Two Novel ALK Mutations Mediate Acquired Resistance to the Next-Generation ALK Inhibitor Alectinib

Ryohei Katayama1,2,3, Luc Friboulet1,2, Sumie Koike3, Elizabeth L. Lockerman1,2, Tahsin M. Khan1, Justin F. Gainor1,2, A. John Iafrate1,4, Kengo Takeuchi5, Makoto Taiji6, Yasushi Okuno7, Naoya Fujita3, Jeffrey A. Engelman1,2, and Alice T. Shaw1,2

Abstract

Purpose: The first-generation ALK tyrosine kinase inhibitor (TKI) crizotinib is a standard therapy for patients with ALK-rearranged non–small cell lung cancer (NSCLC). Several next-generation ALK-TKIs have entered the clinic and have shown promising activity in crizotinib-resistant patients. As patients still relapse even on these next-generation ALK-TKIs, we examined mechanisms of resistance to the next-generation ALK-TKI alectinib and potential strategies to overcome this resistance.

Experimental Design: We established a cell line model of alectinib resistance, and analyzed a resistant tumor specimen from a patient who had relapsed on alectinib. We developed Ba/F3 models harboring alectinib-resistant ALK mutations and evaluated the potency of other next-generation ALK-TKIs in these models. We tested the antitumor activity of the next-generation ALK-TKI ceritinib in the patient with acquired resistance to alectinib. To elucidate structure–activity relationships of ALK mutations, we performed computational thermodynamic simulation with MP-CAFE.

Results: We identified a novel V1180L gatekeeper mutation from the cell line model and a second novel I1171T mutation from the patient who developed resistance to alectinib. Both ALK mutations conferred resistance to alectinib as well as to crizotinib, but were sensitive to ceritinib and other next-generation ALK-TKIs. Treatment of the patient with ceritinib led to a marked response. Thermodynamics simulation suggests that both mutations lead to distinct structural alterations that decrease the binding affinity with alectinib.

Conclusions: We have identified two novel ALK mutations arising after alectinib exposure that are sensitive to other next-generation ALK-TKIs. The ability of ceritinib to overcome alectinib-resistance mutations suggests a potential role for sequential therapy with multiple next-generation ALK-TKIs. Clin Cancer Res; 20(22); 5686–96. ©2014 AACR.

Introduction

The EML4–ALK fusion oncogene was first reported in non–small cell lung cancer (NSCLC) in 2007 (1). Approximately 3% to 5% of NSCLC tumors harbor ALK rearrangements. In addition to EML4, other fusion partners such as TFG, KIF5B, and KLC1 have been identified in NSCLC (2–4). Fusion with any of these partner proteins is believed to mediate oligomerization of ALK, which then leads to constitutive activation of the tyrosine kinase (TK) and aberrant downstream signaling (5). As a result, ALK fusions function as potent “oncogenic drivers,” and cancers harboring these rearrangements are highly sensitive to ALK kinase inhibition.

Crizotinib is a potent small-molecule inhibitor targeting cMET, ALK, and ROS1 tyrosine kinases. On the basis of its activity in phase I, II, and III clinical trials, crizotinib is approved in many countries for the treatment of advanced, ALK-rearranged NSCLC (6–8). Although crizotinib often induces marked and durable responses, most patients will relapse within 1 to 2 years due to the development of resistance.

Multiple different mechanisms of resistance to crizotinib have been reported previously (9–14). In approximately one third of crizotinib-resistant cases, resistance is mediated by a genetic alteration in ALK itself, typically a missense mutation in the tyrosine kinase domain, though amplification of the ALK fusion gene has also been observed. In
Acquired Resistance to Alectinib in ALK-Positive NSCLC

Translational Relevance
The ALK tyrosine kinase inhibitor (TKI) crizotinib is a standard therapy for patients with ALK-rearranged non–small cell lung cancer (NSCLC). However, cancers invariably develop resistance to crizotinib, and this has spurred the clinical development of multiple next-generation ALK-TKIs, including alectinib (RO5424802/CH5424802) and ceritinib (LDK378). To determine how cancers develop resistance to alectinib, we established a cell line model of acquired resistance to alectinib and analyzed a resistant tumor specimen from a patient who had relapsed on alectinib. We identified two novel secondary ALK mutations, both of which were still sensitive to ceritinib in vitro. Importantly, we found that in the patient, ceritinib was able to overcome resistance to alectinib, suggesting a potential role for sequential therapy with multiple next-generation ALK-TKIs.

breakthrough Therapy Designation by the U.S. FDA, and ceritinib was recently approved by the U.S. FDA for ALK-positive patients with crizotinib-resistant or crizotinib-intolerant disease.

