TAS-120 Overcomes Resistance to ATP-Competitive FGFR Inhibitors in Patients with FGFR2 Fusion-Positive Intrahepatic Cholangiocarcinoma

ABSTRACT

ATP-competitive fibroblast growth factor receptor (FGFR) kinase inhibitors, including BGJ398 and Debio 1347, show antitumor activity in patients with intrahepatic cholangiocarcinoma (ICC) harboring activating FGFR2 gene fusions. Unfortunately, acquired resistance develops and is often associated with the emergence of secondary FGFR2 kinase domain mutations. Here, we report that the irreversible pan-FGFR inhibitor TAS-120 demonstrated efficacy in 4 patients with FGFR2 fusion-positive ICC who developed resistance to BGJ398 or Debio 1347. Examination of serial biopsies, circulating tumor DNA (ctDNA), and patient-derived ICC cells revealed that TAS-120 was active against multiple FGFR2 mutations conferring resistance to BGJ398 or Debio 1347. Functional assessment and modeling the clonal outgrowth of individual resistance mutations from polyclonal cell pools mirrored the resistance profiles observed clinically for each inhibitor. Our findings suggest that strategic sequencing of FGFR inhibitors, guided by serial biopsy and ctDNA analysis, may prolong the duration of benefit from FGFR inhibition in patients with FGFR2 fusion-positive ICC.

SIGNIFICANCE: ATP-competitive FGFR inhibitors (BGJ398, Debio 1347) show efficacy in FGFR2-altered ICC; however, acquired FGFR2 kinase domain mutations cause drug resistance and tumor progression. We demonstrate that the irreversible FGFR inhibitor TAS-120 provides clinical benefit in patients with resistance to BGJ398 or Debio 1347 and overcomes several FGFR2 mutations in ICC models.

INTRODUCTION

Intrahepatic cholangiocarcinoma (ICC) is an aggressive malignancy of the liver bile ducts with poor outcomes and rising incidence (1). Most patients are diagnosed with locally advanced or metastatic disease, precluding potentially curative resection. Standard-of-care palliative chemotherapy with gemcitabine and cisplatin offers these patients a median survival of less than one year (2). ICCs exhibit an array of genomic alterations of known oncogenic drivers and tumor suppressors, suggesting the potential of targeted therapies in subsets of patients (3–6). Recurrent genomic alterations that activate the FGFR pathway are present in approximately 20% of ICCs (3, 6–12). The most common alterations are chromosomal fusions consisting of FGFR2 exons 1 to 17, encoding the intact extracellular and kinase domains, fused in-frame to a 3′ partner that possesses a protein dimerization domain. The resulting chimeric FGFR2 proteins are constitutively active and promote proliferation or transformation of several cell types (6, 7, 9). The frequency of FGFR2 fusions in ICC is considerably higher than that reported for any other malignancy (ref. 13; data retrieved from http://www.cbioportal.org). Activating FGFR2 point mutations and amplification or overexpression of FGFR1–3 are also observed in subsets of patients with ICC (8, 14).

Multiple FGFR-selective inhibitors are being tested in clinical trials in patients with ICC harboring FGFR pathway alterations. These second-generation inhibitors represent an improvement over the early generation of multikinase inhibitors with activity against FGFR (e.g., dovitinib and ponatinib), which lack sufficient specificity and potency to effectively treat FGFR-driven tumors. The most clinically advanced FGFR-selective compound in cholangiocarcinoma is the ATP-competitive FGFR1–3 inhibitor BGJ398 (infigratinib), which demonstrated efficacy in a phase II trial of patients with advanced refractory cholangiocarcinoma harboring FGFR2 fusions, amplifications, or point mutations (14). The overall response rate (ORR) in this heavily pretreated patient population was 14.8%, and the disease control rate (DCR) was 75.4% (18.8% and 83.3%, respectively, for patients with FGFR2 fusions). A phase I dose-escalation trial using another ATP-competitive FGFR1–3 inhibitor, Debio 1347 (CHS183284; ref. 15), has also reported early evidence of antitumor activity in a few tumor types including ICC (16). However, rapid emergence of acquired resistance has frequently
been observed, with a 5.8-month median progression-free survival in the BGJ398 trial (14). We recently reported genomic characterization of pre- and post-progression cell-free circulating tumor DNA (cfDNA) and tumor biopsies in 3 patients with FGFR2 fusion–positive ICC treated with BGJ398; this study revealed the emergence of the FGFR2 V563F gatekeeper mutation at progression in all 3 patients, 2 of whom also had additional FGFR2 kinase domain mutations (17). Rapid autopsy in one patient revealed three different FGFR2 kinase domain mutations in spatially distinct metastases, highlighting the additional challenge of interlesional heterogeneity in addressing acquired resistance to an ATP-competitive FGFR inhibitor in ICC.

The third-generation, irreversible FGFR inhibitor TAS-120 covalently binds to a highly conserved P-loop cysteine residue in the ATP pocket of FGFR (C492 in the FGFR2-IIIb isoform; ref. 18). TAS-120 exhibits in vitro potency at low nanomolar concentrations and high specificity against wild-type FGFR1–4 as well as against some FGFR2 kinase domain mutations (19). Preliminary results from a phase 1 basket study of TAS-120 in patients with refractory advanced solid tumors showed an ORR of 25% and a DCR of 78.6% in 28 patients with ICC harboring FGFR2 fusions (20), including some patients who had received prior therapy with an ATP-competitive FGFR inhibitor.

Here, we report the results of clinical and translational studies of TAS-120 in the treatment of patients with FGFR2 fusion–positive ICC who progressed on BGJ398 or Debio 1347, including patients in whom secondary FGFR2 kinase mutations were detected just prior to TAS-120 initiation. We performed complementary studies investigating FGFR2-mediated signaling mechanisms in ICC models and determined the efficacy of these second- and third-generation FGFR inhibitors against clinically observed FGFR2 kinase domain mutations. Our findings reveal genotype–phenotype correlations for drug sensitivity that inform personalized targeted therapy in FGFR-activated ICC.

