Genomic and experimental evidence that $\text{ALK}^{\text{ATI}}$ does not predict single agent sensitivity to ALK inhibitors

**Highlights**

- A method to test rare genomic findings for their relative conditional selection
- $\text{ALK}^{\text{ATI}}$ is not as mutually exclusive with BRAF or NRAS as they are with each other
- $\text{ALK}^{\text{ATI}}$ is not likely to be sufficient for cellular transformation or growth in vitro

Expressing activated oncogenic ALK in BRAFV600E melanoma cells is cytotoxic.
Genomic and experimental evidence that ALK\textsuperscript{ATI} does not predict single agent sensitivity to ALK inhibitors

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SUMMARY
Genomic data can facilitate personalized treatment decisions by enabling therapeutic hypotheses in individual patients. Mutual exclusivity has been an empirically useful signal for identifying activating mutations that respond to single agent targeted therapies. However, a low mutation frequency can underpower this signal for rare variants. We develop a resampling based method for the direct pairwise comparison of conditional selection between sets of gene pairs. We apply this method to a transcript variant of anaplastic lymphoma kinase (ALK) in melanoma, termed ALK\textsuperscript{ATI} that was suggested to predict sensitivity to ALK inhibitors and we find that it is not mutually exclusive with key melanoma oncogenes. Furthermore, we find that ALK\textsuperscript{ATI} is not likely to be sufficient for cellular transformation or growth, and it does not predict single agent therapeutic dependency. Our work strongly disfavors the role of ALK\textsuperscript{ATI} as a targetable oncogenic driver that might be sensitive to single agent ALK treatment.

INTRODUCTION
In cancers, clonal selection influences tumor progression and responses to therapy (Burrell et al., 2013), but in every patient, the process of clonal selection creates independent tumors with independent, parallel evolutionary trajectories. These parallel evolutionary paths can be conditioned upon the occurrence of a previous genetic event, i.e., they can be conditionally selected. Co-occurrence is when the first variant predicts the presence of a second (Mina et al., 2017; Whittaker et al., 2013). Whereas, mutual exclusivity occurs when the presence of the first predicts the absence of the second—and is the primary focus here. Mutual exclusivity is often viewed qualitatively or used to build large scale networks (Ciriello et al., 2012; Mina et al., 2017; Vandin et al., 2012; Zhao et al., 2012). Here we aim to create quantitative guideposts of conditional selection that will allow for direct comparisons of mutual exclusivity relative to known druggable oncogene pairs by controlling for cohort size. These guideposts can be used to triage the translational actionability of rare genomic findings.

A strong example of mutual exclusivity in cancer is seen in the evolutionarily ancient mitogen-activated protein kinase (MAPK) pathway that is present from yeast to metazoans (Widmann et al., 1999). In higher eukaryotes, the MAPK pathway is often canonically activated by receptor tyrosine kinases (RTK) that signal through MAPKs to achieve cellular growth and development (Lemmon and Schlessinger, 2010; Zhang and Liu, 2002). Because of their critical role in growth and division, most known RTK-MAPK mutational events that drive cancer growth are mutually exclusive across patients during parallel evolution. Moreover, many of the most impressive success stories for targeted cancer therapy in the past two decades have centered on this one pathway. Multiple examples of mutual exclusivity have been found between ALK-fusion, epidermal growth factor receptor (EGFR), Kristen rat sarcoma viral (KRAS), and Erb-b2 receptor tyrosine kinase 2 (ERBB2) genes in non-small cell lung cancer (NSCLC) (Takahashi et al., 2010; Torkelson et al., 2015; Varmus et al., 2016). This has led to inhibitors of ALK, EGFR, and recently KRAS\textsubscript{G12C}, offering high rates of single agent therapeutic responses (Bhullar et al., 2018; Fakhri et al., 2019; Muhsein et al., 2003; Shaw et al., 2012; Swaika et al., 2014). In thyroid cancer, BRAF mutations have been found to be mutually exclusive with RET fusions, and both have shown high single agent response rates in clinical trials (Liang et al., 2007; Schlumberger et al., 2014). Similarly, BRAF & NRAS are mutually exclusive in melanoma and single agent use of vemurafenib leads to clinical responses (Edlundh-Rose et al., 2006; Guan et al., 2015; Marzese et al., 2016).
sequencing of the melanoma PDXs expressing ALK ATI revealed that 4 out of 6 of these cell lines had trans-

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noma patients treated with ALK-inhibitors so far, one had a modest response that did not rise to the level

(BRAF and NRAS are with each other. We also show that the relatively rare prevalence of ALK ATI

forming mutations in well-established melanoma oncogenes. Furthermore, out of the two ALK ATI mela-

population. Using this method, we show that ALKATI is not as mutually exclusive with BRAF or NRAS as

sitivity of rare variants versus a positive control pair. Because we use positive controls that are genes in the

MAPK pathway with previously verified single agent responses to targeted therapeutics (Chapman et al.,

ings for potential single agent drug efficacy. If a newly discovered alteration is as mutually exclusive as a

therapeutically established mutually exclusive gene pair, it gives greater confidence in the potential for sin-

gle agent therapy. The decision tree in Figure S1 illustrates the advances in mutual exclusivity analysis that

are enabled by our method.

As a test case for a method that can quantitatively compare mutual exclusivity between gene pairs, we re-

examined the observation that alternative transcription initiation in ALK, termed ALKATI, is a therapeutically

actionable oncogenic target (Wiesner et al., 2015). This novel ALK transcript has a transcription initiation

site in intron 19 of ALK, just upstream of ALK’s kinase domain. Wiesner et al. posited that this transcript ex-

hibits a novel mechanism of oncogenic activation by overexpressing the kinase domain of ALK. Using

in vitro transformation assays and inhibitor treatment, they hypothesize that single agent ALK-inhibitor

therapy can treat the 2–11% (Busam et al., 2016; Wiesner et al., 2015) of melanoma patients expressing

ALKATI. However, Couts et al. reported crizotinib (an ALK inhibitor) sensitivity in melanoma patient derived

xenografts (PDXs) expressing EML4-ALK but not ALK AT (Couts et al., 2017). Exome sequencing of the melanoma PDXs expressing ALKATI revealed that 4 out of 6 of these cell lines had transforming mutations in well-established melanoma oncogenes. Furthermore, out of the two ALKATI melanoma patients treated with ALK-inhibitors so far, one had a modest response that did not rise to the level of an objective response (Wiesner et al., 2015) and the other did not respond (Couts et al., 2017). This contradictory evidence, and the small sample size of the Couts et al. study suggest that reanalysis of the original observation of ALK AT is important to understanding if further investigation is warranted.

Two genes are conditionally selected if their odds ratio differs significantly from 1. In this paper, we develop a simple and user-friendly resampling method to test the relative conditional selection across two gene pairs. Relative conditional selection tests whether a significant difference exists between the odds ratio of two gene pairs. This allows us to quantitatively compare the conditional selection of any gene of interest (GOI) such as ALK AT with positive control genes such as BRAF and NRAS in melanoma. In addition to quantifying levels of relative conditional selection across gene pairs, our approach also detects cases where the lack of conditional selection of a GOI with another gene may be confounded by its low prevalence in the population. Using this method, we show that ALK AT is not as mutually exclusive with BRAF or NRAS as BRAF and NRAS are with each other. We also show that the relatively rare prevalence of ALK AT (~2–10%) is not a major contributing factor behind its lack of mutual exclusivity with BRAF and NRAS. Moreover, by large scale repetition of experiments performed in the Wiesner paper (Wiesner et al., 2015), we uncovered kinase activating mutations in ALK AT that can be selected for during transformation assays. This suggests that it is not sufficient for growth factor independent transformation. We also find that ALK AT
cannot compensate for oncogenic signaling in melanoma cells, and that ALK kinase domain overexpression does not predict ALK inhibitor sensitivity in the cancer cell line encyclopedia (CCLE) (Barretina et al., 2012). Finally, activated ALK actually inhibits the growth of BRAFV600E melanoma cell lines. Given the stunning advances in melanoma therapy, and the rich clinical trial landscape, we suggest that relapse/refractory melanoma patients with ALKAT should not be given ALK inhibitors in an investigational or off-label capacity unless no other options remain.