As with crizotinib, patients eventually develop resistance to next-generation ALK inhibitors. In this study, we have explored acquired resistance to alectinib in a cell line model and in a primary tumor specimen from an alectinib-refractory patient. We have identified two novel secondary mutations within the ALK-TK domain, both of which mediate resistance to alectinib: V1180L, which functions as a gatekeeper such as L1196M, and I1171T, which resides in the αC helix within the ALK-TK domain. The thermodynamic stability of different alectinib–ALK complexes suggests that both V1180L and I1171T substitutions cause resistance by decreasing the binding affinity of alectinib for the mutated kinases. Although cancer cells expressing either V1180L- or I1171T-mutated EML4–ALK are resistant to alectinib as well as crizotinib, they remain sensitive to other structurally distinct ALK inhibitors and to hsp90 inhibitor. Thus, two different therapeutic strategies may be effective in overcoming resistance to alectinib, including the use of an alternative next-generation ALK inhibitor in tumors with susceptible resistance mutations such as V1180L or I1171T.

Materials and Methods

Patients
The ALK-positive NSCLC patient with acquired alectinib resistance underwent biopsy of a resistant tumor in October 2012. Standard histopathology was performed to confirm the diagnosis of malignancy and the histologic subtype. Total nucleic acid was isolated as described. We also performed fluorescence in situ hybridization (FISH) and IHC studies as described below. The cell line was also established when we obtained sufficient tissue. The established ALK-positive patient-derived cell line has been previously tested for mutation status to confirm their authenticity. The electronic medical record was reviewed retrospectively to obtain clinical information under an Institutional Review Board (IRB)–approved protocol.

Reagents
Alectinib and LDK378 were purchased from ActiveBiochem, 17-AAG was from LC-Laboratories, NVP-TAE-684 and ASP3026 were from ChemieTek, and crizotinib was from ShangHai Biochempartner. AP26113 was from Selleck. Each compound was dissolved in DMSO for cell culture experiments.

Isolation of gDNA or total RNA preparation, sequencing of ALK fusion gene
Genomic DNA (gDNA) was isolated from cell pellets or fresh-frozen specimen with the DNaseasy Blood & Tissue Kit (QIAGEN) according to the manufacturer’s protocol. Total RNA was isolated from cell pellets or fresh-frozen specimen with an RNaseasy Mini Kit (QIagen) according to the manufacturer’s instructions. Each exon of ALK kinase domain
(exon 21 to 27) or ALK kinase domain was PCR-amplified from gDNA or cDNA synthesized from total RNA with Oligo dT using KOD Plus ver.2 (TOYOBO), and sequenced bidirectionally by Sanger sequencing.

**Fluorescence in situ hybridization**

Two-color, break-apart FISH to detect ALK rearrangements was performed using formalin-fixed paraffin-embedded tissues with in-house ALK FISH probes [ALK-5’ terminal (red)/ALK-3’ terminal (green)] made from BAC clones. Isolated or split signals indicate ALK rearrangement, while overlapping signals indicate a nonrearranged ALK gene. Images were captured with an Olympus BX51 fluorescent microscope equipped with a charge-coupled device camera (DP71; Olympus).

**Reagents and cell culture conditions**

The H3122 human NSCLC cell line was obtained as previously described (22). PC9 cells were kindly provided by Dr. Kazuto Nishio (Kinki University, Osaka, Japan) and have been previously characterized (23). HCC827 was kindly provided by Dr. Adi Gazdar (University of Texas Southwestern Medical Center, Dallas, TX) and has been previously characterized (24). H460 and A549 human NSCLC cells were obtained from American Type Culture Collection. Ba/F3, immortalized murine bone marrow–derived pro-B cells were obtained from the RIKEN BRC Cell Bank (RIKEN BioResource Center). H3122, H3122-derived alectinib-resistant cells (H3122 CHR-A1), HCC827, PC9, H460, and A549 cells were cultured in RPMI-1640 supplemented with 10% FBS (R-10). Human embryonic kidney 293FT cells were cultured in RPMI-1640 supplemented with 10% FBS (D-10). Ba/F3 cells, immortalized murine bone marrow–derived pro-B cells, were cultured in D-10 with or without 0.5 mg/mL IL3 (Invitrogen).

