ERK Mutations Confer Resistance to Mitogen- Activated Protein Kinase Pathway Inhibitors

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Abstract

The use of targeted therapeutics directed against BRAFV600-mutant metastatic melanoma improves progression-free survival in many patients; however, acquired drug resistance remains a major medical challenge. By far, the most common clinical resistance mechanism involves reactivation of the MAPK (RAF/MEK/ERK) pathway by a variety of mechanisms. Thus, targeting ERK itself has emerged as an attractive therapeutic concept, and several ERK inhibitors have entered clinical trials. We sought to preemptively determine mutations in ERK1/2 that confer resistance to either ERK inhibitors or combined RAF/MEK inhibition in BRAFV600-mutant melanoma. Using a random mutagenesis screen, we identified multiple point mutations in ERK1 (MAPK3) and ERK2 (MAPK1) that could confer resistance to ERK or RAF/MEK inhibitors. ERK inhibitor–resistant alleles were sensitive to RAF/MEK inhibitors and vice versa, suggesting that the future development of alternating RAF/MEK and ERK inhibitor regimens might help circumvent resistance to these agents. Cancer Res; 74(23): 1–11. ©2014 AACR.

Introduction

BRAF is mutated in approximately 50% of melanomas, resulting in constitutive activation of the MAPK pathway ((B)RAF–MEK–ERK; ref. 1). Inhibitors targeting RAF and MEK (a key downstream effector kinase in the pathway) improve the survival of patients with BRAF-mutant melanoma, and are thus used clinically for this indication (2–4). In particular, combined RAF/MEK inhibition appears to improve progression-free survival compared with RAF or MEK monotherapy (2, 5). Despite these therapeutic successes, nearly all patients develop progressive disease that shows resistance to these agents (2, 6).

Multiple studies describing mechanisms of resistance to RAF/MEK inhibition have been published. Reactivation of the MAPK pathway through a variety of means, including alternatively spliced BRAF, NRAS or MEK1/2 mutations, MAP3K8 upregulation, or receptor tyrosine kinase signaling, is the most common clinical drug resistance mechanism (7–11). Because all of these alterations converge on sustained activation of ERK, the clinical development of small-molecule ERK inhibitors is of considerable interest. More generally, ERK signaling represents a key downstream effector of RAS mutations in many cancer types, suggesting that ERK inhibitors might eventually have multiple indications in oncology.

ERK1 and ERK2 proteins are 84% identical and comprise the only known substrates of MEK. MEK activity on ERK results in dual phosphorylation on the Thr-Glu-Tyr motif of ERK1T202/Y204 and ERK2T185/Y187 that fully activates ERK kinase activity. Conversely, ERKs are negatively regulated by dephosphorylation, which is accomplished by dual specificity phosphatases (DUSP; refs. 12, 13). DUSPs also become induced by ERK signaling, creating a negative feedback loop (14). Other downstream ERK effectors include kinases such as RSK and MSK, cytoskeletal molecules, nucleoporins, and transcription factors (e.g., c-FOS, ELK-1, ETS-1, and MITF). Oncogenic dysregulation of this ERK program may profoundly impact cell proliferation and survival (15).

Several small-molecule ERK inhibitors have entered clinical trials (16, 17). Given the importance of secondary kinase mutations as resistance mechanisms in many oncogene-driven cancers, we wished to discover mutations in ERK that confer resistance to ERK inhibitors. We reasoned that at least some such ERK mutations might also confer resistance to RAF/MEK inhibitors. (In this regard, these studies might also identify constitutively active variants of ERK, which have remained elusive despite intensive study.) Random mutagenesis screens have successfully identified clinically relevant resistance alleles (18, 19) in multiple kinase oncogene-driven malignancies. Thus, we employed random mutagenesis in BRAF-mutant melanoma cells to identify mutations in ERK1 or ERK2 that could confer resistance to MAPK inhibitors.