RESULTS

Pairwise comparisons of conditional selection allow direct comparisons between frequent and rare events

Highly mutually exclusive gene pairs such as KRAS/EGFR and EGFR/EML4-ALK in lung cancer, and NRAS/BRaf in melanoma predict the success of ALK, EGFR and BRAF inhibition in the clinic (Gainor et al., 2013; Shaw et al., 2012). These mutually exclusive gene pairs are positive controls that can be compared with any gene/alteration of interest. A direct pairwise comparison would allow us to ask whether a “BRaf/NRas level” of mutual exclusivity exists between any new GOI (such as ALKAT) and BRaf or NRas (BRaf and NRas are an example of a highly mutually exclusive gene pair in melanoma, 20–24). However, the differences in sample size due to mutation frequency complicates our ability to directly compare the odds ratios between distinct gene pairs. If the GOI is much less abundant than the positive control, how often would the positive control have a similar odds ratio if it was only as abundant as the test gene of interest? Resampling will allow us to more directly compare the mutual exclusivities between pairs of genes with unequal abundances in cancer genomes (see STAR Methods and Data S1 for full details).

To directly illustrate this idea, we analyzed how sample size affects the odds ratios observed for a simulated mutually exclusive positive control gene pair. We created a mock clinical cohort with two abundant but mutually exclusive genes (odds ratio of 0.025), and then we performed resampling experiments at different subsample sizes. As expected, decreasing the abundance of one of the genes increases the range in the observed odds ratios (OR) at smaller sample sizes: OR of 0–0.7 for a subsample size of 5 and OR of 0–0.11 for a subsample size of 20 (Figure 1A). Therefore, the range of odds ratios that is observable in highly mutually exclusive genes becomes noisier as sample size decreases. This means that a signal of mutual exclusivity in a pair of genes can be masked if one of the genes is rare. To more directly compare gene pairs with different abundances, we propose adjusting the frequency of the more prevalent genes to a frequency that is matched by the gene of interest.

We refer to this as “pairwise comparisons of conditional selection”. We aim to directly compare two gene pairs while controlling for the differences in sample size. Given two highly prevalent, mutually exclusive positive control genes and a rare GOI with low frequency of mutations, our method adjusts the frequency of mutations of one of the positive control genes to match the frequency of mutations of the rare GOI. This effectively normalizes the prevalence of rare vs common genes. To quantify relative conditional selection, our method compares the odds ratios produced by the two positive control genes to the odds ratios produced by the GOI and one of the positive control genes (Figure 1B). This lets us test the null hypothesis that no difference exists between the odds ratios of the two gene pairs. The higher the difference between these two sets of odds ratios, the more certain one can be that the gene of interest is not as mutually exclusive with one of the positive control genes as the two positive control genes are with each other. Details of the pairwise comparison strategy are included in the STAR Methods, the pseudo-code (Data S1), and our GitHub repository.

To explore the sensitivity and specificity of our resampling approach we generated gene pairs (see STAR Methods, pseudo-code) with varying GOI frequencies, and odds ratios for the two gene pairs (Figure S2A). As expected, when we simulated two gene pairs with similar mutual exclusivities, no significant differences in their odds ratio distributions were detected (Figures 1C and 1D left panel). Furthermore, large differences in odds ratios required relatively few observations of a genetic event (Figures 1C and 1D right panel).

If the simulated gene pairs had a small difference in their odds ratios, whether or not a strong difference was detected depended on the abundance of the GOI in the cohort (Figures 1D middle panel and S2B). For example, in a cohort of 500 patients, if the two positive control genes had an odds ratio of 0.05 and GOI and PC1 had an odds ratio of 0.2, at least 28 GOI patients were needed before a high difference in odds ratios could be detected, i.e., this cohort requires at least 28 GOI patients before we can conclude that the difference in odds ratios between the two gene pairs is real.
Figure 1. Pairwise comparisons of conditional selection between a GOI gene pair and a positive control gene pair in simulated cohorts

(A) Distribution of odds ratio of two mutually exclusive genes when one of the genes is rare (left-side) and abundant (right-side).

(B) A single simulated patient cohort is scored by our pairwise comparisons approach. In this cohort, the odds ratio of PC1 with PC2 was 0.05 and the odds ratio of GOI with both PC1 and PC2 genes was >1. The frequency of GOI was set to 5%. The scoring threshold is the median of the PC1 and PC2 distribution. The score is the percentage of the GOI & PC1 or GOI & PC2 distribution that is greater than this threshold. Odds ratios displayed on the plot are for the comparison between the gene of interest and positive control (gene of interest gene pair).

(C–D) Heatmaps showing the dependence of a high score (blue) on the difference between the odds ratios of the two gene pairs and on the abundance of the GOI. Each tile represents a single simulated cohort. A cohort size of 500 simulated patients was used. A GOI abundance of 25 patients was used in (C).
Our method is sensitive enough to detect meaningful pairwise differences in mutual exclusivity in ~10/500 (2%) patients in a clinical cohort. Given that this sensitivity should be sufficient for the observed frequency of ALK<sup>ATI</sup> (observed at 2-10% frequency), we decided to re-examine the literature controversy surrounding this putatively oncogenic alteration.

**ALK<sup>ATI</sup> is not as mutually exclusive as other established therapeutic targets in melanoma**

The Couts paper contained a sample size of 6 ALK<sup>ATI</sup> patients, but it identified NRAS and BRAF mutations in patients that harbored the ALK<sup>ATI</sup> alteration (Couts et al., 2017). Thus, the lack of mutual exclusivity of ALK<sup>ATI</sup> with the transforming melanoma oncogenes BRAF and NRAS has been suggested, but not conclusively and quantitatively demonstrated with an appropriately powered analysis. Pairwise comparison of conditional selection clearly showed that ALK<sup>ATI</sup> is not as mutually exclusive with BRAF or NRAS as they are with each other (Figures 2A–2C, see STAR Methods for description of data acquisition, sorting, and analysis). No significant difference in mutual exclusivity was detected when BRAF NRAS was compared to another known mutually exclusive gene pair (EGFR and KRAS in lung cancer, Figure 2D). Looking back at our simulated patient cohorts (black cross in Figures 1C and 1D), it is clear that this observed lack of mutual exclusivity of ALK<sup>ATI</sup> is not confounded by its low abundance. If there were 12 ALK<sup>ATI</sup> patients (not 38 ALK<sup>ATI</sup> patients) in our dataset, we would have been unable to make any conclusions about whether a BRAF-NRAS level of mutual exclusivity exists in ALK<sup>ATI</sup>. Thus, ALK<sup>ATI</sup> is significantly less mutually exclusive with BRAF or NRAS than they are with each other.

We also considered the possibility that the lack of mutual exclusivity of ALK<sup>ATI</sup> with BRAF and NRAS was because of the definition of the initial filter cutoffs in the original RNA-seq analysis (the exact filters are mentioned in the STAR Methods). By systematically changing the cutoffs for all of the filters combinatorially, we varied the sensitivity for ALK<sup>ATI</sup> detection by orders of magnitude (Figure 3A, top). More stringent filter sets resulted in fewer ALK<sup>ATI</sup> calls, whereas less stringent filters resulted in more calls (Figure 3A, bottom). Performing pairwise comparisons of conditional selection on all combinations of these filter cutoffs never created mutual exclusivity. This filter analysis indicated that mutual exclusivity in ALK<sup>ATI</sup> could not be observed with any data driven RNA-seq definition.

This analysis shows the utility of tests for pairwise conditional selection to demonstrate quantitative differences in the degree of mutual exclusivity between a positive control gene pair and a new and potentially oncogenic alteration. We also strongly demonstrate a lack of mutual exclusivity between ALK<sup>ATI</sup> and established MAPK pathway driver mutations.

**Kinase domain expression imbalance in ALK is nearly ubiquitous in melanoma and lung cancers**

Given the conclusive lack of mutual exclusivity between ALK<sup>ATI</sup> and transforming melanoma oncogenes, we decided to look deeper at the initial signal, i.e., the bias toward ALK expression in the kinase domain (Exons 20:29). We expected to see a distribution centered at equivalent expression between the kinase domain and the upstream coding region (the diagonal line in Figure 3B). We posited that expression levels in exons 1-19 and 20-29 should be distributed above and below the diagonal, with ALK<sup>ATI</sup> patients being strong outliers from this expected relationship. However, we observed a significant bias toward overexpression of the kinase domain of ALK across all melanoma patients, p value from Kolmogorov-Smirnov test: 2.2 × 10<sup>−16</sup>. In fact, almost all the skin cutaneous melanoma (SKCM, n = 340) and lung adenocarcinoma (LUAD, n = 477) patients in the TCGA (Figures 3B and S3A) expressed higher levels of the ALK kinase domain than exons 1-19. The ubiquity of this deviation in all patients led us to suspect that a systematic error could be at play. Although multiple interpretations of this signal exist, it is concerning that all patients have some propensity to overexpress the kinase domain. As a control, we also examined whether an imbalance of expression toward the kinase domain is a unique feature of ALK. We did not see this propensity for kinase domain expression in EGFR in SKCM or in EGFR in LUAD, p value from KS-test:0.66 (Figure S3B). Hence, the kinase domain expression imbalance is not a generalizable finding and raises concerns of potential systemic biases in exon specific expression levels in ALK in the TCGA.

