full extent of tumor heterogeneity. This is likely because there are known limitations to 10X genomic library preparation. Specifically, there is gene-dropout effects where only the most highly expressed genes are captured by the droplet-based technique. Indeed, 10X genomics technologies have been shown not to detect genes with lower expression(Wang et al., 2021). However, 10X genomics allows you to profile more cells than similar scRNA-seq techniques. Genes with lower expression can be just as important as highly expressed genes for a cell's identity. Actually, transcription factors that determine cell states have been shown to have modest expression[Wang et al. (2021); ([pmid?](#));30205475]. Therefore, we hypothesis that using plate-based scRNA-seq approaches, such as Smart-Seq2, would allow for more gene capture. Thus, Smart-seq2 will enable us to more precisely determine transcriptional heterogeneity and genotype-specific gene expression changes.

Although our analysis was lacking with regards to transcriptional differences between genotypes, we did detect a robust transcriptional change with MAPK blockade(figure4.). Since human lung cancer patients treated with MAPK-pathway inhibitors exhibit a heterogeneity in response, one might hypothesize that each cell responds differently to MAPK-pathway inhibition. In our analysis, we determine that there is generally a homogenous response to MAPK inhibition. One straightforward explanation is that our short term treatment (four days) did not allow for cells to respond to pathway blockade. Perhaps there is an initial response to MAPK-pathway inhibition that is dampened over time that is dependant on transcriptional heterogeneity.

In conclusion, we used 10X genomics to determine the transcriptional heterogeneity of BRAF^{V600E} lung tumors with a five separate cooperating alterations and treatments. We did identify transcriptional heterogeneity that is stereotypic and reproducible across genotypes. However, we failed to identify robust differences between genotypes. Our work highlights the importance of capturing full transcriptomes across multiple timepoints in order to accurately determine cell states throughout BRAF^{V600E}-driven lung tumor evolution.

talk about the batch effects talk about the times similarity to other studies with KRAS talk about homogenous response to D+T

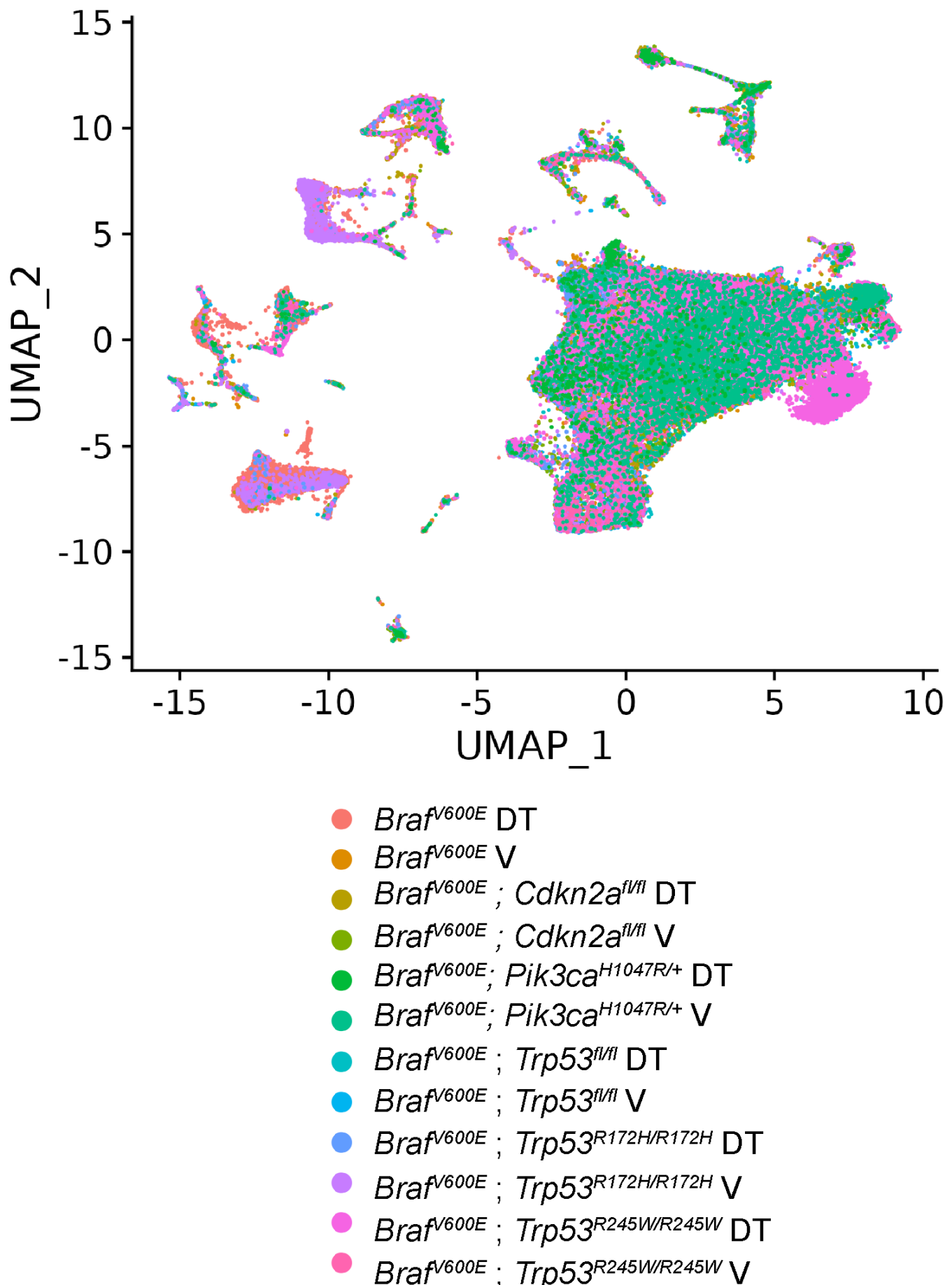


Figure 21: 71,110 cells collected from different genotypes.