**Generation of H3122 CHR-A1 cells**

H3122 CHR-A1 cells were established in the same manner as H3122 CR1 (10). Briefly, H3122 cells were seeded at approximately 50% confluence in 15-cm dishes in R-10. Alectinib was added at a starting concentration of 10 nmol/L, and cells were maintained in fresh drug-containing medium changed every 72 to 96 hours. Cells were passaged once they reached confluence. After every two passages at a given concentration of drug, the concentration of alectinib was increased until a final concentration of 1 μmol/L was achieved. The resulting pool of resistant cells (designated H3122 CHR-A1) was maintained in R-10 with 1 μmol/L alectinib.

**Survival assays**

For 72-hour drug treatments, 2,000 to 3,000 cells were plated in replicates of three to six into 96-well plates. Following drug treatments, cells were incubated with the CellTiter-Glo Assay reagent (Promega) for 10 minutes and luminescence was measured using a Centro LB 960 microplate luminometer (Berthold Technologies). The data were graphically displayed using GraphPad Prism version 5.0 (GraphPad Software). IC50 value was determined by a nonlinear regression model with a sigmoidal dose response in GraphPad.

**Immunoblot analysis**

Lysates were prepared as described previously (ref. 11). Equal amounts of lysates were electrophoresed and immunoblotted with the antibodies against phospho-ALK (Tyr1604), ALK (C26G7), phospho-p42/44 ERK/MAPK (Thr202/Tyr204), p42/44 ERK/MAPK, phospho-Akt (Ser473; D9E), panAkt (C67E7), phospho-S6 Ribosomal Protein (Ser240/244, D68F8), S6 Ribosomal Protein (54D2; Cell Signaling Technology), GAPDH (6C5, Millipore), and β-actin (Sigma).

**Retroviral infection**

cDNAs encoding EML4–ALK variant 1, EML4–ALK variant 1 V1180L, or I1171T mutants were cloned into 1,520 bp retroviral expression vectors (pLenti), and virus was produced as previously described (ref. 10). After retroviral infection, Ba/F3 cells were selected in puromycin (1.0 μg/mL) for 2 weeks. For Ba/F3 cells infected by EML4–ALK variants, IL3 was withdrawn from the culture medium for at least 2 weeks before experiments.

**Statistical analysis**

All data are shown as mean ± SD. Statistical analysis was performed using a two-tailed Student t test. Significance was established for P < 0.05.

**Results**

**Generation and characterization of alectinib-resistant cells**

The EML4–ALK–expressing NSCLC cell line H3122 is more sensitive to alectinib than crizotinib, as reported previously (Supplementary Fig. S1; ref. 11). To explore mechanisms of alectinib resistance, we generated resistant H3122 cells by exposing the sensitive parental cells to increasing concentrations of alectinib for 7 months. The fully resistant cells, H3122 CHR-A1 cells, were maintained in 1 μmol/L of alectinib. H3122 CHR-A1 cells were as resistant to alectinib as cancer cell lines that do not harbor ALK rearrangement (Fig. 1A and B). In contrast to parental H3122 cells, H3122 CHR-A1 cells maintained ALK phosphorylation and downstream AKT and ERK phosphorylation in the presence of 300 nmol/L of alectinib (Fig. 1C).

As amplification of ALK fusion genes has been shown to mediate crizotinib resistance (10), we examined ALK gene copy number by FISH. Like the parental H3122 line, CHR-A1 cells showed no evidence of amplification of the ALK fusion gene (Fig. 1D).

We next examined the entire coding sequencing of EML4–ALK in both H3122 parental and CHR-A1–resistant cells. In the resistant cells, we detected a G to C substitution at nucleotide 3538 of EML4–ALK variant 1, which was not detected in the parental line (Fig. 1D). This 3538 G to C substitution results in a valine to leucine change at codon 1171.
V1180 within the ALK-TK domain. On the basis of structural modeling studies of ALK (25), V1180 is predicted to lie near the L1196 gatekeeper residue, which is mutated in a subset of cell lines and patient biopsies with acquired resistance to crizotinib (9–11, 14, 16). No other tyrosine kinase mutations were detected in resistant cells.