Materials and Methods

Cell lines and reagents

A375, 293T, SKMEL-19, and WM266.4 cells were grown in DMEM with 10% FBS. A375 and 293T cells were acquired from the ATCC. WM266.4 cells were acquired through the Cancer Cell Line Encyclopedia (20). SKMEL-19 cells were a gift from N. Rosen (Memorial Sloan Kettering Cancer Center, New York,
Random mutagenesis

Mutagenesis screens were performed as described previously (19) with additional detail in Supplementary Materials and Methods.

Drug screens with mutagenized libraries

A375 cells (8 × 10^5) expressing tet-inducible GFP, wild-type ERK1/ERK2, or mutant library ERK1/ERK2 were plated in T150 flasks with VX-11e (2 μmol/L), trametinib (3 nmol/L), or trametinib + dabrafenib (1 nmol/L + 10 nmol/L) and doxycycline (DOX; 1 μg/mL) for 2 to 4 weeks until resistant cells emerged. At that time, genomic DNA (gDNA) was isolated (DNeasy kit; Qiagen). Exogenous ERK1/2 was amplified by the PCR using vector-specific primers (Supplementary Materials and Methods) and AccuPrime PFX supermix (Life Technologies). Individual colonies were collected and transferred into 96-well plates and expanded. Genomic DNA was isolated, and PCR was performed as above, then analyzed by Sanger sequencing.

Library generation and massively parallel sequencing

Massively parallel sequencing of PCR products from mutagenesis screens was performed as described previously (19). Additional detail is provided in Supplementary Materials and Methods.

Viability assays

A375, WM266.4, or SKMEL-19 cells were plated in 96-well plates, with six replicates for each drug, VX-11e (2 μmol/L), trametinib (3 nmol/L), dabrafenib (50 nmol/L), or SCH772984 (1 μmol/L or 0.5 μmol/L) were added 24 hours later with or without DOX (1 μg/mL). Drugs were diluted 1:1,000 from stock concentrations in DMSO. Viability was analyzed by CellTitre 96 AQueous One Solution Cell Proliferation assay (Promega). The raw absorbance values were normalized to vehicle-treated controls after background subtraction and graphed as percent viability, with vehicle-treated cells at 100%. Heat maps were created from normalized viability values using GENE-E (21).

Western blotting

A375 cells were exposed to DOX with or without VX-11e (2 μmol/L), or trametinib (3 nmol/L), or trametinib (3 nmol/L), then harvested in RIPA buffer, and run on SDS-PAGE. Proteins were analyzed using the following antibodies: phosphoERK1 T202/Y204/ERK2 T185/Y187, ERK1/2, phosphoELK-1 S383 (Santa Cruz Biotechnology), phosphoERK2 T204, RSK, tubulin, DUSP6 (Cell Signaling Technology), cyclin D1 (Thermo Scientific), and V5 epitope (Abcam). Relative expression was calculated by dividing the relative densities of each phosphoELK-1 band by total ERK (or V5) using Image J. Values were normalized to untreated wild-type, which was set at 1.

Immunoprecipitation kinase assays

Described in detail in Supplementary Materials and Methods.

Site-directed mutagenesis

Primers were designed using the QuickChange Site-Directed Mutagenesis Protocol (Agilent) and are listed in Supplementary Materials and Methods. PCR was performed using iProof DNA polymerase and GC buffer using primers specific for each mutation (Bio-Rad). Mutations were confirmed by Sanger sequencing using vector-specific primers.

Aside from the mutagenesis screens, all experiments were repeated at least three times, with representative graphs or blots shown. Error bars represent SDs.