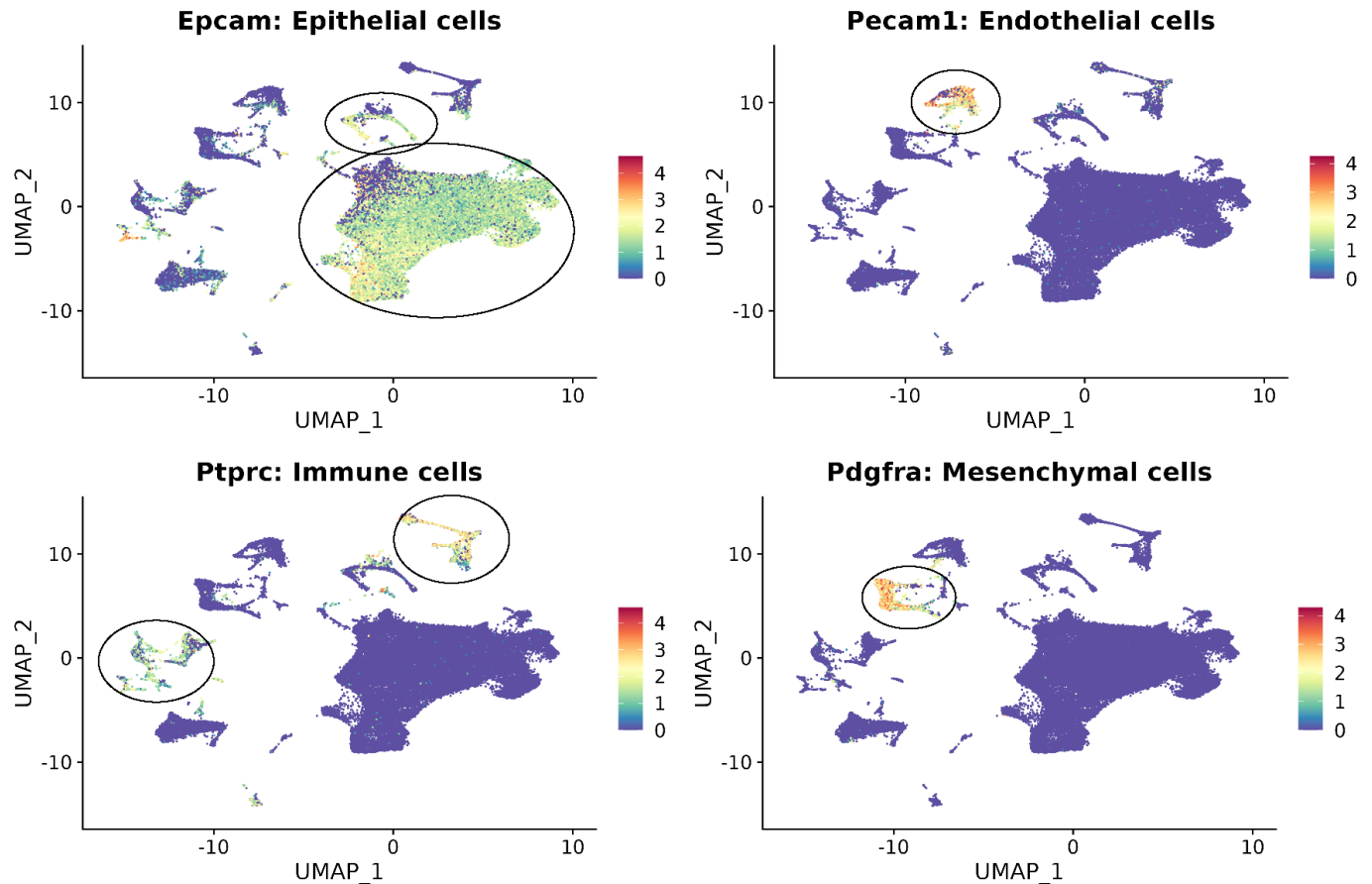


Figure 22: Single cells broadly cluster into epithelial-, immune-, endothelial-, or mesenchymal-like cells.

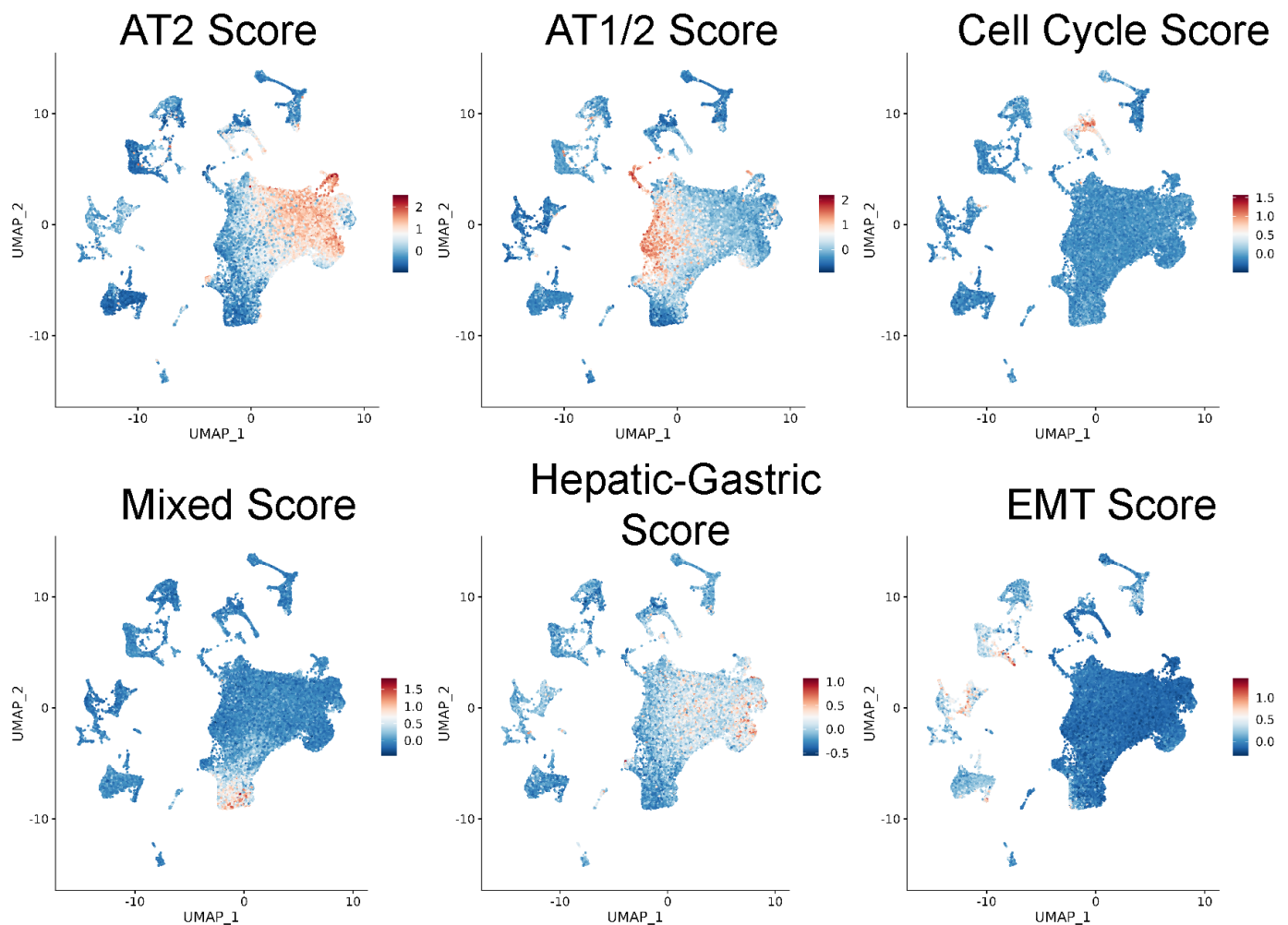


Figure 23: Murine lung adenocarcinoma transcriptional programs in BRAF^{V600E} tumorigenesis