RESULTS

TAS-120 Provides Clinical Benefit in Patients with ICC with Acquired Resistance to BGJ398 or Debio 1347

Among 6 patients with advanced FGFR2 fusion–positive ICC who received care at our institution after progression on BGJ398 or Debio 1347 in clinical trials, 4 subsequently enrolled in the phase 1 trial of TAS-120 (NCT02052778) between November 2015 and November 2017. Each of the 4 patients showed benefit on TAS-120: 2 of these patients achieved a partial response and 2 achieved stable disease by RECIST v1.1 criteria (Fig. 1A) with a duration of benefit of 5.1 to 17.2 months. We highlight these patients to show proof of concept of an irreversible FGFR inhibitor overcoming acquired resistance to an ATP-competitive FGFR inhibitor in the clinic and to elucidate the potential molecular determinants of response for this observation. The patients’ clinical characteristics and FGFR2 gene alterations are summarized in Table 1A and B. No additional cancer-relevant genomic alterations were detected in the pretreatment biopsies, with the exception of copy-number increases of the FGFR1 and MYC loci in the biopsy from patient #3 (see Methods for specific genotyping assays used for the different samples).

Patient 1 is a 74-year-old female with recurrent FGFR2–SORBS1 fusion–positive ICC metastatic to her liver and lymph nodes. On third-line BGJ398 treatment, she achieved a maximum response of −68% followed by progression at approximately 12 months. cfDNA analysis at that time revealed two new FGFR2 kinase domain mutations, K660M and K715R [Fig. 1B; amino acids are numbered according to FGFR2-IIIb splice isoform (NM_00144913.1) because FGFR2 fusions in ICC are expressed in this context (21)]; the equivalent mutations in the one amino acid shorter IIc isoform are K659M and K714R. Biopsy of a single liver lesion at the time of progression showed no FGFR2 kinase domain mutations, suggesting that these mutations were subclonal or that other molecular mechanisms drove resistance in this lesion. The patient subsequently received TAS-120, which resulted in a maximum response of −77% and suppression of K660M and K715R below the level of detection in ctDNA. After nearly 16 months on TAS-120, she had progression in all liver lesions. A third FGFR2 mutation, the gatekeeper V565F, emerged in the ctDNA during the final months of TAS-120 treatment and was detected in a post-progression tumor biopsy.

Patient 2 is a 59-year-old female with FGFR2–ZMYM4 fusion–positive ICC who presented with a dominant 15-cm liver mass and metastases to her liver and lungs. She achieved a maximum response of −50% on second-line BGJ398 treatment. Scans at 6 months showed a mixed response with regression of the dominant mass and progression of satellite liver lesions. ctDNA analysis at that time revealed five mutations in the FGFR2 kinase domain (N550H, N550K, V565F, E566A, and K660M). Two of these mutations were observed in a tumor biopsy of a progressing satellite liver lesion obtained in parallel—V565F and K660M, as previously reported (17). Upon next-line TAS-120 treatment, she achieved stable disease with a best response of +8%. Progression occurred at approximately 7 months, with a mixed response consisting of rebound growth of a previously responsive lung lesion, stability of the dominant mass, and continued progression of the biopsied left lobe liver lesion. Although the spatial location of each mutation was unknown, this heterogeneous response to TAS-120 was reflected in ctDNA analysis where some mutations (N550H, K660M) became undetectable before eventually rebounding at the time of disease progression, and others stabilized (N550K, E566A) or increased (V565F) during therapy (Fig. 1C). A sixth FGFR2 mutation (V563L) emerged in ctDNA during TAS-120 therapy and was detected in a biopsy obtained upon disease progression.

Patient 3 is a 28-year-old male with Crohn disease and FGFR2–INA fusion–positive ICC who presented with a 5.4-cm liver mass concurrently with liver, lung, peritoneal, and lymph node metastases. He received second-line Debio 1347 treatment to a maximum response of −50% followed by disease progression at all sites at nearly 12 months. He then had two post-progression liver biopsies obtained 2.5 months apart on distinct liver lesions with intervening cytotoxic chemotherapy—the first revealed an FGFR2 H683L mutation (CCF = 0.23) and the second revealed three FGFR2 mutations (N550H; CCF = 0.093;
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**Figure 1.** TAS-120 is clinically effective in patients with FGFR2 fusion-positive ICC whose tumors acquired resistance to BGJ398 or Debio 1347.

A, Radiologic scans of patients 1–4 during the course of FGFR inhibitor therapy. B–D, Droplet digital PCR analysis of serial ctDNA samples from patients 1–3. Time periods of therapy with the specific FGFR inhibitors are indicated by shading. Mutations identified in tumor biopsies taken at the indicated times are presented at the bottom of each graph. CCF, cancer cell fraction; MAF, mutant allele frequency.
Table 1A. Clinical data of patients with FGFR2 fusion–positive cholangiocarcinoma receiving FGFR inhibitors