**ALK<sup>ATI</sup> is not sufficient for growth/transformation in vitro**

Our computational reanalysis of ALK expression data in melanoma suggested that ALK<sup>ATI</sup> is significantly less mutually exclusive with NRAS and BRAF than they are with each other. In the original Wiesner et al. study (Wiesner et al., 2015), ALK<sup>ATI</sup> was argued to be sufficient for transformation of Ba/F3s to growth factor independence.
When we transduced Ba/F3s with ALK ATi and EML4-ALK (9 independent replicates each), we found that ALK ATi took significantly longer to grow out (two population doublings in 6.7 ± 0.4 days for EML4-ALK and 10.0 ± 1.0 days for ALK ATi, Figure 4A). However, we also reasoned that longer outgrowth times were indicative of a weaker transforming potential. As such, we scaled up the number of transductions to 48 independent replicates by performing many parallel transductions of ALK ATi, EML4-ALK, and vector. The results were striking. Growth factor independence was observed in 100% of EML4-ALK replicate transductions, but only 37.5% ALK ATi replicate transductions, and in 16% of vector controls (Figure 4B). Viral titers were essentially
Figure 3. Changing the RNA-Seq filter cutoffs does not identify mutual exclusivity in ALK^ATI

(A) Top: Changing the RNA-seq filter cutoffs for RSEM (Li and Dewey, 2011), RPKM (Mortazavi et al., 2008), and count data did not result in mutually exclusive regions of ALK^ATI with BRAF or NRAS. Filters used were RSEM 10-1000, counts data did not result in mutually exclusive regions of ALK^ATI with BRAF or NRAS. Filters used were RSEM 10-1000, counts data did not result in mutually exclusive regions of ALK^ATI with BRAF or NRAS. Filters used were RSEM 10-1000, counts...
confounded by secondary genetic events. To confirm that these mutations in ALK ATI are sufficient for trans-
anoma cell line model. We transduced ALK ATI into two BRAF V600E-harboring melanoma cell lines, and chal-
formation, we added the 3 mutations to the original ALK ATI plasmid via site directed mutagenesis. When
potential should be able to rescue BRAF V600E melanoma from a BRAF V600E inhibitor.

2010; Chapman et al., 2011; Swaika et al., 2014, whereas receptor tyrosine kinases signal through dimeric
RAF family proteins (Freeman et al., 2013). A candidate oncogenic RTK in melanoma with strong transforming
necessary for melanoma cell survival, and sufficient for transformation (Chapman et al., 2011; Sensi et al.,
2012). When we did this, ALK ATI and vector-transduced melanoma cells had statistically indistinguishable
differences in growth factor independence, we decided to test the potential oncogenicity of ALK ATI in a more realistic mel-
Next, we reasoned that growth factor independence in ALK ATI Ba/F3 cells was infrequent because it required a relatively rare second genetic event. To test this hypothesis, we extracted the genomic DNA of the 18 ALK ATI Ba/F3 cell lines that achieved growth factor independence and sequenced their ALK kinase domain. Surprisingly, 3 of the 18 ALK ATI cell lines had well known transforming mutations in the ALK kinase (F1174C, F1174V, and F1174I, Figures 4A and S4B). These point mutations are the primary cancer causing mutations in ALK-mutated neuroblastoma and have been shown to constitutively activate the ALK kinase (Tate et al., 2019). These mutations are also sensitive to some ALK kinase inhibitors. This gives a concrete rationale for why ALK ATI could transform Ba/F3 cells, and that those cells were sensitive to ALK inhibition (Wiesner et al., 2015). We hypothesize that these mutations spontaneously arise in a small subset of trans-
duced Ba/F3 cells, but that IL3 withdrawal strongly selects for the growth benefit conferred by these
artifactual mutations. Although the mutations were not found in all cells, their existence in 3 independent selections strongly suggests that the transformation results and therapeutic treatments of ALK ATI can be confounded by secondary genetic events. To confirm that these mutations in ALK ATI are sufficient for transformation, we added the 3 mutations to the original ALK ATI plasmid via site directed mutagenesis. When we performed new transformation experiments, ALK ATI_F1174_C, V, and I mutant cells transformed Ba/F3’s in a highly efficient manner that resembled our EML4-ALK positive control (Figure S4C). To further confirm that these cells were ALK addicted, we treated Ba/F3 cells with crizotinib and examined their dose-response curves and signaling state via western blots. ALK ATI transformed Ba/F3 cells are sensitive to crizotinib and brigatinib (Figures 4C and S4D). Crizotinib treatment decreases phospho-ALK and phospho-ERK in a manner similar to EML4-ALK (Figure 4D). Together, this data suggests that while ALK ATI transformed Ba/F3 are sensitive to crizotinib, they require a second transforming event (i.e F1174 mutations) to fully transform cells in vitro. These mutations are not seen in melanoma patients. Alongside our re-analysis of the genomic data and the Couts et al data (Couts et al., 2017), our transformation data strongly suggested that ALK ATI is not sufficient for growth/transformation in vitro, and that ALK inhibitors will not be a good single agent treatment strategy in ALK ATI positive melanoma. Though other transformation experiments were performed in (Couts et al., 2017), we argue that the large scale replication of the Ba/F3 results in our lab cast significant concerns on the other transformation studies in ALK ATI.

ALK ATI cannot rescue melanoma cell lines

Although our Ba/F3 analysis suggests that the ALK ATI alterations are not sufficient to transform tool cell lines to growth factor independence, we decided to test the potential oncogenicity of ALK ATI in a more realistic melanoma cell line model. We transduced ALK ATI into two BRAF V600E-harboring melanoma cell lines, and challenged them with a BRAF V600E inhibitor, vemurafenib. The experimental rationale was simple: BRAF V600E is necessary for melanoma cell survival, and sufficient for transformation (Chapman et al., 2011; Sensi et al., 2006). The drug vemurafenib inhibits only the mutant BRAF V600E protein as a monomer (Bollag et al., 2010; Chapman et al., 2011; Swaika et al., 2014), whereas receptor tyrosine kinases signal through dimeric RAF family proteins (Freeman et al., 2013). A candidate oncogenic RTK in melanoma with strong transforming potential should be able to rescue BRAF V600E melanoma from a BRAF V600E inhibitor.

We transduced EML4-ALK, ALK ATI, and vector into two different skin cancer cell lines, SKMEL28 and G361, both of which have a transforming V600E point mutation and are sensitive to vemurafenib (Barretina et al., 2012). When we did this, ALK ATI and vector-transduced melanoma cells had statistically indistinguishable vemurafenib dose responses (p value: 0.49 for SKMEL28 and 0.97 for G361, Figure 5A). Western blots confirmed ALK ATI constructs robustly expressed, and that both melanoma cell lines exhibited strong over-expression of the ALK ATI construct (Figures 5B and SC). The inability of ALK ATI to rescue melanoma cell lines is in line with the notion that ALK ATI is not an oncogenic driver in melanoma.

Figure 3. Continued

100-1000, 20-29/ex20-29 ratio 1-100. Only the minimum p value observed amongst all exon ratios tested is displayed. The midpoint of the p value color gradient is 0.3. Regions that filtered for <10 ALK ATI patients were not included in the pairwise comparison analysis and are colored grey. P-values calculated using a Fisher’s exact test. Bottom: The number of patients that are categorized as ALK ATI positive decreases as the filter stringency increases. (B) The kinase domain of ALK (ex. 20-29) is significantly overexpressed in the majority of SKCM patients (p value is from a χ² test).
We were unable to transduce SKMEL28 and G361 cells with EM4-ALK on multiple transduction attempts. However, we decided to proceed with our vemurafenib challenge in the absence of EML4-ALK. EML4-ALK is an established oncogenic driver in NSCLC and melanoma (Lin et al., 1998; Takahashi et al., 2010, p. 4). We confirmed our viral titer and infectivity by simultaneously transducing and transforming Ba/F3 to IL-3 independence, as well as infecting Hek293T and selecting for puromycin resistance. Interestingly, virus packaged with the EML4-ALK oncogene readily transformed Ba/F3 cells, easily infected Hek293T cells, and selected with puromycin, but we could not successfully select for EML4-ALK containing SKMEL28 or G361 cells (Table S1). Because this was a negative result, we tested the idea that conditional selection via mutual exclusivity might be occurring. Our hypothesis is that simultaneous expression of two growth pathway activating variants in the same cell is not well tolerated, as is previously observed (Ciriello et al., 2012; Cisowski et al., 2016; Mina et al., 2017; Petti et al., 2006). To test this, we performed the same infections that are described above, but during the selection of the infected cells, we performed all selections in both the presence and absence of crizotinib. Whereas vector control and ALK<sup>ATI</sup> cells formed stable cell lines in the presence and absence of crizotinib, we were only able to...
**Figure 5. ALK\(^{ATI}\) cannot replace known oncogenes in melanoma**