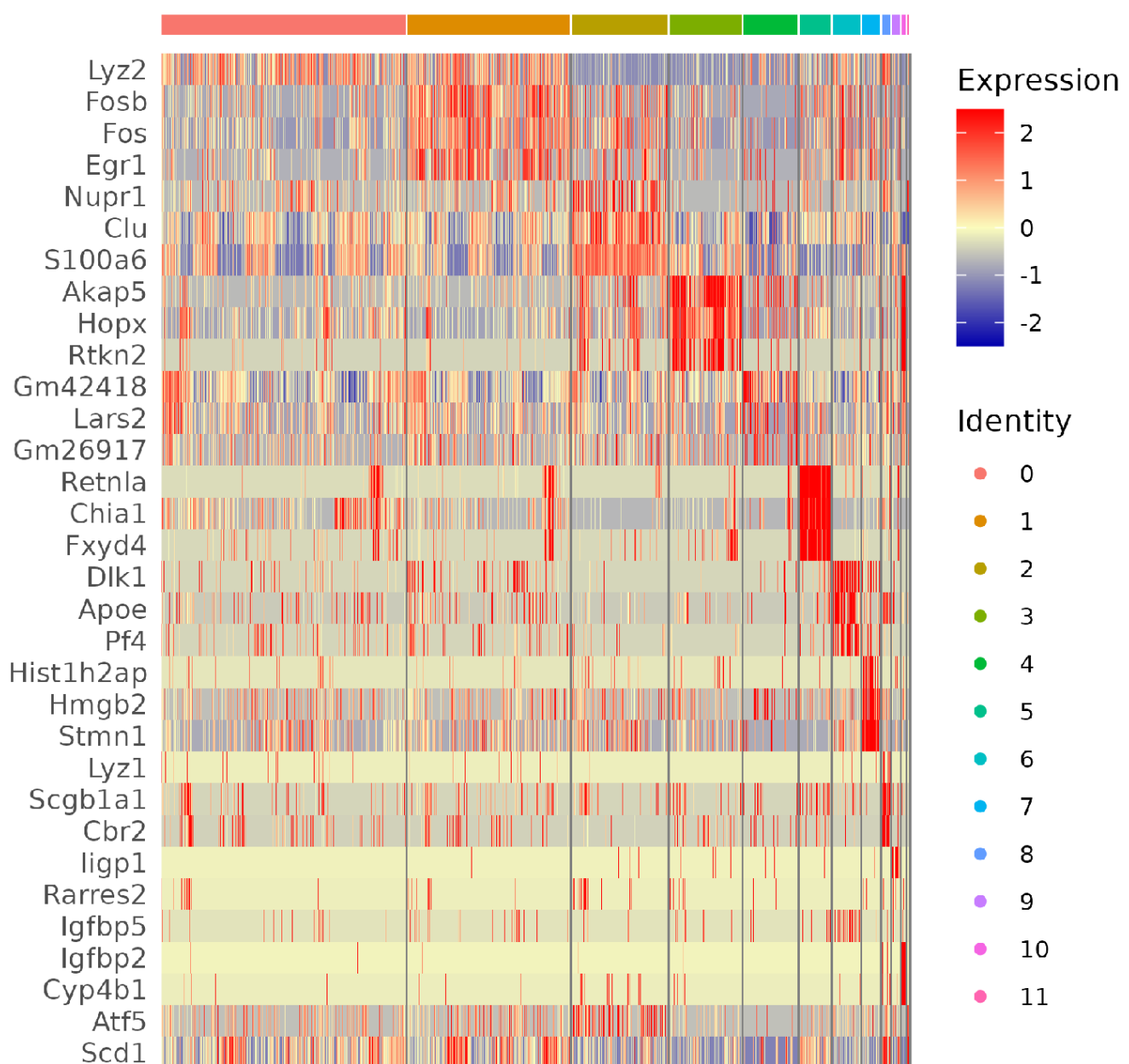
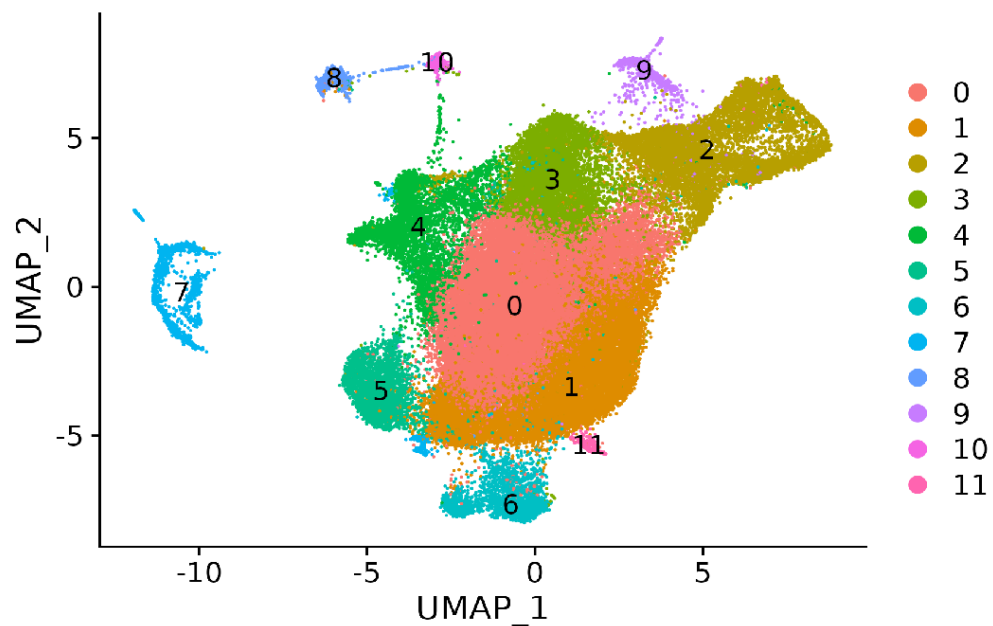
A