<table>
<thead>
<tr>
<th>Patient ID</th>
<th>FGFR2 fusion</th>
<th>First FGFR inhibitor</th>
<th>PFS (months)</th>
<th>BOR</th>
<th>Intervening therapies between 1st and 2nd FGFR inhibitor</th>
<th>Interval between 1st and 2nd FGFR inhibitor (months)</th>
<th>Second FGFR inhibitor</th>
<th>PFS (months)</th>
<th>BOR</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>FGFR2–SORBS1</td>
<td>BGJ398</td>
<td>12.6</td>
<td>−68.2%</td>
<td>None</td>
<td>1.2</td>
<td>TAS-120</td>
<td>15.8</td>
<td>−76.7%</td>
</tr>
<tr>
<td>2</td>
<td>FGFR2–ZMYM4</td>
<td>BGJ398</td>
<td>5.6</td>
<td>−49.9%</td>
<td>None</td>
<td>1.6</td>
<td>TAS-120</td>
<td>7.2</td>
<td>+8.3%</td>
</tr>
<tr>
<td>3</td>
<td>FGFR2–INA</td>
<td>Debio 1347</td>
<td>11.4</td>
<td>−49.5%</td>
<td>Gemcitabine/docetaxel, T11 palliative radiation</td>
<td>3.0</td>
<td>TAS-120</td>
<td>5.1</td>
<td>−22.1%</td>
</tr>
<tr>
<td>4</td>
<td>FGFR2–NRAP</td>
<td>BGJ398</td>
<td>7.1</td>
<td>−40.0%</td>
<td>T8 palliative radiation, pembrolizumab, resection of T8 metastasis, FOLFOX</td>
<td>7.4</td>
<td>TAS-120</td>
<td>17.2</td>
<td>−47.7%</td>
</tr>
</tbody>
</table>

Abbreviations: BOR, best overall response; PFS, progression-free survival.

Table 1B. FGFR2 mutations detected in ctDNA and tumor biopsies

<table>
<thead>
<tr>
<th>Patient ID</th>
<th>FGFR2 fusion</th>
<th>Post-progression BGJ398/Debio 1347, prior to TAS-120</th>
<th>Post-progression TAS-120</th>
</tr>
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<tr>
<td></td>
<td>ctDNA</td>
<td>Tumor biopsy</td>
<td>ctDNA</td>
</tr>
<tr>
<td>1</td>
<td>K660M, K715R</td>
<td>None detected</td>
<td>V565F*</td>
</tr>
<tr>
<td>4</td>
<td>None detected</td>
<td>No biopsy obtained</td>
<td>N550K</td>
</tr>
</tbody>
</table>

NOTE: All mutations were detected on CLIA-certified assays as a routine part of clinical care except those designated as a (detected on droplet digital PCR only) or b (detected on whole-exome sequencing only).
overcome specific resistance mechanisms. The patient eventually experienced growth of a single liver lesion at 17.2 months, and at that time, analysis of ctDNA and tumor biopsy demonstrated the emergence of FGFR2–KIAA1217 fusion (designated MG69; Supplementary Fig. S1G). Treatment of MG69 PDx tumors with TAS-120 (starting when the volume reached ~500 mm3) led to tumor regression and complete proliferative arrest, with prominent effects evident within three days and persisting over a 14-day course (Fig. 2E and F). Moreover, FGFR inhibition suppressed MEK/ERK and SHP2 activity, but not PI3K signaling, in MG69 PDx tumors (Fig. 2G). Thus, FGFR-activated ICC models are highly dependent on FGFR activity to sustain growth and maintain MEK/ERK signaling in vitro and in vivo.

TAS-120 Overcomes Multiple Clinically Observed FGFR Kinase Domain Mutations

To gain insight into the clinical landscape of secondary FGFR2 resistance mutations, we subsequently leveraged our FGFR-driven ICC cell line models to study the spectrum of FGFR2 kinase domain mutations emerging upon clinically acquired resistance to BGJ398 (N550K, V565F, E566A, K660M, and K715R), Debio 1347 (M538I, H683L), or both (N550H, L618V). We engineered these mutations into a retroviral vector expressing the FGFR2–PHGDH fusion, which we observed in a patient with ICC (see Methods). CCLP-1 cells were infected with retroviruses expressing the FGFR2–PHGDH fusion with a WT or mutant FGFR2 kinase domain or empty vector control. Of the mutations that arose in patients treated with BGJ398, N550K, L618V, and K660M resulted in prominent resistance to the drug in vitro (26- to 39-fold increase in IC50), with the V565F gatekeeper conferring the greatest level of resistance (327-fold; Fig. 3A, top; Supplementary Figs. S2A and S2B) show immunoblots for expression of the FGFR2 fusion, which was greatly increased expression of wild-type (WT) FGFR1 (IIIC isoform) as well as the FGFR2 ligand compared with the other cell lines analyzed (Supplementary Fig. S1C–S1E). Thus, biliary tract cancer cell lines with activating molecular alterations in the pathway are specifically dependent on FGFR signaling for growth in vitro.

FGFR signaling engages a series of downstream effectors in different normal and pathologic contexts (22). We examined the principal pathways controlled by FGFR signaling in the ICC13-7 and CCLP-1 cell lines by BGJ398 treatment and immunoblot analysis using phospho-specific antibodies. BGJ398 treatment (50 nmol/L) led to rapid inhibition of the MEK/ERK pathway as reflected by decreased pFRS2 (Y196), whereas minimal effects were observed on the PI3K pathway, as determined by pAKT (T308 and S473; Fig. 2C and D). Dose–response studies showed effective targeting of FGFR2 signaling and downstream inhibition of MEK/ERK at BGJ398 concentrations consistent with the cell viability IC50 data (Supplementary Fig. S1F); comparable data were seen for TAS-120 and Debio 1347. In many types of cancer, strong feedback mechanisms exist to restore MEK/ERK signaling in response to loss of upstream activators of the pathway (23), and these may limit benefit of certain therapeutics that involve MEK/ERK inhibition. Notably, the inhibition of MEK/ERK signaling was durable in both cell lines, with no evidence of pathway reactivation for up to 3 days for BGJ398 treatment (Fig. 2C and D).