Results

ERK1/ERK2 random mutagenesis

To discover mutations in ERK1 or ERK2 that may confer resistance to MAPK inhibitors, a six-part random mutagenesis screen was performed. Here, we mutagenized ERK1 and ERK2 within the pDONR223 vector using a mutator strain of E. coli (22). The resulting mutagenized cDNA library was transferred into a DOX-inducible vector (pCW57.1), which was infected into A375 (BRAF-mutant) melanoma cells that are sensitive to MAP kinase pathway inhibitors. These cells were cultured in the presence of either the ERK inhibitor VX-11e (23), the MEK inhibitor trametinib (GSK1120212), or the combination of trametinib and the RAF inhibitor, dabrafenib (GSK2118436), in the presence of DOX. In all cases, drug-resistant A375 daughter populations emerged within 2 to 4 weeks. Genomic DNA was isolated, and ERK1 or ERK2 cDNA was amplified by the PCR followed by massively parallel sequencing (see Materials and Methods). In parallel experiments, a series of VX-11e–resistant ERK2 colonies were isolated, and ERK2 cDNA was analyzed by Sanger sequencing.

Overall, we identified 33 putative resistance variants spanning 28 amino acids in ERK1, and 24 substitutions affecting 20 amino acids in ERK2 (above a 5% massively parallel sequencing false discovery rate; see Supplementary Materials and Methods; Fig. 1; Supplementary Fig. S1A–F; and Supplementary Table S1). Sanger sequencing of 23 VX-11e–resistant ERK2 colonies identified seven ERK2 mutations in five amino acids, two of which were also identified in the mutagenesis screens. We have no evidence that these mutagenesis screens were saturating; therefore, additional candidate ERK1/2 resistance mutations may remain to be identified. Nonetheless, our screening results suggested that a diverse range of ERK mutations might be associated with resistance to these inhibitors.

We observed five analogous resistance alleles between ERK1 and ERK2 in cells resistant to the ERK inhibitor in both the screen and in isolated colonies (ERK1 Y53H/ERK2 Y361H/N, ERK1 G54A/ERK2 G375S, ERK1 P75L/ERK2 P58L, ERK1 Y181C/ ERK2 Y186N, and ERK1 C82Y/ERK2 C65Y; Fig. 1C), and one analogous residue in MEK inhibitor–resistant cells (ERK1 Y144H and ERK2 Y131H/N/F/C/S; Fig. 1D), suggesting convergent resistance mechanisms between the ERK isoforms. There was relatively little overlap between the ERK inhibitor screens and the RAF/MEK inhibitor screens (three overlapping
alleles of 67 total significant nucleotide alterations), raising
the possibility that mutations identified through RAF/MEK
inhibitor treatment might be resistant to one or both of
these agents, but not to ERK inhibitors (and vice versa).
Several ERK codons had multiple distinct amino acid sub-
stitutions (these include ERK1Y53C/H, ERK1G54A/S,
ERK2Y36N/H, and ERK2P58L/S/T), suggesting the importance
of these residues for inhibition by ATP-competitive ERK
inhibitors.

In general, candidate ERK1/2 resistance mutations arising in
the ERK inhibitor resistance screen clustered within the ATP/
drug binding pocket, suggesting that these variants might
interfere with drug binding. In contrast, the RAF/MEK inhib-
itor resistance alleles were distributed throughout the ERK
proteins, although some mutations clustered in the αC-helix
and the common docking domain. The ERK αC-helix under-
goes a conformational change when phosphorylated that is
necessary for kinase activation, whereas the common docking
domain represents one of the ERK substrate binding domains.
These observations suggested that ERK resistance alleles that
emerged under RAF/MEK inhibition might facilitate kinase
activation. In addition, several of these alterations have been
previously identified through DNA sequencing efforts. Altera-
tions at ERK1R84, ERK1G186, ERK2D321, and ERK2E322 are listed
in the COSMIC database (24), and ERK2E322K is recurrently
mutated in cervical and head and neck cancers (25, 26);
however, none of these variants have been linked to drug
resistance.