**Figure 1.** H3122 CHR-A1 cells are resistant to alectinib and harbor a V1180L mutation in the ALK kinase domain. A, cells were seeded in 96-well black plates and treated with increasing concentrations of alectinib for 72 hours. Cell survival was analyzed using the CellTiter-Glo Assay. Although H3122 cells showed high sensitivity to alectinib (red line), H3122 CHR-A1 cells (green line) were as insensitive to alectinib as non-ALK-rearranged cell lines. B, table summarizing IC50 values (nmol/L) of each cell line against alectinib. C, parental H3122 and CHR-A1 cells were treated with alectinib at the indicated concentrations for 6 hours. Cell extracts were immunoblotted to detect the indicated proteins. D, two-color FISH [ALK-5′ region (red)/ALK-3′ region (green)] analysis was performed on parental H3122 and CHR-A1 cells. Most of the cells harbor isolated or split green signals (arrowhead, EML4–ALK). Shown on the right are electrophoretograms of EML4–ALK cDNA from parental H3122 and CHR-A1 cells. The g3538c mutation within exon 23 results in a V1180L substitution that corresponds to a gatekeeper mutation in ALK.
Identification of a novel ALK resistance mutation in a crizotinib- and alectinib-refractory patient

To identify the mechanisms of alectinib resistance that develop in patients, we examined a resistant tumor sample from a patient with advanced ALK-positive NSCLC who had relapsed on alectinib. This patient had previously received three lines of chemotherapy and had initially responded to crizotinib, as indicated by an improvement in disease burden on CT scans. After 8 months of crizotinib therapy, the patient developed disease progression and was then treated with alectinib (300 mg twice daily). Restaging CT scans demonstrated a response to alectinib, but after 4 months, the patient again experienced disease progression, including worsening liver metastases. The patient was discontinued from alectinib, and underwent biopsy of a resistant liver lesion. Cytopathology of the tumor cells demonstrated adenocarcinoma histology, and ALK FISH analysis confirmed the presence of ALK rearrangement in the resistant tumor specimen. In addition, a cell line was established directly from the biopsy specimen that we designated MGH056-1.

To determine whether a secondary mutation within ALK might underlie the development of alectinib resistance in this patient, we extracted total nucleic acid from a flash-frozen specimen, and the entire tyrosine kinase domain of ALK was amplified from cDNA and sequenced. We identified a T to C missense mutation at codon 3512, leading to an I1171T amino acid substitution (Fig. 2A). The I1171 residue is located in the C helix. Mutation of I1171 has been reported previously in neuroblastoma (26), but not in ALK-rearranged NSCLC. The MGH056-1 cell line also harbored the I1171T mutation.

To determine whether MGH056-1 cells are resistant to ALK tyrosine kinase inhibitor (TKI) therapy, we compared the sensitivity of the parental H3122, H3122 CHR-A1, and MGH056-1 cells to crizotinib and alectinib. As shown in Fig. 2B, both H3122 CHR-A1 and MGH056-1 cells were similarly resistant to crizotinib in cell viability assays. With respect to alectinib, H3122 CHR-A1 cells were highly resistant, whereas the MGH056-1 cells displayed intermediate resistance (Fig. 2B). Furthermore, compared with parental H3122 cells, MGH 056-1 cells required substantially higher concentration of alectinib (or crizotinib) to suppress the phosphorylation of EML4–ALK harboring I1171T, consistent with greater resistance observed in the H3122 CHR-A1 cells compared with the MGH056-1 cells. We next examined the effect of alectinib on ALK phosphorylation in the presence or absence of each resistance mutation. Consistent with the cell survival assays and the immunoblotting results with H3122 CHR-A1 and MGH056-1 cells, alectinib was much less potent at suppressing the phosphorylation of EML4–ALK harboring V1180L and, to a lesser extent, I1171T, compared with WT EML4–ALK (Fig. 3B). Thus, in Ba/F3 cells, both the V1180L and I1171T mutations confer resistance to alectinib, with V1180L appearing to confer a higher degree of resistance than I1171T.