(A) Dose response of ALK\(^{ATI}\) and vector-transduced SKMEL-28 and G-361 cell lines. ALK\(^{ATI}\) does not improve the dose response of SKMEL-28 (p-val: 0.49) and G361 (p-val: 0.99) to vemurafenib. p value calculated using a one-sided paired t test test between ALK\(^{ATI}\) and vector. Error bars represent standard deviation on 3 replicates of 3 independent transductions (9 total replicates per condition per concentration).

|     | Vector | ALK\(^{ATI}\) | EML4ALK |
|-----|--------|---------------|---------|
| G361|        |               |         |
|     | DSMO   | ++            | ++      |
|     | Crizotinib | ++ | ++      |

**Key**

++ Outgrowth during puromycin selection, included in western
++ Some outgrowth during puromycin selection, included in western
- No outgrowth during puromycin selection, unable to include in western

(B) Data from Figure 5A

(C) Representative western blots for G361 and SKMEL28 showing ALK\(^{ATI}\) and EML4ALK knockdown. 

(D) p value: 0.39

(E) p value: 0.93
stably select melanoma cells expressing EML4-ALK when we selected for transductants in the presence of the ALK inhibitor crizotinib at 100nM (Figure 5B, Table S1). This suggests that melanoma cells require ALK inhibition in order to stably express EML4-ALK. The crizotinib dependence of the known oncogene EML4-ALK and the crizotinib independence of ALK ATI in melanoma cells alongside the inability of ALK ATI to rescue a vemurafenib challenge suggest that ALK ATI does not act as a constitutively active oncogene in melanoma cells. It also suggests that the expression of an activated ALK construct and BRAF V600E actually inhibits melanoma growth.

**ALK expression imbalance does not predict transforming potential or single agent therapeutic dependency**

We further probed the transforming potential of ALK ATI by searching for evidence for an ALK ATI dependent transforming potential *in vitro*. To do this, we analyzed BRAF-mutant cell lines (n = 43) from CCLE (Barretina et al., 2012) for their ALK expression and dose response to BRAF inhibitors. In this dataset, 4 of these 43 cell lines were ALK ATI-like (exceeded 2/3 filters). None of these 4 cell lines showed a substantial difference in their sensitivity to 11 distinct BRAF inhibitors (Figures 5D and S5A). Furthermore, the degree of exon imbalance in ALK expression (when treated as a continuous variable) did not predict improved survival against a BRAF-inhibitor challenge.

Although previous experiments suggested that ALK ATI may not be sufficient for oncogenesis, we still wondered if ALK ATI conferred sensitivity to ALK inhibitors in melanoma cell lines *in vitro*. To test this, we analyzed melanoma cell lines (n = 32) from CCLE (Barretina et al., 2012), 4 of which were ALK ATI-like (exceeded 2/3 filters). No evidence of ALK-inhibitor sensitivity was found in any of the 4-ALK ATI-like cell lines (Figures 5E and S5B). Furthermore, the degree of exon imbalance in ALK expression did not predict ALK-inhibitor responses in these 32 cell lines (linear regression p value for crizotinib: 0.93). Hence, the degree of observed exon imbalance is not correlated with single agent sensitivity to ALK-inhibition across 32 melanoma cell lines.

**DISCUSSION**

Conditional selection has been used to study mechanisms of oncogenic activation in a variety of contexts. Previous research has used mutual exclusivity as an indicator of oncogenic network modules and dysregulated pathway analyses, and to identify evolutionary dependencies from alteration occurrences in pan-cancer analyses. Importantly, mutual exclusivity is often used to prioritize rare genomic findings without defining statistical power to detect a given effect size, or to compare to an expected effect size. To our knowledge, no previous method accurately quantifies a negative signal for conditional selection in rare variants. Our code base allows us to quantify a negative finding relative to how often a lack of mutual exclusivity would be seen in a positive control gene pair. Our simulations show that our method for pairwise comparisons is a quantitative and statistically robust method to identify a lack of conditional selection with sufficient statistical power.

We applied our pairwise comparisons of conditional selection to a transcript alteration in ALK ATI because of the controversy in the literature (Couts et al., 2017; Wiesner et al., 2015). Our analysis clearly demonstrates a lack of conditional selection for ALK ATI. ALK ATI is significantly less mutually exclusive than BRAF and NRAS are with each other in melanoma. Moreover, our experiments suggest that ALK ATI is not sufficient for cellular transformation, that kinase domain imbalance does not predict inhibitor response, and that single agent ALK inhibition is unlikely to be therapeutic in melanoma cells.
In their original paper, Wiesner et al. (Wiesner et al., 2015) performed a substantive and detailed description of the ALK<sup>ATI</sup> event. They found enrichment of H3K4me3 and RNA Pol II near the ATI transcription initiation site of tumors expressing ALK<sup>ATI</sup> (Wiesner et al., 2015). Wiesner et al also confirmed that ALK<sup>ATI</sup> is expressed at both ALK alleles by comparing DNA, RNA, and H3k4me3 levels. They also performed gene expression profiling of RNA-Seq datasets showing that ALK<sup>ATI</sup>-like expression is found in 2-11% of melanoma samples and sporadically in various other tumor types and not in normal tissue samples (Busam et al., 2016; Wiesner et al., 2015). The breadth and depth of the analysis leads us to believe that ALK<sup>ATI</sup> is likely a true transcript variant. Moreover, although we believe our work (alongside the work of Couts et al (Couts et al., 2017)) provides strong evidence against single agent therapy, it would be unfair to ignore the potential for therapeutic relevance of ALK<sup>ATI</sup> in contexts other than ALK inhibitor monotherapy. Moreover, from a biological perspective, it would also be unfair to rule out some sort of unknown biological role for ALK<sup>ATI</sup> that does not fit conventional definitions of driver oncogenes. However, combining Couts’ PDX data with the questionable transformation potential of ALK<sup>ATI</sup> (Couts et al., 2017), the lack of objective responses in ALK<sup>ATI</sup> expressing melanomas to ALK inhibitors, and the clear lack of mutual exclusivity of ALK<sup>ATI</sup> in melanoma casts significant doubt upon the single agent therapeutic rationale for ALK<sup>ATI</sup>.

In the melanoma landscape, dramatic responses to approved and investigational immunotherapy agents are yielding important steps forward for patient care. We have systematically shown that the original ALK<sup>ATI</sup> data should be re-evaluated in light of our data reproducing the original finding, and in light of the compelling recent reports in PDX models (Couts et al., 2017; Uguen et al., 2016). Combined with the fact that the patient data in the original manuscript did not achieve the typical clinical criteria for an objective partial response, we strongly recommend that single agent ALK inhibitors receive no further testing in ALK<sup>ATI</sup> patients when other investigational or off label options exist. Given the weight of evidence, it seems unlikely that refractory patients will benefit from this treatment. We also suggest that pairwise tests for conditional selection will be a useful tool to triage any rare genomic finding in the mountains of cancer sequencing data generated in late-stage patients.

**Limitations of study**

4 of the 43 melanoma cell lines in the CCLE (Barretina et al., 2012) matched the expression filters that are characteristic of alternate transcription initiation. The drug sensitivity analysis showed that ALK<sup>ATI</sup>-like expression in these cell lines does not predict sensitivity to ALK-inhibitors (Figure 4E), nor does ALK<sup>ATI</sup> seem to rescue BRAF-mutated melanoma undergoing BRAF inhibition (Figure 4D). Although these results are insightful, a more thorough analysis of the dose-response of ALK<sup>ATI</sup> would require more cell lines to be ALK<sup>ATI</sup>-like. Ultimately, a more thorough analysis will become possible as CCLE continues to add more cell lines to its repository.

**STAR METHODS**

Detailed methods are provided in the online version of this paper and include the following:

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Supplemental information can be found online at https://doi.org/10.1016/j.isci.2021.103343.

ACKNOWLEDGMENTS

This publication was supported, in part, by NIH Grants 5R21EB026617, T32GM108563, 1U01CA265709, R35GM133613, and R35GM124818, and by an Alfred P. Sloan Research Fellowship and additional support from the Simons Center for Quantitative Biology. We would like to thank Scott Leighow for his contribution with idea generation, critical analysis, and proofreading of this manuscript. We would also like to thank Kelly Hartsough, Lauren Randolph, Kyle McIlroy, and Joshua Reynolds for their help revising previous versions of this manuscript.