To corroborate these results in vivo, we screened a collection of patient-derived xenograft (PDX) models of ICC for FGFR alterations and identified a model harboring a FGFR2–KIAA1217 fusion (designated MG69; Supplementary Fig. S1G). Treatment of MG69 PDx tumors with TAS-120 (starting when the volume reached ~500 mm3) led to tumor regression and complete proliferative arrest, with prominent effects evident within three days and persisting over a 14-day course (Fig. 2E and F). Moreover, FGFR inhibition suppressed MEK/ERK and SHP2 activity, but not PI3K signaling, in MG69 PDx tumors (Fig. 2G). Thus, FGFR-activated ICC models are highly dependent on FGFR activity to sustain growth and maintain MEK/ERK signaling in vitro and in vivo.
this drug was considerably less potent against FGFR signaling overall. The most pronounced resistance to Debio 1347 was seen with the N550K, L618V, and K660M mutations (12- to 17-fold increase in IC_{50}), whereas M538L, N550H, and E566A produced intermediate effects (4- to 8-fold). H683L had a modest effect, and K715R did not significantly affect responsiveness to the drug. Moreover, Debio 1347 was relatively effective against the V565F gatekeeper mutation (only 3-fold IC_{50} increase). Notably, TAS-120 showed only minimal or modest changes in activity against each of the acquired FGFR2 mutations (2- to 7-fold IC_{50} increase) with the exception of V565F (103-fold, Fig. 3A, bottom).

To extend these findings, we modeled clonal outgrowth during acquired resistance using a pooled clone system, in which all nine mutant clones were pooled at an initial abundance of 1% amidst a background of cells expressing the WT gatekeeper mutation (only 1% amongst a background of cells expressing the WT gatekeeper mutation).
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Figure 3. FGFR inhibitors have distinct activity profiles against secondary FGFR2 kinase mutations in ICC cell lines that correlate with clinical data. A–D, CCLP-1 cells were engineered by retroviral transduction to express the FGFR2–PHGDH fusion with a WT kinase domain or harboring the indicated mutations, or empty vector (EV). The fusions contain the FGFR2-IIIb splice isoform (NM_001144913.1), and the amino acids are numbered accordingly. A, Graphs of IC₅₀ measurements upon treatment with the indicated FGFR inhibitors. The measured IC₅₀ is also indicated numerically at the right along with the fold change in IC₅₀ of each cell line relative to cell lines expressing the WT fusion. Red shading highlights mutant conferring a greater than 10-fold increase in IC₅₀. B, Pooled CCLP-1 cell clones of all FGFR2 fusion variants were treated with BGJ398, Debio 1347, or TAS-120 at the indicated concentrations over 14 days. The individual clones were monitored using genomic DNA extracted at 14 days, using a droplet digital PCR assay specific to each mutation. Data are mean ± SEM of triplicate determinants of relative change in clonal abundance compared with the start of treatment and are generated from two independent experiments. C, Clonal pools as in B were treated sequentially with 50 nmol/L BGJ398 and 10 nmol/L TAS-120 to mimic the treatment course of patients. Cells were monitored at 7 and 14 days. Data are expressed in relative mutant allele frequency compared with the start of treatment. Data are mean ± SD of triplicate determinants of relative change in clonal abundance compared with the start of treatment and are generated from two independent experiments. D, Immunoblot of CCLP-1 cells expressing the different FGFR2–PHGDH alleles following treatment with the indicated inhibitor concentrations.
FGFR2 fusion (Fig. 3B; Supplementary Fig. S2C). Clonal pools were exposed to different concentrations of each FGFR inhibitor for 14 days, and the change in relative clonal abundance under the selective pressure of therapy was determined by droplet digital PCR (ddPCR; ref. 24). Outgrowth of K715R was not observed under any treatment condition, again suggesting that this mutation is not a functional resistance alteration. Notably, treatment with 50 nmol/L BGJ398 led to outgrowth of the resistance mutations observed in patients 1 and 2 (N550H, N550K, V565F, E566A, K660M) or previously observed (17) in the setting of BGJ398 resistance (e.g., L618V). In contrast, BGJ398 prevented the outgrowth of M538I, which was detected only in patient 3 who was treated with Debio 1347. Conversely, outgrowth of each of these mutations was observed upon treatment with 200 nmol/L Debio 1347, with the exception of V565F, consistent with the clinical course of patient 3. Finally, in the presence of 10 nmol/L TAS-120, only outgrowth of V565F, and to a lesser extent E566A and N550K, was observed. Of note, these were the same three mutations that did not decrease in abundance in patient 2 during TAS-120 therapy (Fig. 1C). Interestingly, higher concentrations of BGJ398 or TAS-120 were able to suppress outgrowth of all resistance mutations with the exception of V565F, highlighting the potential importance of drug exposure in suppressing resistant clones.

We next used the pooled clone system to model the effects of sequential FGFR inhibitor therapy, treating clonal pools sequentially with BGJ398 and then TAS-120 to mirror the clinical course of patients 1, 2, and 4 (Fig. 3C). Three of the mutations (K660M, N550H, and L618V) that emerged during BGJ398 treatment decreased in abundance when treatment was switched to TAS-120, consistent with our ddPCR analyses showing that TAS-120 led to decreases in the clonal abundance in K660M (in patients 1 and 2), N550H (in patient 2), and L618V (in patient 3). Conversely, V565F continued to increase and E566A and N550K levels stabilized, but failed to decrease upon TAS-120 treatment, similar to the clinical observations in ctDNA from patient 2. Thus, our model systems accurately mirrored the clonal dynamics of individual resistance mutations observed in ctDNA analysis from patients treated with TAS-120 after progression on BGJ398 or Debio 1347.