Figure 24: Transcriptional heterogeneity of tumor cells

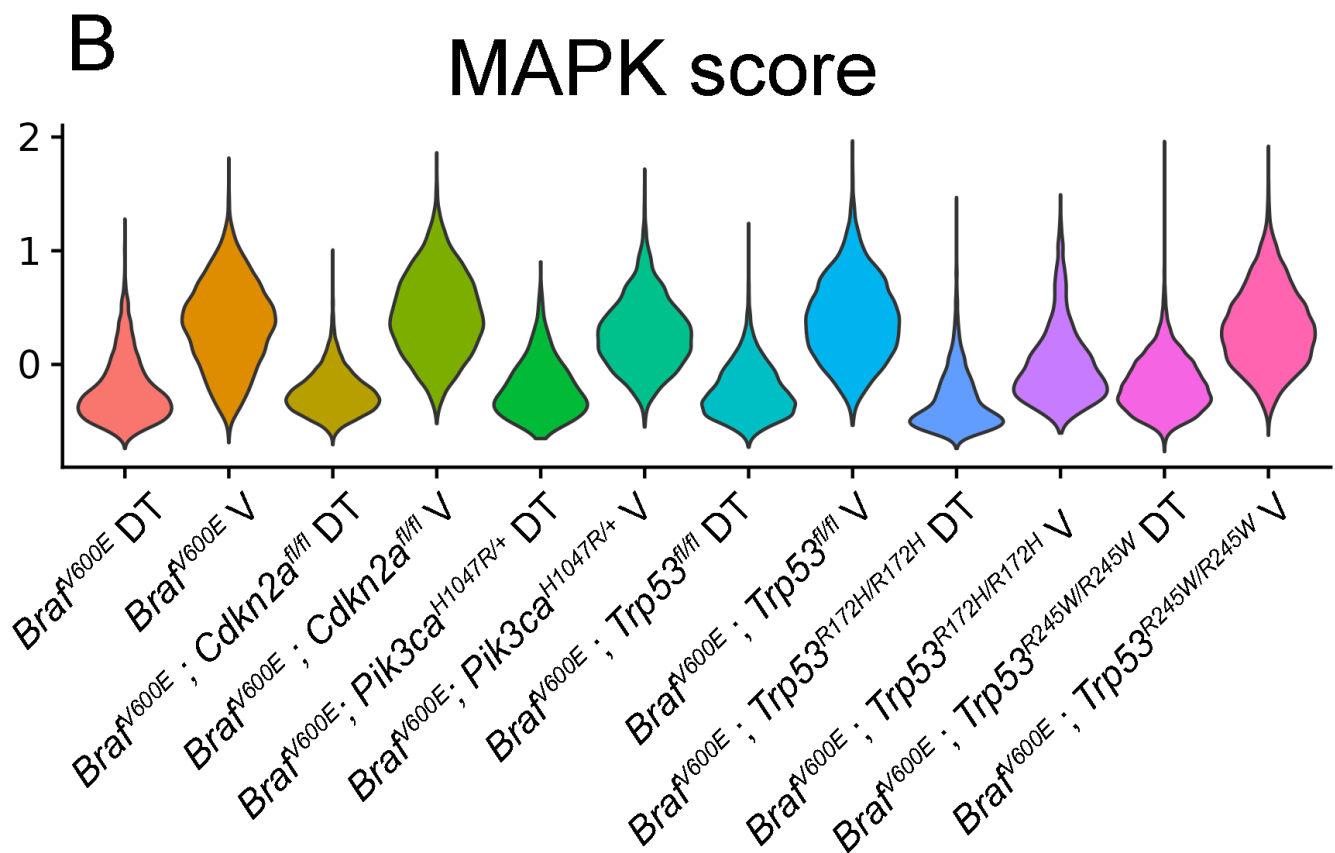
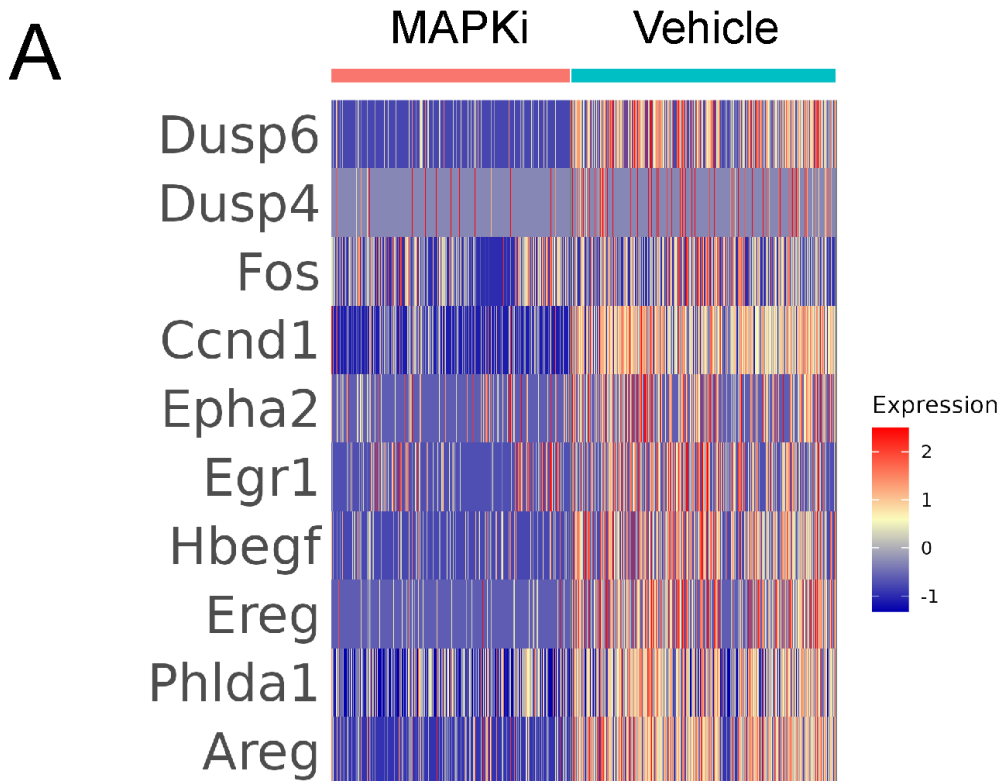
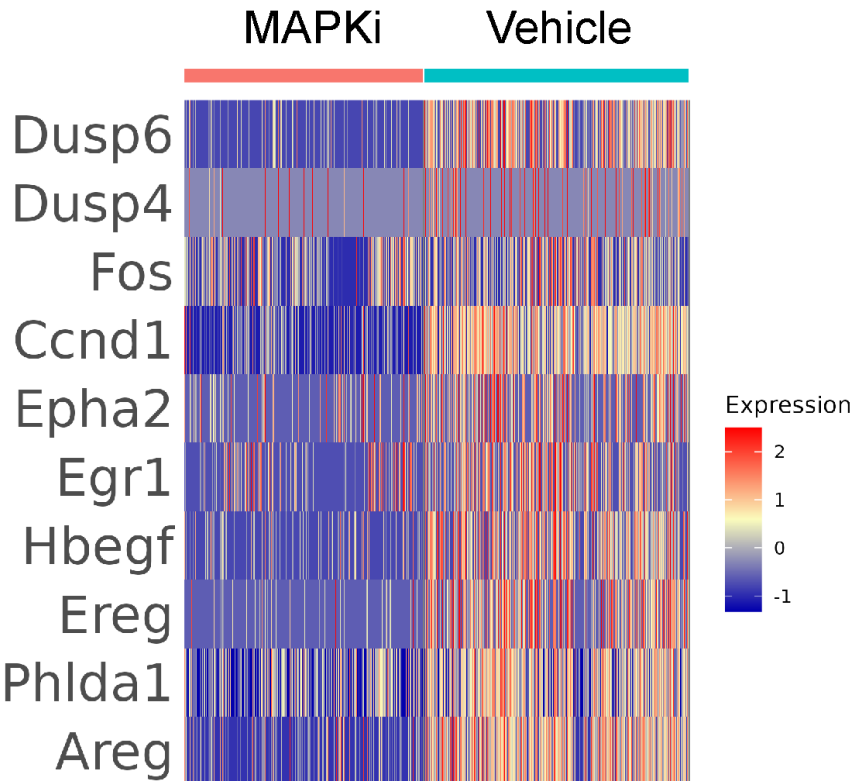


Figure 25: P53 transcriptional signature is identified in BRAF^{V600E}, p53-wildtype tumor cells