AUTHOR CONTRIBUTIONS

HI, IS, AS performed experiments. HI, YR, DM performed computational analyses. HI, IS, YR, FN, EOB, CD, DM, and JRP coauthored the manuscript.

DECLARATION OF INTERESTS

JRP has ownership interests in Theseus Pharmaceutical and MOMA therapeutics.

JRP is a co-founder of Theseus pharmaceuticals.

JRP has ownership interests in Theseus Pharmaceuticals and MOMA Therapeutics.

Received: October 22, 2020
Revised: June 17, 2021
Accepted: October 22, 2020
Published: November 19, 2021

REFERENCES

Barretina, J., Caponigro, G., Stransky, N., Venkatesan, K., Margolin, A.A., Kim, S., Wilson, C.J., Lehár, J., Kryukov, G.V., Sonkin, D., et al. (2012). The Cancer Cell Line Encyclopedia enables predictive modelling of anticancer drug sensitivity. Nature 483, 603–607. https://doi.org/10.1038/nature10913.

Bhullar, K.S., Lágró, N.O., McGowan, E.M., Parmar, I., Jha, A., Hubbard, B.P., and Rupasinghe, H.P.V. (2018). Kinase-targeted cancer therapies: progress, challenges and future directions. Mol. Cancer 17, 48. https://doi.org/10.1186/s12943-018-0804-2.

Bischak, J.D., Carbonetto, P., and Stephens, M. (2019). Creating and sharing reproducible research code the workflowr way. F1000Res 8. https://doi.org/10.12688/f1000research.20843.1.

Bollag, G., Hirth, P., Tsai, J., Zhang, J., Ibrahim, P.N., Cho, H., Spevak, W., Zhang, C., Zhang, Y., Habets, G., et al. (2010). Clinical efficacy of a RAF inhibitor needs broad target blockade in BRAF-mutant melanoma. Nature 467, 596–599. https://doi.org/10.1038/nature09454.

Burrell, R.A., McGranahan, N., Bartek, J., and Swanton, C. (2013). The causes and consequences of genetic heterogeneity in cancer evolution. Nature 501, 338–343. https://doi.org/10.1038/nature12625.

Busam, K.J., Villain, R.E., Lum, T., Busam, J.A., Hollmann, T.J., Saw, R.P.M., Coit, D.C., Scolyer, R.A., and Wiesner, T. (2016). Primary and metastatic cutaneous melanomas express ALK through alternative transcriptional initiation. Am. J. Surg. Pathol. 40, 766–795. https://doi.org/10.1097/PAS.0000000000000611.

Chapman, P.B., Hauschild, A., Robert, C., Haenen, J.B., Ascieto, P., Larkin, J., Dummer, R., Garbe, C., Testori, A., Maio, M., et al. (2011). Improved survival with vemurafenib in melanoma with BRAF V600E mutation. New Engl. J. Med. 364, 2507–2516. https://doi.org/10.1056/NEJMoa1103782.

Ciriello, G., Cerami, E., Sander, C., and Schultz, N. (2012). Mutual exclusivity analysis identifies oncogenic network modules. Genome Res. 22, 398–406. https://doi.org/10.1101/gr.125567.111.

Cisowski, J., Sayin, V.I., Liu, M., Karlsson, C., and Bergo, M.O. (2016). Oncogene-induced senescence underlies the mutual exclusive nature of oncogenic KRAS and BRAF. Oncogene 35, 1328–1333. https://doi.org/10.1038/onc.2015.186.

Couts, K.L., Remis, J., Turner, J.A., Bagby, S.M., Murphy, D., Christiansen, J., Hintzsche, J.D., Le, A., Pitts, T.M., Wells, K., et al. (2017). ALK inhibitor response in melanomas expressing EML4-ALK fusions and alternate ALK isoforms. Mol. Cancer Ther. https://doi.org/10.1158/1535-7163.MCT-17-0472.

Edlundh-Rose, E., Egyházi, S., Omholt, K., Mansson-Brahme, E., Platz, A., Hansson, J., and Lundéberg, J. (2006). NRAS and BRAF mutations in melanoma tumours in relation to clinical characteristics: a study based on mutation screening by pyrosequencing. Melanoma Res. 16, 471–478. https://doi.org/10.1097/01.cmr.0000252300.22052.86.

Fakih, M., O’Neil, B., Price, T.J., Falchuck, G.S., Desai, J., Kuo, J., Govindan, R., Rasmussen, E., Morrow, P.K.H., Ngang, J., et al. (2019). Phase 1 study evaluating the safety, tolerability, pharmacokinetics (PK), and efficacy of AMG 510, a novel small molecule KRASG12C inhibitor, in advanced solid tumors. JCO. 37, 3003. https://doi.org/10.1200/JCO.2019.37.15_suppl.3003.

Freeman, A.K., Ritt, D.A., and Morrison, D.K. (2013). The importance of Raf dimerization in cell signaling. Small GTPases 4, 180–185. https://doi.org/10.4161/sgtp.26117.

Gainor, J.F., Varghese, A.M., Ou, S.-H.I., Kabraji, S., Awad, M.M., Katayama, R., Pawlak, A., Minokudison, M., Yeap, B.Y., Rielypress, G.J., et al. (2013). ALK rearrangements are mutually exclusive with mutations in EGFR or KRAS: an analysis of 1,683 patients with non–small cell lung cancer. Clin. Cancer Res. https://doi.org/10.1158/1078-0432.CCR-13-0318.

Guan, J., Gupta, R., and Filipp, F.V. (2015). Cancer systems biology of TCGA SKCM: efficient detection of genomic drivers in melanoma.
Poynter, J. N., Elder, J. T., Fullen, D. R., Nair, R. P., Soengas, M. S., Johnson, T. M., Redman, B., Thomas, N. E., and Gruber, S. B. (2006). BRAF and NRAS mutations in melanoma and melanocytic nevi. Melanoma Res. 16, 267–273. https://doi.org/10.1097/01.mcr.0000226690.73179.f3

Roskoski, R., and Sagedhi-Nejad, A. (2018). Role of RET protein-tyrosine kinase inhibitors in the treatment RET-driven thyroid and lung cancers. Pharmacol. Res. 128, 1–17. https://doi.org/10.1016/j.phrs.2017.12.021

Schluumberger, M., Tahara, M., Winth, L. J., Robinson, B., Brose, M. S., Elisei, R., Dutcus, C. E., de las Heras, B., Zhu, J., Habra, M. A., et al. (2014). A phase 3, multicenter, double-blind, placebo-controlled trial of lenvatinib (E7080) in patients with 131I-refractory differentiated thyroid cancer (SELECT). JCO 32. https://doi.org/10.1200/jco.2014.32.18_suppl.lba6008

Sensi, M., Nicolini, G., Petti, C., Bersani, I., Lozupone, F., Molla, A., Vegetti, C., Nonaka, D., Mortarini, R., Parmiani, G., et al. (2006). Mutually exclusive NRASQ61R and BRAFV600E mutations at the single-cell level in the same human melanoma. Oncogene 25, 3357–3364. https://doi.org/10.1038/sj.ong.1209379

Shaw, A.T., Camidge, D.R., Engelman, J.A., Solomon, B.J., Kwak, E.L., Clark, J.W., Salgia, R., Shapiro, G., Bang, Y.-J., Tan, W., et al. (2012). Clinical activity of crizotinib in advanced non-small cell lung cancer (NSCLC) harboring ROS1 gene rearrangement. JCO 30, 7508. https://doi.org/10.1200/jco.2012.30.15_suppl.7508

Swaiwa, A., Crozier, J.A., and Joseph, R.W. (2014). Vemurafenib: an evidence-based review of its clinical utility in the treatment of metastatic melanoma. Drug Des. Devel. Ther. 8, 775–787. https://doi.org/10.2147/dddt.S31143

Takahashi, T., Sonobe, M., Kobayashi, M., Yoshizawa, A., Menju, T., Nakayama, E., Mino, N., Iwaki, S., Sato, K., Miyahara, R., et al. (2010). Clinicopathologic features of non-small-cell lung cancer with EML4–ALK fusion gene. Ann. Surg. Oncol. 17, 889–897. https://doi.org/10.1245/s10434-009-0808-7

Tate, J. G., Barnford, S., Jubb, H.C., Sundka, Z., Beare, D. M., Bindal, N., Boutsakas, H., Cole, C. G., Creature, C., Dawson, E., et al. (2019). COSMIC: the catalogue of somatic mutations in cancer. Nucleic Acids Res. 47, D941–D947. https://doi.org/10.1093/nar/gkx1015