Signaling studies corroborated the cell viability findings. CCLP-1 cells expressing N550K, V565F, L618V, and K660M retained robust levels of pFRS2, pSHP2, pMEK, and pERK upon treatment with 50 nmol/L BGJ398, whereas signaling by the other mutants was inhibited partially (N550H, E566A) or strongly (H683L; Fig. 3D; Supplementary Fig. S2A). Treatment with TAS-120 (50 nmol/L) effectively suppressed signaling by all mutants except V565F. Finally, Debio 1347 (200 nmol/L) showed reduced potency against most of the mutants but remained relatively active against the V565F gatekeeper mutation compared with the other two inhibitors. All three inhibitors were effective against K715R. We confirmed our findings for a subset of the FGFR2 mutants in ICC13-7 cells via cell viability assays and immunoblot for signaling proteins (Supplementary Fig. S2D–S2F). Thus, we demonstrate in relevant in vitro ICC models that TAS-120 has activity against multiple secondary FGFR2 resistance mutations, which likely accounts for the benefit of TAS-120 seen in patients who previously progressed on BGJ398 or Debio 1347.

We conducted in silico structural modeling to gain insight into the molecular basis for the drug response profiles. TAS-120 docks into the ATP-binding pocket of FGFR2, with its acrylamide group forming a covalent bond with the sulfhydryl group of FGFR2-C492 (Fig. 4A and B). As with BGJ398 (17, 25), the dimethoxy phenyl group of TAS-120 is in close contact with the V565 gatekeeper residue. Accordingly, modeling data indicate that TAS-120 and BGJ398 resistance to V565F is due to steric clash preventing access of these drugs into the ATP-binding pocket. TAS-120 remains effective against V565I (19), likely due to less severe hindrance caused by the smaller isoleucine side chain. Debio 1347 lacks the bulky dimethoxy

Figure 4. Structural modeling of secondary FGFR2 kinase domain mutations with TAS-120. A, Model showing TAS-120 docked into the binding pocket of WT FGFR2. Amino acid residues corresponding to mutations conferring resistance to ATP-competitive FGFR inhibitors are highlighted. Structural representations were prepared using PyMOL. B, A close-up view of TAS-120 in ATP-binding pocket of WT FGFR2. The gatekeeper residue (V565) is in close proximity to the dimethoxy phenyl group of TAS-120.
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improved potency and specificity afforded by its covalent bind-
come FGFR kinase domain mutations, taking advantage of the
ATP-competitive inhibitors. TAS-120 was designed to over-
target’s kinase domain has limited the durability of benefit to
resistance associated with recurrent acquired mutations in the
subset of ICC and support the clinical utility of TAS-120
mutations in this setting. Collectively, these data high-
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pathway are specifically dependent on FGFR signaling for
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a molecular basis for these differences. Finally, we demon-
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pathway are specifically dependent on FGFR signaling for
growth and sustained SHP2/MEK/ERK signaling, and that
TAS-120 retains efficacy against FGFR2 kinase domain
mutations differs across agents and that
structural studies of these agents bound to FGFR provide
a molecular basis for these differences. Finally, we demon-
strate that preclinical ICC models with activation of the
pathway are specifically dependent on FGFR signaling for
growth and sustained SHP2/MEK/ERK signaling, and that
TAS-120 retains efficacy against FGFR2 kinase domain
mutations in this setting. Collectively, these data high-
light the FGFR-driven oncogene addiction of a defined
subset of ICC and support the clinical utility of TAS-120
in patients with acquired resistance to second-generation
FGFR inhibitors.

The efficacy seen across several early-phase clinical trials of
FGFR2 inhibitors in patients with advanced refractory ICC
(14, 28–30) represents a breakthrough in a disease with no
FDA-approved targeted therapies to date. However, as seen
with other tyrosine kinase inhibitors, the rapid emergence of
resistance associated with recurrent acquired mutations in the
target’s kinase domain has limited the durability of benefit to
ATP-competitive inhibitors. TAS-120 was designed to over-
come FGFR kinase domain mutations, taking advantage of the
improved potency and specificity afforded by its covalent bind-
mode and distinct orientation in the ATP-binding pocket
of FGFRs. This irreversible binding also permanently disables
FGFR2 enzymatic activity, thus providing the potential advan-
tage of extended pharmacodynamic duration without the need
for maintaining high drug levels. Covalent small-molecule kinase
inhibitors have demonstrated success in multiple malignancies
and have gained FDA approval in EGFR-mutant non–small
cell lung cancer (afatinib, osimertinib), ERBB2/HER2-mutant
breast cancer (neratinib), and various non-Hodgkin lymphomas
(ibrutinib; ref. 31).

We evaluated the efficacy of TAS-120, BGJ398, and Debio
1347 against the spectrum of nine clinically observed secondary
FGFR2 kinase domain mutations using ICC cell lines and
serial ctDNA analysis. The inhibitors exhibit unique in vitro
profiles, and the key findings included: (i) the mutations
that conferred greatest resistance to BGJ398 were N550K,
V565F, L618V, and K660M; (ii) the mutations that conferred
interest to Debio 1347 were N550K, L618V, and K660M; (iii) Debio 1347 largely retained activity against the
V565F gatekeeper mutation; and (iv) TAS-120 remained active
against all mutations except V565F, with modest reduction
in activity against E566A and N550K. Additional studies will
be required to determine the impact of these kinase domain
mutations on FGFR2 fusion protein stability and turnover
and also on the kinetics of signaling reactivation upon inhibi-
tor withdrawal. Moreover, it will be important to establish
the extent to which preexisting FGFR2 mutations affect time
to treatment failure, as observed in EGFR-mutant non–small
cell lung cancer (32).