ERK1 and ERK2 mutations confer resistance to ERK
inhibitors by interfering with drug binding

We first characterized ERK1/2 mutations that arose in the
setting of ERK inhibition. To verify their resistance effects, we
engineered mutations into the DOX-inducible vector by site-
directed mutagenesis. The resulting constructs were stably introduced into A375 cells and evaluated for growth in the presence of VX-11e. In total, we tested 20 candidate DOX-inducible ERK1/2 resistance alleles in cell growth inhibition assays, of which 16 conferred ≥4-fold increased cell viability compared with cells cultured without DOX (Fig. 2A; Supplementary Table S1; Supplementary Fig. S1G). In these experiments, wild-type ERK1 and ERK2 conferred a minor growth advantage (1.5-fold) in the presence of the ERK inhibitor, whereas kinase-dead ERK1 and ERK2 (27) and GFP had no effect (Fig. 2A).

To determine whether these ERK1 and ERK2 resistance alleles might confer cross-resistance to other ERK inhibitors, we also performed cell growth inhibition assays using the ERK inhibitor SCH772984. For ERK1, we chose four alleles that conferred growth in the presence of the VX-11e, three of which conferred maximal growth (>90%) and one that conferred moderate resistance (>75%). We also included the three ERK2 alleles identified by the random mutagenesis screen. Indeed, these alleles were also resistant to SCH772984, as measured by >4-fold increases in viability following mutant allele expression. One allele, ERK1Y537F, showed only a 2.5-fold increase in viability at 1 μmol/L SCH772984, but conferred a 5-fold growth advantage at 0.5 μmol/L SCH772984, similar to the other ERK1 alleles (Fig. 2B). Conceivably, this difference may relate to compound-specific differences in ERK binding associated with this particular residue.

Next, we examined downstream ERK signaling from ten validated alleles upon DOX induction. As a control, we included wild-type ERK1 and ERK2, both of which produced robust RSK phosphorylation and elevated DUSP6 expression in this setting (Supplementary Fig. S2). In contrast, kinase-dead ERK1 and ERK2 showed no increased RSK phosphorylation, but did induce DUSP6 expression, suggesting a possible kinase-independent function (Supplementary Fig. S2). In the absence of ERK inhibitor, all validated ERK resistance alleles examined augmented RSK phosphorylation, DUSP6 expression, or both (Supplementary Fig. S2).

Although the ERK inhibitor–resistant ERK alleles conferred growth advantage in the presence of 11e, some alleles produced discordant effects on ERK signaling. For example, both ERK1G54A and the analogous variant ERK2G37S had minimal discordant effects on ERK signaling. For example, both growth advantage in the presence of 11e, some alleles produced controls. In addition, ERK signaling was maintained in the presence of drug compared with controls. In addition, ERK signaling was maintained in the presence of SCH772984 (Supplementary Fig. S3A). In contrast to VX-11e, which does not alter phosphorylation of ERK (Fig. 2), SCH772984 decreased phosphorylation of wild-type ERK1/2, while having no effect on phosphorylation of the ERK2-resistant alleles (Supplementary Fig. S3A). Thus, these resistance alleles can maintain ERK signaling in the presence of an ERK inhibitor.

To determine whether the intrinsic kinase activity of the ERK1/2 alleles was maintained with inhibition of ERK, we examined the kinase activity of the ERK alleles via phosphorylation of an Elk-1 peptide using immunoprecipitation (IP) kinase assays and immunoblotting (see Materials and Methods). As expected, VX-11e blocked wild-type ERK1- and ERK2-mediated ELK-1 phosphorylation. In contrast, the ERK resistance alleles showed persistent ERK kinase activity in the presence of either VX-11e (Fig. 3A and B) or SCH772984 (Supplementary Fig. S3B). As expected, kinase-dead ERK1/2 did not phosphorylate the ELK-1 peptide (Fig. 3A and B). Thus, validated ERK resistance alleles maintained kinase activity even with concomitant ERK inhibitor exposure.