Uguen, A., Uguen, M., and Guibourg, B. (2016). ALK expression in melanomas: Looking for a needle in a Haystack. Am. J. Surg. Pathol. 40, 1437. https://doi.org/10.1097/PAS.000000000000686

Unni, A.M., Lockwood, W.W., Zejnullahu, K., Lee-Lin, S.-Q., and Varmus, H. (2015). Evidence that synthetic lethality underlies the mutual exclusivity of oncogenic KRAS and EGFR mutations in lung adenocarcinoma. Elife 4, e06907. https://doi.org/10.7554/eLife.06907

Vandin, F., Upfal, E., and Raphael, B.J. (2012). De novo discovery of mutated driver pathways in cancer. Genome Res. 22, 375–385. https://doi.org/10.1101/gr.120477.111

Varmus, H., Unni, A.M., and Lockwood, W.W. (2016). How cancer genomics drives cancer biology: does synthetic lethality explain mutually exclusive oncogenic mutations? Cold Spring Harb. Symp. Quant. Biol. 81, 247–255. https://doi.org/10.1101/cshperspect.a016866

Wells, S.A., Robinson, B.G., Gagel, R.F., Dralle, H., Fagin, J.A., Santoro, M., Baudin, E., Elisei, R., Jarzab, B., Vasselli, J.R., et al. (2012). Vandetanib in patients with locally advanced or metastatic medullary thyroid cancer: a randomized, double-blind phase III trial. J. Clin. Oncol. 30, 134–141. https://doi.org/10.1200/JCO.2011.35.5040

Whitaker, S.R., Theurillat, J.-P., Allen, E.V., Wagle, N., Hsiao, J., Cowley, G.S., Schadendorf, D., Root, D.E., and Garraway, L.A. (2013). A genome-scale RNA interference screen implicates NF1 loss in resistance to RAF inhibition. Cancer Discov. 3, 350–362. https://doi.org/10.1158/2159-8290.CD-12-0470

Widmann, C., Gibson, S., Jarpe, M.B., and Johnson, G.L. (1999). Mitogen-activated protein kinase: Conservation of a three-kinase module from yeast to human. Physiol. Rev. 79, 143–180. https://doi.org/10.1152/physrev.1999.79.1.143

Wiesner, T., Lee, W., Obernauf, A.C., Ran, L., Murali, R., Zhang, Q.F., Wong, E.W.P., Hu, W., Scott, S.N., Shah, R.H., et al. (2013). Alternative transcription initiation leads to expression of a novel ALK isoform in cancer. Nature 526, 453–457. https://doi.org/10.1038/nature15258

Zhang, W., and Liu, H.T. (2002). MAPK signal pathways in the regulation of cell proliferation in mammalian cells. Cell Res. 12, 9. https://doi.org/10.1038/sj.cr.7290105

Zhao, B., and Pritchard, J.R. (2014). Inherited disease genetics improves the identification of cancer-associated genes. PLOS Genet. 12, e1006081. https://doi.org/10.1371/journal.pgen.1006081

Zhao, J., Zhang, S., Wu, L.-Y., and Zhang, X.-S. (2012). Efficient methods for identifying mutated driver pathways in cancer. Bioinformatics 28, 2940–2947. https://doi.org/10.1093/bioinformatics/bts564
STAR METHODS

KEY RESOURCES TABLE

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| Antibodies          |        |            |
| Mouse monoclonal anti-ALK | CST     | Cat#3791; RRID: AB_1950402 |
| Rabbit monoclonal anti-pALK | CST     | Cat#3341; RRID: AB_331047 |
| Rabbit monoclonal anti-pERK 1/2 | CST     | Cat#4370; RRID: AB_2315112 |
| Rabbit anti-ERK 1/2 | CST     | Cat#9102; RRID: AB_330744 |
| Rabbit monoclonal anti-β-Actin | CST     | Cat#4970; RRID: AB_2223172 |
| Anti-rabbit IgG     | CST     | Cat#7074; RRID: AB_2099233 |
| Anti-mouse IgG      | CST     | Cat#7076; RRID: AB_330924 |

| Chemicals, peptides, and recombinant proteins |        |
|------------------------------------------------|--------|
| Vemurafenib                                     | Sellekchem | Catalog No.S1267 |
| Crizotinib                                      | Sellekchem | Catalog No.S1068 |
| Brigatinib                                      | Sellekchem | Catalog No.S8229 |

| Critical commercial assays                      |        |
|------------------------------------------------|--------|
| CellTiter Glo                                   | Promega | G7570 |
| QuikTiter Lentivirus Quantitation Kit (HIV p24) | Cell BioLabs | VPK-107 |
| QS Site-directed mutagenesis kit                | NEB    | E05545 |

| Deposited data                                  |        |
|------------------------------------------------|--------|
| Deposited data                                  | Github Repository | NA |

| Experimental models: Cell lines                 |        |
|------------------------------------------------|--------|
| BaF3 Cells                                      | DSMZ    | ACC 300 |
| SK-MEL-28                                       | ATCC    | HTB-72 |
| G-361                                           | ATCC    | CRL-1424 |
| HEK-293T                                        | ATCC    | CRL-1573 |

| Recombinant DNA                                 |        |
|------------------------------------------------|--------|
| pLVX-IRES-Puro                                  | Clontech | Cat#632183 |
| mRNA Sequence ALK<sup>ATI</sup>                 | European Nucleotide Archive | LN864494.1 |
| mRNA sequence for EML4-ALK                      | GenBank | AB274722.1 |

RESOURCES AVAILABILITY

Lead contact
Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact, Justin Pritchard (jp94@psu.edu).

Material availability
All unique/stable reagents generated in this study (including AL<sup>ATI</sup> BaF3 cell lines) are available from the Lead Contact with a completed Materials Transfer Agreement.

Data and code availability
Our Github repository includes all the data used and generated during our analyses (in the data and output directories).
The Github repository also includes all the code used to parse, analyze, and plot the data (in the code and analysis directories).

The GitHub repository is also available as a Github page, which contains the output of the Rmarkdown analyses. The pairwise comparisons page was created with the help of the r package WorkflowR (Blischak et al., 2019).

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Cell culture

Cell lines used were: SK-MEL-28 (ATCC HTB-72), G-361 (ATCC CRL-1424), BaF3 (DSMZ ACC-300), HEK-293T (ATCC CRL-1573). Prior to use, all cell lines were tested to be free of mycoplasma using a biochemical-based test (Mycoalert plus, Lonza). BaF3 cell lines were cultured in RPMI while HEK-293T, Skmel-28 and G361 cell lines were cultured in DMEM. Media was supplemented with a final concentration of 10% FBS, and 1% Penicillin/Streptomycin/L-Glutamine. WT Ba/F3 cell lines were grown in media supplemented with 10ng/mL IL3 (Pepro-tech). Stable transductions were verified for puromycin resistance by using kill curve concentrations ranging from 0.25ug/mL to 2ug/mL. Cells were split when they were 70-80% confluent. The subculture ratios used for all of our cell lines included splitting ratios of 1:5 every 2-4 days when cells were 70-80% confluent.

METHOD DETAILS

Frequency correction in gene pairs

In these methods, frequency of an event refers to its relative frequency (empirical probability), not absolute frequency (count).

Suppose we have a joint table from our positive controls with frequencies

| Positive Control 2 | Yes | No |
|--------------------|-----|----|
| Positive Control 1 |     |    |
| Yes                | p₁₁ | p₁₀ |
| No                 | p₀₁ | p₀₀ |

Table: A 2x2 Contingency table of the positive control 1 and positive control 2 genes. Where \( p_{11} \) refers to the frequency of the population having mutations in both the positive control 1 gene and the positive control 2 gene; \( p_{10} \) refers to the population having a mutation in the positive control 1 gene only, and so on.

The odds ratio, defined as the strength of association between two events, is \( (p_{11} + p_{00}) / (p_{10} + p_{01}) \) for Table above.

Also suppose we have a GOI with frequency \( p_{GOI} \). We would like to transform the above table to a new table such that Positive Control 1 positive individuals appear at frequency \( p_{GOI} \), but where the odds ratio remains the same. We can do this by adding or removing the appropriate fraction of Positive Control 1 positive individuals without regard to their status with respect to Positive Control 2. This results in a new table with cell probabilities given by:

| Positive Control 2 | Yes | No |
|--------------------|-----|----|
| Positive Control 1 |     |    |
| Yes                | \( p_{GOI} \) \( p_{11} / (p_{10} + p_{11}) \) | \( (1 - p_{GOI}) \) \( p_{01} / (p_{00} + p_{01}) \) |
| No                 | \( (1 - p_{GOI}) \) \( p_{00} / (p_{00} + p_{01}) \) | \( p_{GOI} \) \( p_{01} / (p_{00} + p_{01}) \) |

Table: A 2x2 Contingency table of the positive control 1 and positive control 2 genes with the frequency of mutations in the positive control 1 gene corrected to the frequency of mutations in the gene of interest.
This table is derived by multiplying each row in the original table by an appropriate constant (e.g., the first row is multiplied by the ratio between the desired fraction of Positive Control 1 positive individuals \( p_{\text{GOI}} \) and the observed fraction in the \( p_{10}+p_{11} \) from the original table). It is easy to verify that the Table is a valid probability distribution with cells that sum to 1, that Positive Control 1 is now at frequency \( p_{\text{GOI}} \), and that the odds ratio is unchanged from the original table.