Structural analysis of resistance mutations by computational simulation with MP-CAFEE

To understand the structure–function relationship of the two alectinib-resistance mutations in the ALK kinase domain, we performed computational simulation analysis with MP-CAFEE. To evaluate the thermodynamic stability of each complex of alectinib with WT, I1171T-, and V1180L-mutated ALK, we performed molecular dynamics simulation and estimated the free energies of each complex using MP-CAFEE method (27). Supplementary Figure S2 shows the comparison of experimental IC50 values and calculated free energy values. A linear correlation between experimental and calculated values was observed. This result indicates that the molecular dynamic simulation and free energy estimation by MP-CAFEE could correctly predict the resistant mechanism of each resistant mutant. In the MP-CAFEE method, the free energy value is estimated by summation of Coulomb and van der Waals potential energies. Figure 3C shows the estimated interaction energies of alectinib with WT, I1171T, or V1180L ALK. Interestingly, the decrease in binding affinities of I1171T-alectinib and V1180L-alectinib, compared with WT ALK-alectinib, are caused by decreases in Coulomb and van der Waals interactions, respectively. As shown in Fig. 3D, the I1171T mutation distorts the C-helix, shifting the position of the glutamic acid at 1167 inferiorly. As the cyano group of alectinib forms a hydrogen bond with E1167, this downward shift may disrupt binding of alectinib to the I1171T mutant. In contrast, for the V1180L mutant, our modeling predicts that the methyl group of the leucine residue clashes with the multicyclic rings of alectinib (Fig. 3D). This steric interference likely decreases the van der Waals interaction between V1180L and alectinib. Taken together, these results suggest that for both I1171T and V1180L mutations, drug resistance is due to decreased binding affinity of alectinib for the mutant kinases.

ALK I1171T and V1180L mutations mediate resistance to alectinib in Ba/F3 models

To determine whether the I1171T and V1180L mutations are sufficient to cause resistance to alectinib, we engineered Ba/F3 cells expressing EML4–ALK harboring each mutation. In cell survival assays, Ba/F3 lines expressing the mutant EML4–ALK proteins were significantly less sensitive to alectinib than those expressing wild-type (WT) EML4–ALK (Fig. 3A). Ba/F3 cells with EML4–ALK V1180L were more resistant than Ba/F3 cells with EML4–ALK I1171T (Fig. 3A), consistent with greater resistance observed in the H3122 CHR-A1 cells compared with the MGH056-1 cells. We next examined the effect of alectinib on ALK phosphorylation in the presence or absence of each resistance mutation. Consistent with the cell survival assays and the immunoblotting results with H3122 CHR-A1 and MGH056-1 cells, alectinib was much less potent at suppressing the phosphorylation of EML4–ALK harboring V1180L and, to a lesser extent, I1171T, compared with WT EML4–ALK (Fig. 3B). Thus, in Ba/F3 cells, both the V1180L and I1171T mutations confer resistance to alectinib, with V1180L appearing to confer a higher degree of resistance than I1171T.

Structurally distinct next-generation ALK inhibitors overcome alectinib-resistant mutations

In addition to alectinib, several other ALK-TKIs are currently in clinical development. To determine whether these next-generation ALK inhibitors may have activity in alectinib-resistant cancers, we used the Ba/F3 cells expressing EML4–ALK harboring the I1171T or V1180L mutation.
controls, we also tested Ba/F3 cells expressing WT EML4–ALK and parental, IL3-dependent Ba/F3 cells. We focused on four ALK inhibitors: the tool compound NVP-TAE684 (28), and three drugs currently in clinical trials or clinically available for ALK-rearranged NSCLC, ceritinib (LDK378; refs. 17, 21), AP26113, and ASP3026. As shown in Fig. 4A, NVP-TAE684 demonstrated potent activity against Ba/F3 cells expressing V1180L EML4–ALK or the I1171T EML4–ALK. The clinically available ALK inhibitors ceritinib and AP26113 showed slightly different selectivity profiles against the different ALK resistance mutations. Ceritinib was highly active against V1180L, even more so than against WT EML4–ALK (Fig. 4A and Supplementary Fig. S3). The I1171T mutation was also sensitive to ceritinib, demonstrating a dose-response similar to WT EML4–ALK. In contrast, AP26113 and ASP3026 were similarly active against both V1180L-mutant and WT EML4–ALK, but AP26113 was a little less active against I1171T-mutated EML4–ALK, whereas ASP3026 was inactive against I1171T (Supplementary Figs. S3 and S4). The suppression of phospho-ALK by the different inhibitors across the various mutations was consistent with the potencies observed in these Ba/F3 cells.