Finally, we mapped the validated ERK1 and ERK2 alleles within the three-dimensional crystal structure of ERK2 bound to a structurally similar ERK inhibitor (VX-9a, PDB:3I60; ref. 23). The resistance alleles lie in close proximity to the ATP/drug binding pocket, albeit within distinct functional domains (Fig. 2D). In particular, they occupy the glycine-rich loop (ERK1A148N, ERKY153H, ERK1G154A, ERK2Y56H, and ERK2T37S), an interval between the β-strand and α-helix (ERK1S74G, ERK1P75L, and ERK2P58L), the α-helix itself (ERK1Y81C, ERK1K92Y, ERK2Y64N, and ERK2G37S), and the activation loop (ERK1G186D). Presumably, these substitutions interfere with engagement of the ATP binding cleft by ERK inhibitors.

**ERK mutations that confer resistance to RAF/MEK inhibition preserve ERK activity despite MEK inhibition**

Next, we validated a series of candidate ERK resistance alleles that emerged from the RAF/MEK inhibitor mutagenesis screens. After cloning and expressing these mutations as described above, we performed cell growth inhibition assays using the MEK inhibitor trametinib and the RAF inhibitor dabrafenib, either alone or in combination. Of 17 mutated alleles examined, 10 were confirmed as resistant to each agent and the combination, conferring >2.5-fold growth advantage in the presence of the inhibitor(s) compared with controls (Fig. 4A; Supplementary Table S2). Of the 7 alleles that failed validation, at least two may have been passenger mutations that co-occurred with bona fide resistance alleles during the mutagenesis procedure (See Supplementary Materials and Methods and Supplementary Fig. S4). Because we observed robust growth advantage for two ERK1 alleles (ERK1A206V and ERK1S219F), we engineered the analogous mutations within ERK2 (because these were not observed in the primary screen). The resulting alleles (ERK2A192V and ERK2S195F) also conferred resistance to RAF/MEK inhibition (Fig. 4A).

The observation that validated ERK1/2 alleles conferred cross-resistance to all RAF/MEK inhibitor conditions raised the possibility that in contrast to the ERK inhibitor resistance mutations (which may primarily interfere with inhibitor binding), these might represent activating events. This notion was...
Figure 2. Pharmacologic and biochemical validation of ERK resistance alleles from ERK inhibitor screens. A, viability of A375 cells expressing tet-inducible ERK inhibitor resistance mutations exposed to VX-11e with or without DOX is shown as a heat map. GFP, wild-type ERK1/2, and kinase-dead ERK1K71R/ERK2K54R served as controls. Alleles are sorted by sensitivity to VX-11e + DOX. ERK2 alleles identified by sequencing drug-resistant colonies (black asterisks), or in both drug-resistant colonies and the ERK inhibitor random mutagenesis screen (red asterisks), are indicated. B, A375 cells expressing ERK inhibitor resistance mutations were treated as above except with SCH722984 (SCH'984). Viability was depicted as in A. Gray squares indicate mutations that were not tested. C, expression of phospho-ERK (pERK), phospho-RSK (pRSK), DUSP6, and cyclin D1 was examined in A375 cells expressing ERK resistance mutations or controls. Cells were treated with DOX in the presence or absence of VX-11e. D, structural localization of validated ERK1 and ERK2 mutations (spheres) mapped onto available ERK2 crystal structure data (cartoon) cocrySTALLized with VX-9a, a parent compound of VX-11e [PDB:3I60]. The glycine-rich loop (teal), the αC-helix (magenta), and the activation loop (blue) are indicated.
butteressed by the observation that the ERK2^{E322K} mutation—a known activating allele that occurs in cervical cancers (26)—also arose in this context. Moreover, ERK1/2^{A206V/A189V} and ERK1/2^{S219P/S202P} occur in the activation lip of ERK1/2 and in close proximity to the two phosphorylation residues that confer full ERK kinase activity when phosphorylated (ERK1^{Y148H}, ERK2^{Y131F}, ERK2^{D321G}, and ERK2^{E322K}). Also known as the D-site recruitment site, this ERK1/2^Y148_Y131 have not been described, but may in principle due to reduced DUSP binding (28, 29).