The clonal dynamics observed with serial ctDNA analysis
may hold important implications for the clinical manage-
ment of patients with these resistance alterations. ddPCR
analysis of ctDNA showed that the mutation allele frequen-
cies for several FGFR2 mutations decreased upon TAS-120
treatment—K660M in patient 1, N550H and K660M in
patient 2, and L618V and H683L in patient 3—pointing to
the activity of TAS-120 against these alleles in the clinic. Sim-
ilarly, the sustained increase or emergence of V565F upon
TAS-120 in patients 1–3 is consistent with the in vitro resist-
ance studies, as is the lack of reduction in levels of E566A
and N550K. These data, if validated prospectively in larger
clinical cohorts, may provide support for a new paradigm
in which particular FGFR resistance mutations, detected in
serial ctDNA or tumor biopsies, could inform the choice of
subsequent FGFR-targeted therapies. The precedent for this
is emerging in advanced ALK fusion–positive NSCLC where
specific ALK kinase domain mutations that arise at the time
of crizotinib resistance suggest which second-generation
inhibitor should be used for the next-line treatment (33).
To guide such strategies in ICC, it will be important to also
establish the full spectrum of mechanisms of resistance to
TAS-120, including validating the functional impact of
V563L, which emerged upon progression on TAS-120 treat-
ment in patient 2. Notably, whereas resistance to other irre-
versible kinase inhibitors frequently arises due to mutations
of cysteine residues that mediate covalent binding (34, 35),
no mutations at the covalent binding site of TAS-120 (C492)
were identified in any of the 4 patients studied in the current
report.

A key challenge in the administration of pan-FGFR inhibi-
tors remains hyperphosphatemia-related dose holds and dose
reductions. One patient in this study had such a dose hold on TAS-120 (see Methods) and two had such dose holds on BQJ398. Hyperphosphatemia is a class effect of FGFR inhibitors arising from on-target pathway blockade of FGFR2–FGFR1 signaling in the renal tubule (22, 36). Notably, we found that clonal outgrowth of multiple mutations occurred more readily at lower concentrations of BQJ398 (Fig. 3B), highlighting that reduced drug exposure may play an important role in the emergence of resistance. Although further studies are needed to establish the impact of toxicity-related drug modifications on treatment response, the clinical experience highlights the importance of aggressive hyperphosphatemia management and the urgency to develop FGFR2-selective agents.

Although ctDNA analysis serves as a useful, noninvasive tool for diagnosing resistance and monitoring response to therapy, our studies also illustrate the importance of a comprehensive approach to studying drug resistance. In patient 3, the three FGFR2 mutations identified in the baseline TAS-120 liver biopsy sample went undetected by both targeted sequencing and ddPCR in the corresponding plasma sample, possibly reflecting low tumor shedding and emphasizing the complementary benefits of tumor biopsy and ctDNA analysis. In patient 1, two FGFR2 kinase domain mutations arose at the time of progression on BQJ398 but only one conferred resistance in functional modeling, underscoring the importance of functionally validating putative resistance mutations discovered on ctDNA or tumor tissue analysis.

In keeping with the clinical data, our preclinical studies demonstrate that ICC models with constitutive activation of FGFR signaling are strongly dependent on the pathway. FGFR inhibitor treatment of FGFR-driven ICC cell lines and a PDX model led to growth inhibition as well as potent and durable inactivation of the SHP2/MEK/ERK pathway. Unlike breast and gastric cancers with high-level FGFR2 amplification (37), FGFR signaling in these ICC models was not additionally coupled to the PI3K/akt pathway. These findings highlight the distinct signaling outputs of oncogenic FGFR signaling in different cancer contexts and point to SHP2/MEK/ERK as a likely major effector of the pathway in ICC. The data are consistent with SHP2/MEK/ERK activation being the principal effector of FGFR in normal physiology (38) and with the frequent presence of concurrent PIK3CA activating mutations with FGFR2 fusions in ICC, indicating the potential independence of these pathways (14). Although our studies suggested that FGFR2 fusions with different partners had comparable outputs, further studies will be required to fully address the potential differential impact of N-terminus partners on oncoprotein localization, inhibitor sensitivity, and downstream signaling targets, as reported for fusions involving the ROS1 receptor tyrosine kinase (39).

In summary, we demonstrate that strategically sequencing FGFR inhibitors can prolong the duration of benefit from FGFR inhibition in patients with FGFR2 fusion–positive ICC. Moreover, resistance profiles differ across agents and may evolve under the selective pressure of sequential FGFR inhibitors. However, FGFR2-independent resistance may emerge as an additional issue that can limit the potential of these next-generation inhibitors. As the clinical development of FGFR inhibitors pushes forward, it will be critical to incorporate tumor biopsies at baseline and progression and serial ctDNA analysis into clinical trials. These complementary analyses can facilitate our understanding of resistance and elucidate the biology underlying heterogeneous responses. Overall, this approach of tailoring FGFR-targeted strategies based upon resistance mechanisms detected in serial ctDNA and tumor biopsies may provide a new standard of care for this disease.

METHODS

Patients

Patients provided written informed consent to treatment on the phase II trial of BQJ398 (NCT02160041), phase I trial of Debio 1347 (NCT01948297), and phase I trial of TAS-120 (NCT02052778). On the phase II trial of BQJ398, the patients received 125 mg orally daily on days 1 to 21 of each 28-day cycle. On the phase I trial of Debio 1347, the patients received 110 mg orally daily continuously on days 1 to 28 of each 28-day cycle. On the phase I dose-escalation or expansion phase of the trial of TAS-120, the patients received 16 to 24 mg orally daily of TAS-120 continuously on days 1 to 21 of each 21-day cycle. All dose reductions and safety assessments were performed per protocol for all three trials. The TAS-120 dosing for each patient was as follows: patient #1 (16 mg); patient #2 (24 mg); patient #3 (20 mg); and patient #4 (16 mg). The timing, reason, and duration of the first dose hold for each patient is as follows: patient #1 (cycle 14, day 1; grade 3 motor neuropathy; 15 days); patient #2 (cycle 1, day 8; hyperphosphatemia; 7 days); patient #3 (cycle 3, day 1; grade 2 ALT and AST elevation; 7 days); and patient #4 (cycle 14, day 19; palliative radiation therapy; 12 days). The timing, reason, and dosing for the first and subsequent dose reductions for each patient is as follows: patient #1 (cycle 15, day 1, reduced to 12 mg orally every day for grade 4 creatine kinase elevation and remained at this dose until progression); patient #2 (cycle 3, day 1, reduced to 16 mg orally every day due to hyperphosphatemia and again on cycle 3, day 15 to 8 mg orally every day due to hyperphosphatemia; remained at this dose until progression); patient #3 (cycle 3, day 15, reduced to 16 mg orally every day due to grade 2 AST and ALT elevation, remained at this dose until progression); and patient #4 (no dose reductions).