Figure 2. Discovery of the ALK I1171T secondary mutation in a patient with acquired resistance to crizotinib and alectinib. A, secondary I1171T mutation was only detected in the post-alectinib-treated specimen. Shown are electrophoreograms of ALK kinase domain cDNA from the post-alectinib treatment. B, sensitivity of MGH056-1 cells to crizotinib or alectinib. MGH056-1 cells, parental H3122, and CHR-A1 cells were treated with indicated concentrations of crizotinib or alectinib. Cell survival was analyzed with the CellTiter-Glo Assay. C, MGH056-1 cells were exposed to increasing concentrations of crizotinib or alectinib for 6 hours. Cell lysates were immunoblotted to detect the indicated proteins.
Parental H3122 cells, as well as CHR-A1 cells harboring the V1180L EML4–ALK mutation. 684, crizotinib, and ceritinib) in alectinib-resistant H3122 mutations.

resistance to alectinib mediated by either I1171T or V1180L tors—ceritinib and AP26113—may be able to overcome suggest that at least two of the next-generation ALK inhibi- studies (Supplementary Figs. S3 and S5). These results suggest that at least two of the next-generation ALK inhibitors—ceritinib and AP26113—may be able to overcome resistance to alectinib mediated by either I1171T or V1180L mutations.

We next tested the efficacy of several ALK inhibitors (TAE-684, crizotinib, and ceritinib) in alectinib-resistant H3122 CHR-A1 cells harboring the V1180L EML4–ALK mutation. Parental H3122 cells, as well as KRAS- or EGFR-mutant cancer cell lines (A549, H460, PC-9, and HCC827), were used as controls. As shown in Fig. 4B, TAE-684 markedly suppressed cell growth in both sensitive H3122 and alectinib-resistant H3122 CHR-A1 cells, but had almost no effect on the viability of other, non–ALK-dependent cancer cell lines. TAE-684 suppressed ALK phosphorylation and downstream signaling (Fig. 4C, left), and induced apoptosis (Supplementary Fig. S6). In contrast, crizotinib treatment was significantly less effective against H3122CHR-A1 cells compared with H3122 parental cells (Fig. 4B and C). Like TAE-684, ceritinib demonstrated potent activity against both parental H3122 and alectinib-resistant H3122 CHR-A1 cells, decreasing cell growth (Fig. 4B), suppressing ALK phosphorylation (Fig. 4C), and inducing apoptosis (Supplementary Fig. S6).

Figure 3. Biochemical and structural basis of alectinib resistance mediated by ALK I1171T and V1180L mutations. A, Ba/F3 cells expressing WT, I1171T, or V1180L EML4–ALK were seeded in 96-well plates and treated with the indicated concentrations of alectinib for 72 hours. Cell viability was analyzed using the CellTiter-Glo Assay. B, inhibition of phospho-ALK by alectinib in Ba/F3 models. WT or mutated EML4–ALK-expressing Ba/F3 cells were exposed to increasing concentrations of alectinib for 2 hours. Cell lysates were immunoblotted to detect the indicated proteins. C, estimated interaction energies of the indicated complexes [alectinib with WT ALK (green), ALK I1171T (cyan), or ALK V1180L (magenta)] by MP-CAFEE methods are shown. In the MP-CAFEE methods, the free energy value ($\Delta G$) is estimated by summation of coulomb and van der Waals (vdw) potential energies. D, shown are the complex average structures in the equilibrated molecular dynamics simulation for WT ALK (green), ALK I1171T (cyan), and ALK V1180L (magenta).

The hsp90 inhibitor 17-AAG overcomes alectinib resistance in H3122 CHR-A1 cells

A number of hsp90 inhibitors have demonstrated clinical activity in ALK-rearranged NSCLC, including in 1 patient with acquired resistance to crizotinib (29, 30). We therefore tested whether the alectinib-resistant H3122 CHR-A1 cells might be sensitive to the hsp90 inhibitor 17-AAG. Compared with KRAS or EGFR-mutant cancer cell lines (A549, H460, PC-9, and HCC827), CHR-A1 cells were highly sensitive to 17-AAG treatment, and nearly as sensitive as parental H3122 cells (Supplementary Fig. S7). On the basis of immunoblotting, 17-AAG treatment reduced EML4–ALK protein levels in both parental H3122 and H3122CHR-A1 cells to similar extents, as well as downstream signaling. Thus, Hsp90 inhibition may represent an alternative therapeutic strategy for overcoming resistance to alectinib due to acquisition of a resistance mutation.