**Pairwise comparisons of gene pairs**

In order to estimate the typical range of odds ratios likely to be produced under the hypothesis that GOI and PC2 are as mutually exclusive as PC1 and PC2 while controlling for the observed frequency of mutations in the GOI, we can thus construct samples from the above table of size \( N \), where \( N \) is the number of patients in this cohort. Specifically, we construct these samples as a draw from a multinomial distribution with \( N \) trials and probabilities given by the above table of frequencies (Table) where the expected frequency of Positive Control 1 positive individuals has been set to \( p_{\text{GOI}} \). Independently drawing 1000 such tables and calculating the odds ratio for each table, we calculate the percent of PC1 vs PC2 odds ratios that have a lower odds ratio than the odds ratio for PC1 and GOI (labeled as score in Figure 1). The higher this score, the lower the overlap is between the odds ratios of the two gene pairs. A score of >95% was set as the threshold for rejecting the hypothesis that GOI and PC1 are as mutually exclusive as PC1 and PC2. Section 1 of the pseudo-code (Data S1) contains a step-by-step implementation of this process.

**Generating simulated cohorts at various mutual exclusivities and GOI frequencies**

We generated cohorts of gene pairs to test our pairwise comparisons method and to characterize its sensitivity. Suppose we would like to generate a joint table (such as Table above) from our positive controls with frequencies \( p_{11}, p_{10}, p_{01}, \) and \( p_{00} \). Also suppose we would like to generate a joint table with the gene of interest and positive control 1 gene with frequencies \( q_{11}, q_{10}, q_{01}, \) and \( q_{00} \).

Here, we outline a method to calculate the frequencies of these two gene pairs using three inputs:

- The odds ratio of the positive control genes, \( OR_p \), which can be written in terms of the frequencies in Table:

  \[
  OR_p = \frac{p_{11} + p_{00}}{p_{01} + p_{10}} \quad \text{(Equation 2.1)}
  \]

- The odds ratio of the gene of interest and positive control 1, \( OR_q \), which can be written in terms of its frequencies:

  \[
  OR_q = \frac{q_{11} + q_{00}}{q_{01} + q_{10}} \quad \text{(Equation 2.2)}
  \]

- The frequency of the gene of interest, \( p_{\text{GOI}} \):

  \[
  p_{\text{GOI}} = q_{11} + q_{01} \quad \text{(Equation 2.3)}
  \]

Since we are generating valid probability distributions, the frequencies of each gene pair sum to 1. Hence,

\[
q_{11} + q_{10} + q_{01} + q_{00} = 1 \quad \text{(Equation 2.4)}
\]

\[
p_{11} + p_{10} + p_{01} + p_{00} = 1 \quad \text{(Equation 2.5)}
\]

For simplicity of these derivations, we assume that the frequency of patients without a mutation in either gene is the same as the frequency of patients with a mutation in positive control 1 only:

\[
q_{00} = q_{10} \quad \text{(Equation 2.6)}
\]

\[
p_{00} = p_{10} \quad \text{(Equation 2.7)}
\]

This assumption was made to simplify the derivations in Equations 2.8, 2.9, and 2.10 below. Note that we also tested our pairwise comparisons method using a cohort generation strategy that does not make the
simplifying assumptions in Equations 2.6 and 2.7. i.e., \( p_{00} \) and \( q_{00} \) were pre-defined as inputs alongside \( \text{OR}_p \), \( \text{OR}_q \), and \( p_{\text{GOI}} \). Adjusting the frequencies of \( p_{00} \) and \( q_{00} \) this way did not significantly change the outcome of our simulation studies and did not change the conclusion of our simulation studies (results not shown here but are available on our GitHub page).

Equation 2.6 can be substituted into Equation 2.2:

\[
\text{OR}_q = \frac{q_{11} + q_{00}}{q_{01} + q_{10}} = \frac{q_{11}}{q_{01}}
\]  
(Equation 2.8)

Similarly, Equation 2.7 can be substituted into Equation 2.1:

\[
\text{OR}_p = \frac{p_{11} + p_{00}}{p_{01} + p_{10}} = \frac{p_{11}}{p_{01}}
\]  
(Equation 2.9)

Equations 2.3, 2.4, 2.6, and 2.8 can be arranged as a system of linear equations and solved using Gaussian elimination:

\[
\begin{bmatrix}
q_{11} + q_{10} + q_{01} + q_{00} = 1 \\
q_{10} - q_{00} = 0 \\
q_{11} - q_{01} + \text{OR}_q = 0
\end{bmatrix}
\rightarrow
\begin{bmatrix}
1 & 0 & 1 & 0 & \ p_{\text{GOI}} \\
1 & 1 & 1 & 1 & 1 \\
0 & 1 & 0 & -1 & 0
\end{bmatrix}
\rightarrow
\begin{bmatrix}
p_{\text{GOI}} + \text{OR}_q \\
1 - p_{\text{GOI}} \\
0 & 0 & 1 & 0 & \ p_{\text{GOI}} \\
0 & 0 & 1 & 0 & \ 1 + \text{OR}_q \\
\frac{1 - p_{\text{GOI}}}{2}
\end{bmatrix}
\]

This way, the frequencies of the GOI vs PC1 gene pair are solved as functions of the inputs:

| Gene of interest | Yes | No |
|------------------|-----|----|
| Positive Control 1 | \( \frac{p_{\text{GOI}} + \text{OR}_q}{1 + \text{OR}_q} \) | \( \frac{1 - p_{\text{GOI}}}{2} \) |
| | \( \frac{p_{\text{GOI}}}{1 + \text{OR}_q} \) | \( \frac{1 - p_{\text{GOI}}}{2} \) |

Table: A 2x2 contingency table of the positive control 1 gene and the gene of interest in which each frequency is a function of the input variables \( \text{OR}_q \) and \( p_{\text{GOI}} \).

Since we assume that the data for both the gene pairs comes from the same patient cohort, the overall frequency of mutations in the positive control 1 gene, termed \( p_{\text{PC1}} \), is the same in both the gene pairs. Therefore, \( p_{\text{PC1}} \) can be rewritten as a function of the inputs \( p_{\text{GOI}} \) and \( \text{OR}_q \):

\[
p_{\text{PC1}} = p_{11} + p_{10} = q_{11} + q_{10} = \frac{p_{\text{GOI}} + \text{OR}_q}{1 + \text{OR}_q} + \frac{1 - p_{\text{GOI}}}{2}
\]  
(Equation 2.10)

Therefore, Equations 2.5, 2.7, 2.9, and 2.10 can be arranged as a system of linear equations and solved using Gaussian elimination:

\[
\begin{bmatrix}
p_{11} + p_{10} = p_{\text{PC1}} \\
p_{11} + p_{10} + p_{01} + p_{00} = 1 \\
p_{10} - p_{00} = 0 \\
p_{11} - p_{01} + \text{OR}_p = 0
\end{bmatrix}
\rightarrow
\begin{bmatrix}
1 & 1 & 0 & 0 & \ p_{\text{PC1}} \\
1 & 1 & 1 & 1 & 1 \\
0 & 1 & 0 & -1 & 0
\end{bmatrix}
\rightarrow
\begin{bmatrix}
p_{\text{PC1}} \\
1 - p_{\text{PC1}} \\
0 & 0 & 1 & 0 & \ 1 - \text{OR}_p \\
0 & 0 & 0 & 1 & \ 1 - \text{OR}_p
\end{bmatrix}
\]
This way, the compartments of the positive control gene pair are solved:

| Positive Control 1 | Positive Control 2 | No |
|--------------------|--------------------|----|
| Yes                | \( \frac{O_{R_p} \times (1 - 2p_{PC1})}{1 - OR_p} \) | \( \frac{O_{R_p} (p_{PC1} - 1) + p_{PC1}}{1 - OR_p} \) |
| No                 | \( \frac{1 - 2p_{PC1}}{1 - OR_p} \) | \( \frac{1}{1 - OR_p} \) |

Table: A 2x2 contingency table of the two positive control 1 genes in which each frequency is a function of the input variables \( OR_p \) and \( p_{PC1} \).