CT and/or MRI scans were performed at baseline and every 6 to 9 weeks to assess for tumor response by RECIST version 1.1 criteria. The clinical and genomic data relating patient #2’s treatment prior to TAS-120 therapy have been reported previously (17). All biopsies, tumor specimens, and peripheral blood draws for plasma isolation were collected and analyzed in accordance with Institutional Review Board (IRB)–approved protocols, to which patients provided written informed consent, and all studies were conducted in accordance with the Declaration of Helsinki.

Reporting of FGFR2 Mutations

We report FGFR2 kinase domain mutations as the amino acid number of the FGFR2-IIIb splice isoform (NM_001144913.1), which is the primary isoform expressed in FGFR2 fusion–positive ICC (21), Commercial genotyping tests (e.g., Guardant) and our prior report (17) designate mutations using the one amino acid shorter FGFR2-IIIc isoform (NM_000141.4) as the reference sequence.

Targeted Sequencing of Tumor Tissue

DNA derived from the primary tumor and metastases were analyzed using deep-coverage targeted sequencing of key cancer-associated genes.
genes. Targeted sequencing was performed in the setting of clinical care via the SNaPshot platform, the FoundationOne platform, or MSK-IMPACT, and the methodology has been described previously (40, 41). Clinical genotyping platforms utilized on biopsies after progression on the ATP-competitive inhibitor were as follows: patients #1 and #2 (FoundationOne), patients #3 (MSK-IMPACT), and patient #4 (no biopsy performed). After progression on TAS-120, the assays used on biopsies were as follows: patient #1 (MGH SNaPshot), patient #2 (FoundationOne), patient #3 (no biopsy performed), and patient #4 (MGH SNaPshot). In patients #1, #2, and #4, no other cancer-relevant genomic alterations besides the FGFR2 fusion were detected in the treatment-naive tissue sample taken at initial diagnosis or at diagnosis of advanced disease; MGH SNaPshot was used for patients #1 and #2 and MSK-IMPACT was used for patients #3 and #4.

Solid Fusion Assay

Our internal tumor-profiling assay was performed on RNA extracted from formalin-fixed paraffin-embedded specimens as part of routine clinical care. The solid fusion assay is a targeted RNA-sequencing method of Anchored Multiplex PCR to detect FGFR2 fusions, and the methodology has been described previously (42). Mutational profiling was performed at the Clinical Laboratory Improvement Amendments (CLIA)-certified Translational Research Laboratory at the Massachusetts General Hospital (MGH) Cancer Center (Boston, MA).

ddPCR

DNA template (8 to 10 μL) was added to 10 μL of ddPCR Supermix for Probes (Bio-Rad) and 2 μL of the custom primer/probe mixture. This reaction mix was added to a DGB cartridge together with 60 μL of Droplet Generation Oil for Probes (Bio-Rad) and used for droplet generation. Droplets were then transferred to a 96-well plate (Eppendorf) and then thermal cycled with the following conditions: 5 minutes at 95°C, 40 cycles of 94°C for 30 seconds, 55°C for 1 minute followed by 98°C for 10 minutes (ramp rate 2°C/second). Droplets were analyzed with the QX200 Droplet Reader (Bio-Rad) for fluorescent measurement of FAM and HEX probes. Gates was performed on the basis of positive and negative controls, and mutant populations were identified. The ddPCR data were analyzed with QuantaSoft analysis software (Bio-Rad) to obtain fractional abundance of the mutant DNA alleles in the WT/normal background. The quantification of the target molecule was presented as the number of total copies (mutant plus WT) per sample in each reaction. Fractional abundance is calculated as follows: F.A. % = (Nmut/Nmut + Nwt) x 100, where Nmut is the number of mutant events and Nwt is the number of WT events per reaction. ddPCR analysis of normal control plasma DNA (from cell lines) and no DNA template controls were always included. Probe and primer sequences are available upon request.

Targeted Sequencing of ctdNA

Cell-free DNA was extracted from whole blood, and 5 ng to 30 ng of ctdNA was isolated. Sequencing libraries were prepared with custom in-line barcode molecular tagging, and complete sequencing at 15,000 × read depth of the critical exons in a targeted panel of 70 genes was performed at a CLIA-certified, College of American Pathologists-accredited laboratory (Guardian Health; ref. 43).

Whole-Exome Sequencing

Whole-exome sequencing (WES) was performed by the Broad Institute sequencing platform. WES of matched pretreatment and post-progression biopsies and normal blood was performed as described previously (44).