Ceritinib is active in a cell line model and a patient with alectinib resistance

We also tested the efficacy of different ALK inhibitors in the MGH056-1 cells. As shown in Fig. 5A, ceritinib, but not crizotinib or alectinib, markedly suppressed the cell growth of MGH056-1 cells. Ceritinib also suppressed ALK
Figure 4. Ceritinib overcomes alectinib resistance in Ba/F3 models and H3122 CHR-A1 cells. A, parental Ba/F3 cells and Ba/F3 cells expressing WT, I1171T, or V1180L EML4-ALK were seeded in 96-well plates and treated with the indicated concentrations of TAE-684 (left), crizotinib (center), or ceritinib (right) for 72 hours. Cell viability was analyzed using the CellTiter-Glo Assay. B, cancer cell lines, including parental H3122 and alectinib-resistant H3122 CHR-A1 cells, were seeded in 96-well plates and treated with increasing concentrations of TAE-684 (left), crizotinib (center), or ceritinib (right) for 72 hours. Cell viability was measured using the CellTiter-Glo Assay. Both parental H3122 cells and H3122 CHR-A1 cells showed marked sensitivity to TAE-684 and ceritinib. Non-ALK-rearranged cell lines (A549, H460, HCC827, and PC-9 cells) showed minimal growth inhibition when exposed to ALK inhibitors. C, suppression of ALK signaling by ALK inhibitors (TAE684, crizotinib, or ceritinib) in parental H3122 and CHR-A1 cells. Cells were exposed to increasing concentrations of TAE684, crizotinib, or ceritinib for 6 hours. Cell lysates were immunoblotted to detect the indicated proteins.
phosphorylation and downstream AKT and ERK phosphorylation to a greater extent than crizotinib and alectinib (Figs. 5B and 2C). Consistent with these results, treatment of patient MGH056 with ceritinib led to significant tumor regression (Fig. 5C), with a confirmed partial response lasting over 7 months. These preclinical and clinical results suggest that ceritinib may be effective in treating cancers that have become resistant to alectinib due to a secondary mutation such as I1171T or V1180L.

Discussion

ALK-rearranged NSCLC represents one of the newest oncogene-addiction paradigms in clinical oncology. As with other oncogene-addicted cancers, ALK-rearranged lung cancers are initially sensitive to the first-generation targeted agent, but eventually become resistant through a variety of different mechanisms. Remarkably, the vast majority of crizotinib-resistant tumors remain ALK-dependent and re-respond to more potent, next-generation ALK inhibitors such as alectinib and ceritinib (21, 31). However, despite their promising activity in early-phase studies, resistance to next-generation ALK inhibitors invariably develops and ultimately limits the clinical benefit afforded by these new agents.

In this study, we focused on acquired resistance to alectinib, one of the most advanced of the next-generation ALK inhibitors in the clinic. We identified two novel ALK mutations, V1180L in a cell line made resistant to alectinib, and I1171T in a tumor specimen from a patient who had relapsed on alectinib. Both mutations confer resistance to crizotinib as well as alectinib, and hence add to the growing list of secondary ALK mutations that can mediate crizotinib resistance. Mutation of the V1180 residue has not yet been reported in patients, but was observed at very low frequency in an in vitro mutagenesis screen for crizotinib-resistant mutants in EML4-ALK (25). On the basis of the crystal structure of ALK (32), V1180 resides at the back of the ATP pocket and likely makes direct contact with crizotinib and alectinib, similar to the L1196 gatekeeper residue. Our computational modeling revealed weaker binding of alectinib to the V1180L-mutant compared with WT ALK, supporting the notion that substitution of leucine for valine at this residue interferes with the ability of alectinib to bind effectively to the kinase.

The second alectinib-resistant mutation, I1171T, has not been previously reported in crizotinib-resistant patients, but mutation of this residue has been described in 2 patients with neuroblastoma (26). In neuroblastoma, mutation at
this residue results in an I1171N amino substitution and leads to activation of ALK, though the mutant kinase is unable to transform Ba/F3 cells (33). On the basis of our computational modeling studies, we predict that I1171T disrupts a hydrogen bond between alectinib and E1167 (Fig. 3D), destabilizing the complex of alectinib with the mutant kinase. Whether ALK I1171T is weakly oncogenic like I1171N and whether this could contribute to crizotinib resistance is unknown.