A**B**

MAPK score

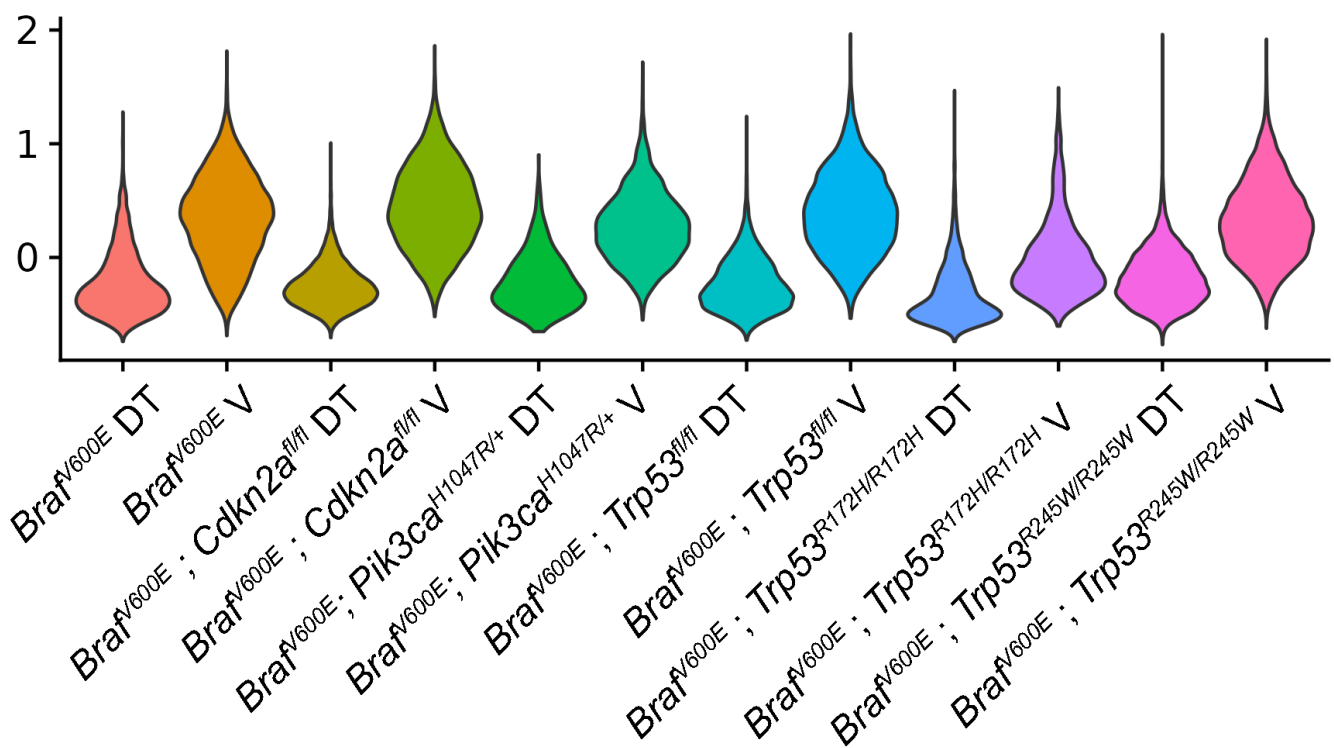


Figure 26: P53 transcriptional signature is identified in *BRAF^{V600E}*, p53-wildtype tumor cells

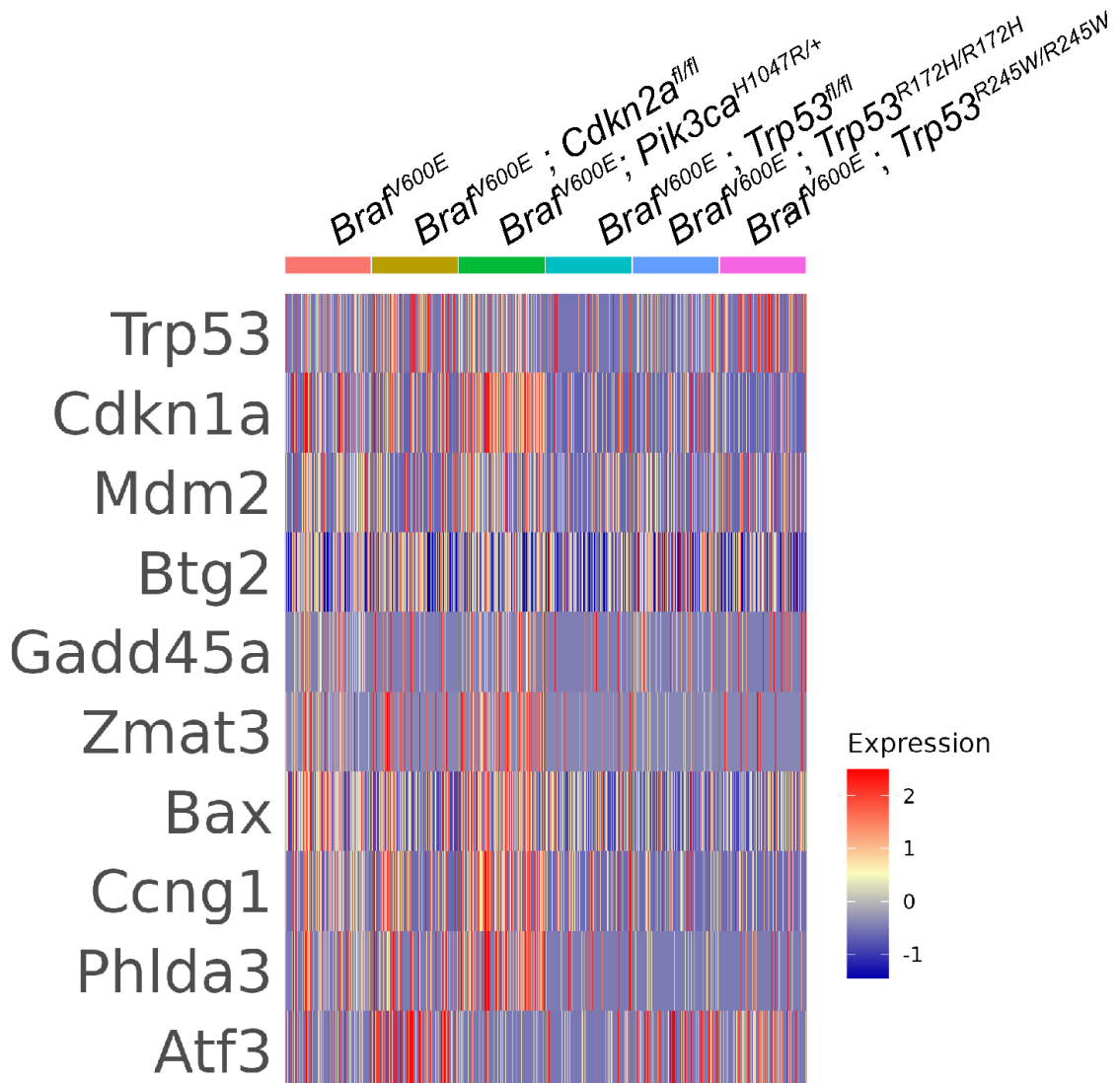
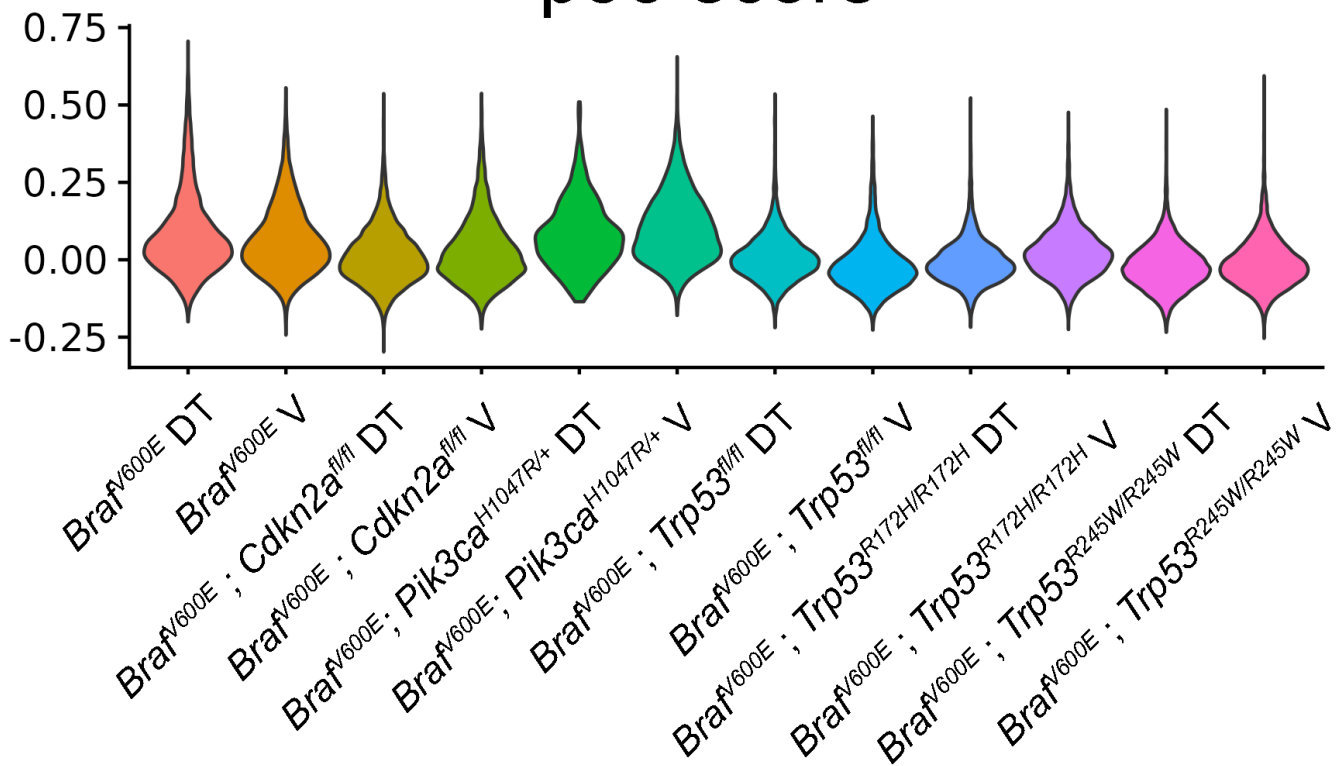
A**B****p53 score**