Having established a method to calculate the frequencies of the two gene pairs using the three inputs \( OR_p \), \( OR_q \), and \( p_{GOI} \), we calculated these frequencies at various inputs. Each set of these input variables generates a contingency table containing the frequencies of the positive control 1 gene and the positive control 2 gene, and another contingency table containing frequencies of the positive control 1 gene and the gene of interest (see example output 2 in pseudo-code Supplemental information section 2). The odds ratio of the positive control, genes \( OR_p \), was varied from 0.01 (very mutually exclusive) to 0.1 (less mutually exclusive). The odds ratio of the gene of interest gene with the positive control 1 gene, \( OR_q \), was varied from 0.01 (mutually exclusive) to 1 (not mutually exclusive). The frequency of a mutation in the gene of interest, \( p_{GOI} \), in the cohort was varied from 0.005 (rare) to 0.5 (abundant). Figure S2A provides a visual illustration of how these input variables affect the simulated cohorts.

Once these contingency tables with frequencies of gene pairs were generated, the previously described pairwise comparisons method was used to correct the frequency of the positive control 1 gene. Next, simulations were performed for each of these frequency-adjusted contingency tables by drawing counts from a multinomial distribution with central tendencies around their respective frequencies using \( N \) trials, where \( N \) is the cohort size of the simulated cohort (example output 1 in the pseudo-code). We varied the cohort size of the simulations from 100 to 1,000 patients. Finally, the percentage of the odds ratios from the GOI and positive control 1 gene pair that are greater than the odds ratios of the pair of two positive control genes was calculated. The higher this percentage, the more certain one can be that the gene of interest is not as mutually exclusive with the positive control 1 gene as the two positive control genes are with each other. Supplemental information section 2 of the pseudo-code contains a step-by-step implementation of this process.

Analysis of public data sets

We downloaded our level 3 TCGA data from the Broad Institute TCGA GDAC Firehose (http://gdac.broadinstitute.org/) 2016_01_28 run. In our original ALK expression filters (Figure 2), ALK\(^{ATI} \) candidates were identified as samples with an ALK expression level of RSEM \( \geq 100 \), \( \geq 500 \) total reads across all ALK exons, and \( \geq 10 \)-fold greater average expression in exons 20–29 compared to exons 1–19. When varying filters, ALK\(^{ATI} \) patients were identified as samples with an ALK expression level of RSEM \( \geq 100 \), number of ALK reads \( \geq 500 \), and Exon20-29/Exon1-19 RPKM ratio \( \geq 10 \) (Figure 3). Cell line expression and drug sensitivity data (Figure 4) was downloaded from the cancer cell line encyclopedia (https://portals.broadinstitute.org/ccle/data). Melanoma cell-lines were classified as ALK\(^{ATI} \)-like if they matched 2/3 ALK\(^{ATI} \) filters.

Cell line expression and drug sensitivity data: CCLE 2019 RNA-seq gene expression data for 1019 cell lines were downloaded from the Broad Institute CCLE website (https://portals.broadinstitute.org/ccle). We used this data to identify ALK\(^{ATI} \)-like cell lines for their sensitivity to BRAF and ALK-inhibitors in melanoma. For 49 melanoma cell lines, ALK expression data was extracted and ALK\(^{ATI} \) expression was detected using two criteria: 1) ALK is expressed in the cell line with RSEM \( > 100 \), and read counts \( > 500 \), and 2) Since the ATI-site resides in intron 19 of ALK, a 10-fold greater expression of ALK exon20-29 than exon1-19 was expected for ALK\(^{ATI} \)-like cell lines.
For the identification of EML4-ALK cell lines, CCLE 2019 Fusion calls for 1019 cell lines were downloaded from CCLE. The EML4-ALK fusion calls were identified in 2 lung cancer cell lines (NCIH3122 and NCIH2228) and a pancreas cancer cell line (SNU324). For the identification of BRAF-mutant cell lines: 2019 cancer cell line mutant calls were downloaded from CCLE. A total of 111 BRAF mutants cell lines were identified that were targeted by 11 BRAF inhibitors.

**Plasmid generation**

The sequence of ALKATI was obtained from the European Nucleotide Archive under the accession number LN864494. The sequence of EML4-ALK was obtained from Genbank AB274722.1. The coding sequences, cloned into a pLVX-IRES-Puro backbone, were prepared by Genscript Gene Services. Please refer to Table S2 for the sequence of both of these plasmids.

**Lentiviral transduction**

Lentiviral constructs were co-transfected in HEK293T cells using calcium phosphate alongside third-generation packaging vectors that were pseudotyped with VSV-G. Viral supernatant was collected at 24 hours. All transductions were conducted limiting MOI to achieve the lowest viral titer required to produce IL-3 independence. After transduction with an oncogene, cell lines were selected with Puromycin to test transduction efficiency and subsequently with -IL3 to assess growth factor independence. All engineered ALK cell lines were sequenced using sanger sequencing to confirm their identity.

**In vitro transformation and drug treatment assays**

We used a standard Ba/F3 transformation protocol for lentiviral transduction. Lentiviral particles were made by transfecting HEK293T cells with the plasmid of interest and with packaging plasmids (3rd generation Lentiviral system from Addgene). Replication incompetent virus was collected using a BL2+ safety protocol. Upon virus collection, Ba/F3 cells at 500k/mL in 4mLs were infected with an equal volume of virus. A minimum of three replicates were used for each infection condition. Three days after infection, the Ba/F3s cells were spun out of virus and selected for IL3 independence and/or puromycin resistance (0.5ug/mL Puromycin was used). An infection was determined to be successful Ba/F3 cells for a given construct grew out for both IL3 independence and puromycin resistance. During selection, cell growth was quantified by doing daily counts of live/dead cells on a daily basis. These live/dead analyses were performed by supplementing 20uLs of cell culture with 0.4% trypan blue and subsequently counting live cells using a hemocytometer. Alternatively, live dead cells were counted using flow cytometry analysis (BD Accuri C6 Plus).

**Lentiviral particle quantification**

For each infection, before transducing Ba/F3s with virus, 500uL of virus was set aside and frozen at −20C. The number of lentiviral particles were quantified using the QuikTiter Lentivirus Quantitation Kit from Cell Biolabs (HIV P24 ELISA). All reported viral titers were within the linear range of the standard curve made using a positive control. Transduction efficiency was verified by counting outgrowth rates using puromycin selections across multiple infections.

**Immunoblots**

Cells were lysed in RIPA buffer (#9806S, CST) with 1 mM PMSF (P7626, MilliporeSigma), and Phosphatase Inhibitor Cocktail 2 (P5726, MilliporeSigma). After quantifying total protein concentration with the BCA assay (23225, ThermoFisher), 10 μg of each sample was boiled at 95°C for 5 min and loaded onto the 4-12% Bis-Tris Gel (NP0336, ThermoFisher). Gels were run for 1 hour and 15 min at 120 V in with NuPage MES Buffer (NP0002, ThermoFisher). Proteins were then transferred onto polyvinylidene difluoride membrane (IPVH15150, MilliporeSigma) for 1 hour at 30 V in NuPage Transfer Buffer (NP0006, ThermoFisher). Membranes were checked for successful and adequate transfer with a Ponceau S stain (P7170, Millipore-Sigma). Primary antibodies were diluted 1:1,000 (1 ug/mL) in blocking buffer (927-40000, Licor), and secondary antibodies were diluted to 1:10,000 (100 ng/mL) in 2.5% BSA (0332, VWR) in Tris buffered saline tween. All antibody incubations were performed overnight at 4°C. The primary antibodies used were rabbit antibody #9102L (CST) for ERK1/2, rabbit antibody #4370 (CST) for pERK, mouse antibody #31F12 (CST) for ALK in its kinase domain, rabbit antibody #3341S (CST) for pALK, and rabbit antibody #13E5 (CST) for βActin. Anti-rabbit IgG #7074S (CST) and anti-mouse IgG #7076S (CST) secondary antibodies were used to detect the primary antibodies. Chemiluminescent signal was generated by SuperSignal West Pico PLUS Chemiluminescence substrate (345777, ThermoFisher), and detected on a BioRad ChemiDoc.
Imaging System. Membranes were stripped (21059, ThermoFisher) for 30 min, and re-blocked for 1 hour at room temperature.

QUANTIFICATION AND STATISTICAL ANALYSIS
All statistical tests accompany a description of the test in the figure legends and the corresponding STAR Methods Section. Data is reported as mean with standard error of the mean (SEM) or as boxplots when noted, with boxplots designating the 25th-75th percentiles, median value, minimum value, and maximum value.