Compared with V1180L, which conferred high-level TKI resistance, the I1171T mutation was associated with intermediate resistance to alectinib in cell line studies. This mutation was discovered in a patient who had relapsed after 4 months of alectinib therapy. Before alectinib, the patient had received crizotinib with a response lasting 8 months. Although no tumor specimens were available before or after crizotinib therapy for genetic analyses, we suspect that the I1171T mutation emerged during the course of alectinib treatment, given his previous durable response to crizotinib followed by a re-response to alectinib. Of note, this patient was treated with alectinib at a dose of 300 mg twice daily. This represents half of the recommended phase II dose (RP2D) of alectinib established in a recent phase I study (NCT01588028; ref. 20). The relatively low drug exposure at this dose may have been a factor in selecting and/or expanding a clone with intermediate resistance to alectinib. As higher drug exposures are predicted at the RP2D, we speculate that patients treated at this dose could develop more highly resistant mutations such as V1180L.

Recently, several other mechanisms of resistance have been reported in patients who have relapsed on next-generation ALK inhibitors. The solvent front mutation G1202R, first discovered in a crizotinib-resistant tumor (11), appears to mediate high-level resistance to both alectinib and ceritinib. In one case, a patient who had relapsed on crizotinib was treated with alectinib at the RP2D and showed no evidence of response, consistent with intrinsic resistance (34). Molecular studies performed on a resistant specimen revealed the G1202R mutation. Similarly, in a series of 11 ceritinib-resistant tumors, three were found to harbor a new G1202R mutation and two had acquired a mutation at residue F1174 (17). Importantly, neither of these mutations was detectable in biopsies taken before ceritinib treatment. In the absence of a secondary ALK mutation, activation of alternative signaling pathways could also mediate resistance to alectinib. Indeed, in 1 patient who had relapsed on alectinib, amplification of cMET was reported in a resistant specimen, although it is unknown if it was driving resistance (35). In the case of our alectinib-resistant patient with I1171T, cMET was likely not driving the development of resistance, because ceritinib, which has no anti-cMET activity, was able to induce a durable response lasting over 7 months.

The observation that other structurally distinct, next-generation ALK inhibitors may overcome alectinib resistance in vitro and in vivo is clinically significant. Currently, nine next-generation ALK inhibitors have entered the clinic, with several showing potent activity in both crizotinib-naïve and crizotinib-resistant patients (19–21). One of the nine next-generation ALK inhibitors has already been approved by the U.S. FDA for the treatment of advanced, crizotinib-resistant, ALK-positive NSCLC. Until now, it was unknown whether patients could continue to derive benefit from ALK inhibition after failure of a next-generation ALK inhibitor. Our results suggest that patients may benefit from multiple, sequential ALK inhibitor therapies, depending on the underlying mechanism of resistance. In those cases with susceptible resistance mutations, such as V1180L and I1171T, ceritinib may be highly effective, even in a third-line, post-crizotinib, post-alectinib setting. However, in cases where resistance is mediated by a highly recalcitrant mutation, such as G1202R or by a completely different tyrosine kinase, ceritinib may not be helpful, and alternative treatment strategies, such as hsp90 inhibition or combinatorial therapeutics, may be required. Overall, these findings highlight the importance of serial biopsies to track the dynamic evolution of drug resistance, and to enable the rational selection of therapies most likely to be effective based on the underlying molecular alterations.

Disclosure of Potential Conflicts of Interest
J. Iafrate is a consultant/advisory board member for Chugai and Pfizer. K. Takeuchi is a consultant/advisory board member for and reports receiving a commercial research grant and speakers bureau honoraria from Chugai. J.A. Engelman reports receiving a commercial research grant from Novartis and is a consultant/advisory board member for Chugai, Genentech, Novartis, and Ventana. A.T. Shaw is a consultant/advisory board member for Ignyta, Genentech, Novartis, and Pfizer. No potential conflicts of interest were disclosed by the other authors.

Authors’ Contributions

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Ryohei Katayama, Luc Friboulet, Sumie Koike, et al.


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