Alterations at in vivo ERK1/2^S219_P202 occur in the activation lip of ERK1/2 and dissection of isoform-specific may contain distinctive functional attributes, thus allowing highly homologous, “activating” variants of ERK1 and ERK2 to confer full ERK kinase activity when phosphorylated (ERK1^{Y148H}, ERK2^{Y131F}, ERK2^{D321G}, and ERK2^{E322K}). Also known as the D-site recruitment site, this docking domain represents the means by which ERK1/2 binds to some substrates. Mutations in the common docking domains (ERK1^{Y148H}, ERK2^{D321G}, and ERK2^{E322K}) confer elevated activity in vivo due to reduced DUSP binding (28, 29). Alterations at ERK1/2^{Y148/Y131} have not been described, but may in principle impair phosphatase and substrate binding considering their three-dimensional localization near the common docking domain (Supplementary Fig. S5B).

Although our mutagenesis screens were unlikely to be saturating, we only observed αC-helix resistance mutations in ERK1, whereas common docking domain alterations (ERK2^{D321G} and ERK2^{E322K}) were only found in ERK2 (Fig. 4B). Interestingly, the analogous common docking domain mutations in ERK1 (ERK1^{D338N} and ERK1^{E339K}) did not confer resistance to RAF/MEK inhibition in our hands (Supplementary Fig. S6). This suggests that although ERK1 and ERK2 are highly homologous, “activating” variants of ERK1 and ERK2 may contain distinctive functional attributes, thus allowing dissection of isomrf-specific functions in the future.

Next, we analyzed ERK signaling after expression of the validated alleles. As with the ERK inhibitor resistance mutations, DOX-induced expression of RAF/MEK inhibitor resistance alleles in both ERK1 and ERK2 activated ERK signaling, as evidenced by RSK or DUSP6 expression (Supplementary Fig. S2). The common docking domain mutations (e.g., ERK2^{D321G} and ERK2^{E322K}) did not induce RSK phosphorylation, consistent with published observations that such alterations may disrupt RSK binding (30). When these same ERK mutations were expressed in the presence of the MEK inhibitor, DUSP6 and cyclin D1 expression were maintained, whereas RSK phosphorylation was more variable (Fig. 4C and D). Thus, ERK mutations arising in the setting of RAF/MEK inhibition could maintain ERK signaling, even in the presence of upstream pathway inhibition.

To determine whether the intrinsic kinase activity of the RAF/MEK inhibitor–resistant ERK alleles was maintained despite MEK inhibitor exposure, we analyzed ERK kinase activity by IP kinase assays. Here again, all ERK resistance alleles examined maintained ERK kinase activity (as measured by ELK-1 peptide phosphorylation) in the presence of MEK inhibitor, even though MEK-dependent (and activating) ERK phosphorylation was decreased (Fig. 5, input). ELK-1 binds to both the F-site recruitment motif and the common docking domain within ERK. Therefore, mutations confined to the common docking domain (ERK1^{Y148H}/ERK2^{Y131F}, ERK1^{D338N}/ERK2^{E322K}) should not be sufficient to abolish ERK-ELK-1 interactions (30, 31). However, we observed diminished residual kinase activity in the setting of trametinib exposure with the common docking domain mutants compared with other alleles (Fig. 5). Altogether, these results suggest that ERK resistance alleles arising in the setting of RAF/MEK inhibition maintain sufficient kinase activity to rescue BRAF-mutant melanoma cell growth—even in the presence of upstream pathway inhibition.