Figure 27: Genotype-specific gene expression changes.

Materials and Methods

Animal Husbandry All mice were housed in an environmentally controlled room, and all animal care and experimental procedures were approved by (and in accordance with) the Institutional Animal Care and Use Committee Office of the Huntsman Cancer Institute at the University of Utah. Genetically engineered mouse breeding and genotyping was conducted as previously described (van Veen et al., 2019). The BRAF^{CAT} and *Trp53*^{f/f} mice were previously described (Dankort et al., 2007; Jonkers et al., 2001; Shai et al., 2015; van Veen et al., 2019). All viruses were administered in a Biosafety Level 2+ room, as is regulated by the Institutional Biosafety Committee Guidelines. Adeno-SPC-CRE virus (University of Iowa) was delivered through nasal instillation. The *Trp53*^{wMR172H} and *Trp53*^{wMR245W} mice and respective genotyping protocols were gifted by Gigi Lozano and her laboratory and previously published (Kim et al., 2021; Zhang et al., 2018).

Tuba-Seq *Braf*^{CAT/+} and *H11b*^{LSL-CAS9/+} mice have been described(Chiou et al., 2015; van Veen et al., 2019) and maintained on a mixed background. approximately equal numbers of male and female mice were used in the study. Lung tumors were initiated by intratracheal intubation of lentiviral Cre vectors as previously described(DuPage et al., 2009). Barcoded lentiviral vectors were supplied by Monte Winslow as previously described(Winters et al., 2017). Briefly, Lentiviral vectors that contain an eight nucleotide sequence that defines the CRISPR-targeted followed by a 15 nucleotide sequence that is unique only to each initiated tumor. Lentivirus was generated in HEK 293T cells in DMEM supplemented with 10% FBS by standard lentiviral generation and titered as previously described(Winters et al., 2017). Genomic DNA was isolated from bulk lung tissue after weighing. Briefly, benchmark control cell lines were spiked-in to enable calculation of the absolute number of tumor cells from the number of barcode reads. After tissue homogenization and 12-hour proteinase-K digestion, genomic DNA isolation was extracted using standard Phenol-Chloroform and precipitated by ethanol. Sequencing libraries were prepared by PCR amplifying the sgRNA-BC sequencing in eight reactions in parallel with Q5 master mix and purified with AMPure beads and quantified with the Qubit instrument. Computational analysis was performed in house by the Monte Winslow laboratory.

Cell Line Generation Tumor bearing mice were euthanized and lungs dissected. Lungs were then minced and incubated in digestive enzymes Collagenase I, dispase, elastase, for 30 minutes at 37 degrees. Single-cell suspensions were then resuspended in complete DMEM medium and filtered through a 100 micron filter and centrifuged at 300 x g for 5 minutes. Single-cell suspensions were then incubated in red blood cell lysis buffer for 5 minutes at room temperature, then resuspended in complete RPMI and placed in a tissue culture incubator. Cells were then passaged once a week with differential trypsinization to purify tumor cells from viable stromal cells. p53 status and BRAF^{V600E} status (tdTomato+) were assessed by western blot and flow cytometry.

Tissue Culture Murine tumor derived cell lines and HCC-3564 cells were routinely cultured in DMEM supplemented with 10% heat-inactivated FBS and 1% penicillin/streptomycin. Cells were routinely tested for mycoplasma.

Immunoblotting Cells destined for western blot were scraped in ice cold PBS then centrifuged to pellet cells. After supernatant aspiration, cells were lysed with RIPA buffer supplemented with a protease/phosphatase inhibitor cocktail (Halt). Cell lysate protein concentration was determined with BCA assay. 40 microgram/lane was mixed with 4X SDS buffer and 10X sample buffer. Samples were run through a 4-12% Bis-tris gel at 200V for approximately 75 minutes then transferred to a PVDF membrane using the iBlot2 transfer apparatus. Membranes were then incubated in Odyssey blocking buffer for 30 minutes before being incubated overnight with primary antibody. Membranes were then washed 3X with TBS-T and incubated in secondary antibody for 2 hours before being washed 3X and imaged on the Licor Odyssey clX.

RNA Sequencing Cells lines were routinely cultured for approximately 10 passages before RNA purification using Qiagen Mini kit from one million cells. Libraries were generated with Illumina TruSeq Stranded mRNA Library Prep with UDI kit and sequenced on a Nova-Seq targeting 25 million reads per sample. fastq files were processed on Galaxy.org using FastQC, HISAT2, MultiQC, FeatureCounts, and DESeq2.

Immunohistochemistry and Immunofluorescence Immunohistochemistry was performed as previously described, with the rabbit primary antibody against P53, SPC, phospho-ERK1/2, NKX2-1, HMGA2, p19/ARF (Mollaoglu et al., 2017; van Veen et al., 2019).

Slide Imaging and Quantification Hematoxylin and eosin (H&E) and IHC stained slides 1 were scanned automatically by the 3D Histech Pannoramic MIDI scanner (Thermo Fisher). Slides were imaged and analyzed using Caseviewer Software or QuantCenter analytical center provided on the 3D Histech Slide Scanner at the indicated magnification. Tumor burden was manually calculated on each lung lobe and total tumor area was compared to total lung area. Tumor diameters were measured using QuantCenter software from 3D Histech.