**RAF/MEK inhibitor–derived ERK resistance alleles are sensitive to ERK inhibition and vice versa**

Previous studies have shown that several clinically validated genetic mechanisms of resistance to RAF and MEK inhibitors...
remain sensitive to ERK inhibition (Supplementary Fig. S7A; refs. 7, 8, 32). In our hands, RAF or MEK inhibitor–resistant BRAF-mutant melanoma cell populations generated by continuous exposure to tool compound inhibitors (PLX-4720 and AZD6244, respectively) following N-ethyl-N-nitrosourea (ENU) mutagenesis were also sensitive to ERK inhibition (Supplementary Fig. S7B). Because ERK resistance mutants arising from ERK inhibitor screens were largely distinct from those that emerged from the RAF/MEK inhibitor screens, we wished to determine whether either category of ERK mutations might confer cross-resistance to the alternative MAP kinase pathway inhibitor(s).

To investigate this question, we performed cell growth inhibition assays in the presence of RAF, MEK, or ERK inhibitors. In general, ERK resistance mutants arising from RAF/MEK inhibitor screens were sensitive to ERK inhibition. Conversely, the ERK mutations from ERK inhibitor screens remained sensitive to RAF and MEK inhibition (or both; Fig. 6A). The same was true in two additional cell lines (SKMEL-19 and WM266.4); however, the overall magnitude of resistance was lower in these lines (Fig. 6B and C and Supplementary Fig. S8A). Interestingly, one ERK mutation (ERK1C82Y) conferred resistance to all MAP kinase inhibitors tested—though the magnitude of the resistance phenotype was more modest [45% (RAF/MEK inhibitors) and 75% (ERK inhibitor) viability compared with untreated controls]. Overall, ERK resistance mutations arising from ERK inhibition remained vulnerable to combined RAF/MEK inhibition, and the converse was also true.

ERK1/2 overexpression reduces the viability of some BRAF-mutant melanoma cells

While performing these studies, we noticed that ERK overexpression was lethal to A375 cells in the absence of MAP
Overexpression of ERK1 G54A and ERK2 G37S, which exhibit immunoblotting, produced the greatest reductions in viability (Fig. 7B, ERK mutations arising in the setting of RAF/MEK inhibition (Fig. 3), also had no effect on viability (Fig. 7A). In contrast, reduced basal kinase activity compared with wild-type ERK (ERK1K71R/ERK2K54R) had no effect. Kinase activity of ERK mutants from RAF/MEK inhibitor resistance alleles or controls (wild-type and kinase-dead ERK1/2 (ERK1K71R/ERK2K54R) treated with trametinib (GSK'212). pERK and pRSK were monitored in the input cell lysates by immunoblotting.

**Discussion**

In melanoma, clinical resistance to RAF/MEK inhibition often involves reactivation of the MAPK pathway. Although multiple distinct mechanisms to achieve this effect have been described, ERK activation stands as their chief point of convergence. Thus, ERK represents an attractive therapeutic target in MAP kinase-driven cancers, and several ERK inhibitors have entered clinical trials. However, even if ERK inhibitors are successful clinically, resistance to these agents will likely emerge, as has been seen with most other kinase-based anticancer therapeutics.

Secondary mutations within the kinase target represent a common resistance mechanism for this class of therapeutics. The present study used random mutagenesis screens to identify mutations in ERK1 and ERK2 that confer resistance to tool compounds that bear structural similarity to ERK inhibitors in clinical trials. Multiple resistance mutations were identified in both ERK isoforms, and additional mutations could emerge in the future if such screens are carried out to saturation. ERK mutations we identified using the VX-11e compound were cross-resistant to SCH772984, suggesting that these resistance alleles might represent generalizable resistance mechanisms to ATP-competitive ERK inhibitors.

In addition to identifying mutations in ERK that confer resistance to ERK inhibitors, we identified several mutations in both ERK1 and ERK2 that retain robust ERK signaling despite pharmacologic RAF/MEK inhibition. Although we have not formally proved that these represent bona fide constitutive active ERK isoforms, the mutations may nonetheless prove useful for future biochemical studies.