Single Cell RNA-Seq analysis Mice were initiated as described previously using ad5-Spc-Cre. Four days prior to tissue harvesting, mice were either treated with corn oil control or corn oil with dabrafenib (150mg/kg) and trametenib(2mg/kg) once daily and again two hours prior to tissue harvest. single cell suspensions were made using enzymatic digestion, as previously detailed in cell line generation. Single-cell suspensions were then sorted with the BD ARIA II flow cytometer. 10X genomics library prep and sequencing was performed by HCL high throughput genomics core. Seurat objects where then created for downstream differential gene expression analysis and visualization with ggplots2.

Table 1: Critical reagents

Reagents List	Reagent Type	Name	Source	Catalog number
antibody	beta-Actin	CST	3700T	wb: 1:10,000
antibody	Phospho ERK1/2	CST	4370	wb:1:2000, IHC:1:1000
antibody	ERK1/2	CST	4696	wb: 1:2000
antibody	P53	Leica	cm-5	wb: 1:2000, IHC: 1:1000, IF:1:250
antibody	NKX2-1	Abcam	ab76013	IHC: 1:1000
antibody	Pro-SPC	Millipore	Cat# AB3786	IHC: 1:2000
antibody	HMGA2	CST	8179S	IHC 1:2000
antibody	p19/ARF	Chuck Sherr	(Bertwistle et al., 2004)	
Cell Line	293T	ATCC	CRL-3216	
Cell Line	HCC-364	Bivona lab	(Lin et al., 2014)	
Recombinant DNA	Pmd2.g	Addgene	#12259	
Recombinant DNA	pspax2	Addgene	#12260	
Recombinant DNA	P53-R175H	Addgene	#22936	
Chemical Compound	dabrafenib	Selleck	S2807	

Reagents List	Reagent Type	Name	Source	Catalog number
Chemical Compound	PLX4720	Plexxikon/ Research Diets		
Chemical Compound	trametinib	Selleck	S2673	
Other	Collagenase type 1	Thermo Fischer Scientific	17100017	
Other	Dispase	Corning	354235	
other	Elastase	Worthington Biochem	LS006365	
other	ad5spc-Cre	university of Iowa	VVC-Berns1168	

Chapter Six: Future Perspectives

This work's general purpose was to understand better how $BRAF^{V600E}$ tumors form and progress to lung adenocarcinoma. Our strategy was (1) to determine additional tumor suppressor genes in $BRAF^{V600E}$ lung tumorigenesis, (2) to evaluate the effects of different mutant-p53 alleles, and (3) to profile the tumor cell heterogeneity using scRNA-seq. To achieve these goals, we heavily relied on genetically engineered mouse models of B_BRAF^{V600E} lung tumors to characterize and compare the contribution of candidate-based tumor suppressors. This chapter will examine the exciting possibilities that arose due to our work.

Every year in the United States, approximately 4000 patients succumb to $BRAF$ -mutant lung cancer. Unlike melanoma or thyroid cancer, $BRAF^{V600E}$ is not the most common $BRAF$ alteration. Less potent MAPK activating mutations such as $BRAF^G469A$ are more common(Bracht et al., 2019; [omid:30770389?](#)). It is likely that less potent MAPK activators serve as support signaling molecules for oncogenes like $KRAS(G12D)$, which fine-tune pathway activation. However, little is known about how vital non-V600E $BRAF$ mutations are in the evolution of lung adenocarcinoma. Therefore, preclinical models such as genetically engineered mice will be vital to determine the role of non- $BRAF^{V600E}$ mutations in lung adenocarcinoma. Previous work from our lab has established that $BRAF^{V600E}$ plays a substantial role in the early formation of lung benign lung tumors but fails to form lung adenocarcinoma. Rather, benign to malignant lung cancer progression depends on additional genetic alterations(Trejo et al., 2013).

The mechanisms by which the mutationally activated MAPK-pathway promotes lung tumorigenesis remain incompletely understood. However, recent work suggests that the levels of MAPK-pathway activation correlate with tumor initiation and progression(Cicchini et al., 2017; Feldser et al., 2010). Therefore, there must be secondary events after mutational activation of KRAS or BRAF that alter the tone of the MAPK pathway. Recent work has highlighted potential mechanisms tumor cells adapt to increase pathway activation. Amplification of the KRAS locus occurs during early murine tumorigenesis initiated by KRAS-G12D(To et al., 2011). During late tumorigenesis, MAPK-pathway hyperactivation can occur through loss of DUSP6 regulation by NKX2-1(Ingram et al., 2021). However, NKX2-1 loss is also known to affect tumor cell identity and is associated with dedifferentiation(Mollaoglu et al., 2018; Zewdu et al., 2021); Snyder et al. (2013)]. For increased MAPK pathway activation to drive tumor progression, p53 function needs to be inhibited. Therefore, there are two main barriers to tumor progression in lung adenocarcinoma: (1) MAPK pathway activation and (2) subversion of p53-mediated tumor suppression.

We initially hypothesized that the barrier to tumor progression was the initiation of oncogene-induced senescence. Previous studies have linked oncogenic MAPK signaling to p53-mediated tumor suppression(Woods et al., 1997; Zhu et al., 1998). One hypothesis is that MAPK pathway activation crescendos during tumorigenesis until p19/ARF expression is induced, which triggers p53 activation and tumor suppressor. However, it is unclear if p53 needs to be silenced, as work from our lab and others have provided evidence that WNT signaling is also a barrier(Juan et al., 2014; Tammela et al., 2017). Whether increased WNT-signaling can overcome p53-induction remains to be understood. However, it is clear that p53 suppression and MAPK activation are both frequent events in human cancers and cooperate to form malignant lung adenocarcinoma in mice. Therefore, more research is needed to determine the precise mechanisms governing p53 tumor suppression and tumor progression.

The diversity of p53 mutations in human cancers is unlike other tumor suppressor genes. Whereas mutations in tumor suppressors APC, RB, and BRCA1, are inactivated thru aberrant stop-codons or gene deletion events, mutations in p53 are often missense mutations in the DNA-binding domain(Hainaut and Pfeifer, 2016). Although mutational inactivation of p53 is important for $BRAF^{V600E}$

driven lung tumor progression, common p53 alterations have not been tested for cooperating with BRAF^{V600E}. Instead, null alleles of p53 have been studied. This is undoubtedly due to the poor design of hotspot mutants of p53, such as p53-R175H, p53-R273H, and p53-R248W. These alleles of mutant p53 are designed to be null for p53 prior to recombination, and mice harboring these alleles are prone to developing cancers that have nothing to do with MAPK activation in the lung epithelium. Therefore, we utilized the recently generation wild-type-to-mutant p53 alleles *TRp53*^{R172H} and *Trp53*^{R245W} (*Tp53*^{R175H} and *Tp53*^{R248W} in humans)(Zhang et al., 2018). Part of this work has been dedicated to characterizing the effect of *Trp53*^{f/f}, *Trp53*^{R172H}, and *Trp53*^{R245W} on *Braf*^{V600E} lung tumorigenesis. Strong evidence and careful characterization of mutant-p53 alleles have determined that not all p53-mutations are created equal. We have shown that p53^{R172H} and p53^{R245W} differentially alter BRAF^{V600E} tumor progression. The mechanism by which p53-missense mutants abrogate tumor suppression is two-fold. Mutant-p53 inactivates wild-type p53 through acting as a dominant-negative(Chan et al., 2004). More controversial is the ability of mutant-p53 to exert gain-of-function effects. Our studies failed to determine a molecular explanation as to why P53^{R172H} and P53^{R245W} are more potent drivers of cancer growth and drug resistance. However, we did determine that cell lines derived from mouse lung tumors have altered transcriptional profiles. Recently, evidence has emerged suggesting aneuploidy is a common event in mutant-p53 cells(Redman-Rivera et al., 2021). Specifically, copy number alterations were more common in mutant-p53 cells than in p53-null cells in an isogenic study(Redman-Rivera et al., 2021). Therefore, the phenotypic diversity underlying many p53 gain-of-function studies could reflect inherent genomic instability imparted by p53-mutations. This idea could help explain our data, as loss of mutant-p53 had no apparent phenotype in mutant-p53 cells. Therefore, it is of great interest to determine the copy number variation in our various p53-mutant models.