One of the RAF/MEK inhibitor–resistant alleles detected in our screen, MAPK1 E322K (ERK2), was identified in cervical and head and neck carcinoma (25, 26). Although ERK2 E322K does not exhibit increased basal kinase activity in vitro, our data are consistent with the notion that this variant may remain activated in vivo because of reduced DUSP binding and hence loss of negative regulation (35), thereby conferring resistance to RAF/MEK inhibitors. Conceivably, treatment of patients containing this residue with RAF and/or MEK inhibitors may prove ineffective.

Our results raise the possibility that tumors with somatic ERK mutations arising in the setting of ERK inhibition might still be sensitive to RAF/MEK inhibition. Reciprocally, ERK mutations that confer resistance to RAF/MEK inhibition...
could remain sensitive to ERK inhibitors. If such mutations were to arise clinically, one could speculate that an alternative RAF/MEK and ERK dosing regimen might be attempted to achieve more prolonged disease control. Indeed, anecdotal reports of successful alternating targeted therapeutic regimens have emerged for other oncogene-driven cancers, such as EGFR-driven lung cancer. Of course, such speculation must be tempered by the fact that somatic ERK mutations have not yet been described as a clinical resistance mechanism to RAF/MEK inhibition in melanoma cells (of note, BRAF-amplification is concordant with recent observations that “acute” withdrawal of vemurafenib from a patient-derived xenograft tumor model has progressed to drug resistance produced a temporary tumor regression (34). This has also been termed the “drug holiday” effect (33). Presumably, the flare of MAP kinase signaling that ensues upon drug discontinuation is deleterious to melanoma cells (of note, BRAF amplification was a resistance mechanism in this study, making an acute signaling flare even more likely upon drug removal).

If this general phenomenon (e.g., lethality of acute drug withdrawal) were broadly relevant at some point during the treatment course, one could envision intermittent therapeutic strategies for RAF/MEK inhibition in melanoma. Indeed, discontinuous vemurafenib allowed for disease stabilization in continuous vemurafenib allowed for disease stabilization in
the aforementioned study by Das Thakur, Stuart and colleagues (34). There have also been reports of successful rechallenge with vemurafenib in patients that initially relapsed on the drug (36–38). Moreover, recent clinical trial results, in which vemurafenib was administered chronically, while the MEK inhibitor was given intermittently, suggest that intermittent MEK inhibition was no worse than chronic administration of both drugs, but MEK inhibitor toxicity may have been reduced by intermittent dosing (39). The ERK resistance alleles identified herein may provide a useful set of tools for detailed preclinical evaluation of intermittent ERK inhibition alternating with “acute” ERK hyperactivation as a novel therapeutic approach.

In summary, this study has identified a set of mutations in ERK1 and ERK2 that confer resistance to MAP kinase pathway inhibition at the level of RAF/MEK and ERK. These findings may anticipate future resistance mechanisms, enable new biochemical studies of ERK isoforms, and support preclinical studies of novel therapeutic strategies. They could also inform the development of “next-generation” ERK inhibitors that could be developed for melanoma and many other MAP kinase–driven cancers.

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Disclosure of Potential Conflicts of Interest
L.A. Garraway received a commercial research grant from Novartis. He is also a consultant/advisory board member for Novartis and Foundation Medicine. No potential conflicts of interest were disclosed by the other authors.

Authors’ Contributions
Conception and design: E.M. Goetz, L.A. Garraway Development of methodology: E.M. Goetz, N. Wagle, L.A. Garraway Acquisition of data (provided animals, acquired and managed patients, provided facilities, etc.): E.M. Goetz, D.J. Treacy Analysis and interpretation of data (e.g., statistical analysis, biostatistics, computational analysis): E.M. Goetz, M. Ghandi, N. Wagle, L.A. Garraway Writing, review, and/or revision of the manuscript: E.M. Goetz, M. Ghandi, N. Wagle, L.A. Garraway Study supervision: L.A. Garraway

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