This dissertation support previous studies that found gain-of-function effects imparted by mutant-p53(Pfister and Prives, 2017). However, the field of mutant-p53 gain-of-function is still murky with controversy. In the future, it is essential to resolve our understanding of mutant-p53 proteins and how they influence the properties of cancer cells. We believe the diversity of mutant-p53 mechanisms can be explained by complex interactions between tumor types and each mutant-p53. Therefore, careful analysis of each p53-mutant in each cancer type might be required. Therefore, we propose to combine Tuba-Seq with CRISPR/CAS9-mediated homology-directed repair to determine the fitness landscape of every p53 mutation possible, similar to what has been performed with *Kras* in lung cancer and more recently in colorectal cancer (Cai et al., 2021; Zafra et al., 2020). This will allow for the unbiased quantification of the growth advantage imparted by individual p53-mutants in a single animal.

Our studies also shed insight into p53 stability in BRAF^{V600E} lung cancer. We have observed that mutant-p53 accumulation correlates with p19/ARF, consistent with previously published research(Feldser et al., 2010; Junntila et al., 2010). Therefore, p19/ARF may play a dichotomous role in tumor development. In the context of wild-type p53, p19/ARF might play a tumor-suppressive role in response to overactive MAPK signaling. In mutant-p53 expressing tumors, p19/ARF can stabilize mutant-p53, potentiating cancer progression. In order to fully understand p53 tumor suppression and mutant-p53 dominant-negative effects, we propose to study p19/ARF-specific deletion in the context of wild-type and mutant p53 in BRAF^{V600E} tumorigenesis. Since the *CDKN2A* locus encodes p19/ARF, it is difficult to determine the effect of p19-specific roles in genetically-engineered mice. However, one could still test the role of p19/ARF in mutant-p53 stabilization *in vivo* and *in vitro* using p19/ARF specific RNA-interference such as siRNAs or short-hairpin RNAs.

Pathway Targeted Therapy

Resistance is inevitable to pathway-targeted therapy. Drug resistance is a consequence of complex tumor heterogeneity that drives tumor evolution. Interestingly, it has been suggested that the baseline mutation rate of tumor cells is high enough that in a sufficiently large tumor, all possible

single nucleotide variants have likely occurred(Ruan et al., 2020.; Tomlinson et al., 1996) Furthermore, the more prolonged the therapy, the more likely that resistant cells can propagate and fuel tumor growth after an initial tumor regression. In terms of MAPK-pathway inhibition in melanoma and lung cancer, many resistance mechanisms have been identified(Maynard et al., 2020; Rudin et al., 2013). Not surprisingly, many resistance mechanisms involved reactivation of the MAPK by mutational activation of upstream or downstream regulators(Holderfield et al., 2014; Lin et al., 2014; Rudin et al., 2013). Our work has provided evidence that p53-mutations can alter responses to pathway-targeted therapy. Our data indicate that *Trp53*-mutant cell lines have greater basal MAPK pathway activation. Moreover, the MAPK pathway is still active in response to MAPK targeted therapy. Therefore, we hypothesize that mutant-p53 can alter MAPK pathway activity to drive therapeutic resistance. However, we have yet to determine a mechanism to explain how mutant-p53 alters MAPK signaling. Thus, understanding the molecular mechanisms governing mutant-p53 cell biology and biochemistry is warranted.

Currently, p53 mutational status is not a contraindicator of MAPK targeted therapy in *BRAF^{V600E}* mutant lung cancer(Matsumura et al., 2018). Patients are often stratified into wild-type or mutant p53 during diagnosis. However, Our data suggest that there are substantial differences between p53-mutants regarding tumor progression and therapeutic resistance. There are complex gene-gene interactions at play that affect therapeutic resistance. Therefore, careful analysis of cooperating genomic events' contribution to pathway-targeted therapy needs to be tested in preclinical models.

Since reactivation of the MAPK is a common mechanism of dabrafenib plus trametinib resistance, further vertical inhibition of the pathway is considered an attractive therapeutic strategy. Previous research has shown that triple combination (BRAFi+MEKi+ERKi) imposes a more substantial threshold for pathway reactivation and subsequent resistance(Xue et al., 2017). Although more effective at tumor regression and evasion of resistance, vertical inhibition undoubtedly causes severe side effects that can not be tolerated. Therefore, more combination strategies are likely required. Fortunately, exciting therapeutic strategies are emerging from our lab that appears to be efficacious in MAPK-driven tumors, such as the combination of MAPK-pathway inhibitors and autophagy inhibition(Foth and McMahon, 2021; Kinsey et al., 2019; Truong et al., 2020).

In parallel to my efforts to characterize three p53-mutations in *BRAF^{V600E}* lung cancer, we determined the contribution of a broad range of tumor suppressor proteins to tumor progression. Our work has highlighted the role SETD2 plays in suppressing lung tumorigenesis. However, we currently do not understand the mechanism behind SETD2 tumor suppression. One hypothesis is that SETD2 prevents spurious gene transcription. Thus when SETD2 function is lost, tumor cells adopt multiple cell states due to altered gene expression. The more cell states a tumor cell can access increases the odds of adopting pro-growth characteristics best for the tumor environment. In order to test this hypothesis, we can profile the transcriptomes of single cells that have intact SETD2 and SETD2 deficient tumor cells. Importantly, we must consider alternative transcriptional start sites and alternative splicing. Since p53 dysregulation is at the heart of tumor progression, we would like to know if SETD2 affects p53-stability and target-gene expression. Alternatively, SETD2 loss might circumvent p53 activation through some undescribed mechanism.

Overall, lung adenocarcinoma is a disease that is characterized by a high mutational burden that results in mutational activation of the MAPK pathway and tumorigenesis. Large-scale genomic studies have provided great insight into the molecular underpinning of lung adenocarcinoma. However, due to the complex interactions between gene products, the future of personalized medicine still relies on rigorous testing in preclinical models like genetically-engineered mice. These efforts of this work will aid in the ultimate understanding of cooperating genetic events in lung adenocarcinoma. Ultimately, this dissertation has helped answer several questions regarding *BRAF^{V600E}* tumorigenesis and cooperating events for tumor progression. Additionally, this work has also generated new questions

that need to be answered to fully understand how BRAF^{V600E} and p53 mutations cooperate to form deadly lung cancers.

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