Cell Culture

Established cell lines were obtained from the following sources: Riken Bioresource Center (HuCCT1, RBE, SSP-25, HuH-28) and Korean Cell Line Bank (SNU1079). We are grateful for the kind gifts of CCLP-1 and CCSW-1 cells from Dr. P. Bosma (Academic Medical Center, Amsterdam, the Netherlands), SG231 from Dr. A.J. Demetris (University of Pittsburgh, Pittsburgh, PA), and MMNK-1 from Dr. J. Luyendyk (University of Kansas Medical Center, Kansas City, KS). Cell lines were grown at 37°C under 5% CO2 in their required growth medium (Gibco) supplemented with 10% FBS and 1% penicillin and streptomycin. To generate patient-derived biliary tract cancer cell lines (ICC13-7, ICC14, ICC15, ICC16, ICC17, ICC18, GB2), we utilized resection or autopsy specimens directly or following growth as PDXs, as per our IRB-approved protocol (DFCIC, #13-162). Samples were minced with sterile razor blades, digested with trypsin for 30 minutes at 37°C, and then resuspended in RPMI supplemented with 20% FBS, 1% l-glutamine (Gibco, #25300-081), 1% MEM nonessential amino acids solution (Gibco, #11140-050), 0.1% sodium pyruvate (Gibco, #11360-070), 0.5% penicillin/streptomycin, 10 μg/mL gentamicin (Gibco, #15710-064), and 0.2 U/mL human recombinant insulin (Gibco, #12858-014) and seeded on plates coated with rat tail collagen (BD Biosciences). Cells were passaged by trypsinization, adapted to RPMI supplemented with 10% FBS and 1% penicillin/streptomycin, and transferred to uncultured tissue culture plates prior to functional studies. They were routinely checked to be Mycoplasma free. HuCCT1, RBE, SSP-25, HuH-28, and SNU1079 cells were authenticated by short tandem repeat (STR) DNA profiling by the cell line bank from which they were obtained. Authentication by STR DNA profiling through the ATCC was performed for CCLP-1, CCSW-1, SG231, MMNK-1, ICC2, and ICC4 (between December 2015 and March 2016) and ICC13-7, ICC14, ICC15, ICC16, ICC17, ICC18, and GB2 (between January and April 2018). All cell lines were used within 20 passages of establishment from patients or receipt from repositories.

Generation of WT and Mutated FGFR2–PHGDH-Expressing Cell Lines

An FGFR2–PHGDH fusion construct, containing exons 1–17 of FGFR2-IIIb fused to PHGDH (NM_006623.3) exons 6–12, was amplified from reverse-transcribed cDNAs from an ICC patient sample and inserted into the pMSCV vector using the same kit. Targeted Sanger sequencing was done to confirm the mutations generated. Retrovirus was generated by transfecting the pMSCV constructs and packaging plasmids into 293T cells. After collection of retrovirus, transfected 293T cells were collected to confirm protein expression from each construct. Viral infections of CCLP-1 and ICC13-7 cells were performed in the presence of polybrene. Infected cells were selected in 6-15 μg/mL for one to two weeks. For both cell lines, the period of time in culture between thawing, infection, selection, recovery, and experimental setup and completion was less than 10 passages.

Cell Viability Assay

For IC50 measurement using the FGFR inhibitors, cells were dissociated into single cells and seeded into a 384-well tissue culture plate, each well with 200 viable cells and 40 μL of growth medium. After 24 hours, compounds were added to each well over a 15-point concentration range, along with DMSO controls, using a Tecan D300e digital drug dispenser. Cells were cultured for 5 days in the presence of compound before assessing viability by adding 15 μL of CellTiter-Glo (Promega) to each well, incubating for 20 minutes at room temperature on a shaker, and measuring luminescence using an Envision Digital Drug Dispenser.
plate reader. Each condition was performed in five replicates, and each dose point was normalized to DMSO controls to estimate relative viability. At least two independent experiments were performed for each compound and cell line. IC₅₀ values were determined by GraphPad Prism using a 4-parameter dose–response model. Crystal violet staining assays were done by seeding cells into 6-well plates one day before addition of drug. Cells were grown in the presence of drug for 4 days (CCLP-1 cells) or 2 weeks (ICCP-13-7 cells), then washed with PBS, fixed with cold methanol, stained with 0.5% crystal violet solution (Sigma-Aldrich), and washed extensively under tap water.

**Immunoblot Analysis**

Cells were treated with drugs in 6-well plates for 5 to 8 hours or as indicated. Cell protein lysates were prepared in ice-cold RIPA lysis buffer (50 mmol/L Tris-HCl, pH 7.4, 150 mmol/L NaCl, 2 mmol/L EDTA, 1% NP-40, 0.1% SDS, 0.5% sodium deoxycholate, containing Roche protease inhibitors and Calbiochem phosphatase inhibitor cocktail set I and II). Debris was removed by centrifugation in a microfuge at maximum speed for 10 minutes at 4°C. Protein concentration in clarified lysate was determined by Pierce BCA Protein Assay (Thermo Fisher Scientific). Ten micrograms of protein was used to perform analysis by SDS-PAGE, electrotransfer, and immunoblotting with specific antibodies. The following antibodies were used: from Cell Signaling Technology (all at 1:1,000 dilution), phospho-FRS2 Y196 (3868S), SHP2 (3397S), phospho-MEK1/2 S217/221 (9154S), MEK1/2 (4694S), phospho-ERK1/2 T202/Y204 (9106S), ERK1/2 (4695S), phospho-AKT S473 (4060S), AKT (2472S), phospho-FGFR1 (5774S), FGFR1 (9740S), from Abcam (1:3,000 dilution), FRS2 (ab183492), phospho-SHP2 Y542 (ab63232), from Abcam (1:100 dilution), FGFR2 (ab75984); from Sigma (1:20,000 dilution), β-actin (A5316).

**Quantitative RT-PCR**

Total RNA from cell lines was extracted using the RNeasy Mini Kit (Qiagen). RNA (1 μg) was reverse-transcribed using SuperScript II Reverse Transcriptase (Invitrogen) or reagents from the QuantiTect Rev. Transcription Kit (Qiagen), according to the manufacturer’s instructions. Quantitative RT-PCR (qRT-PCR) was performed on a CFX384 Real-Time PCR Detection System (Bio-Rad) with iTaq Universal SYBR Green Supermix, 2X (Bio-Rad). FGFR1–3 and FGF20 primer sets were analyzed by qRT-PCR. Data were normalized for expression of the housekeeping gene ribosomal 18S. Primer sequences are provided in Supplementary Table S1.

**PDX Treatment Studies**

Mice were housed in pathogen-free animal facilities. All experiments were conducted under protocol 2005N000148 approved by the Subcommittee on Research Animal Care at MGH. To develop an FGFR2 fusion human PDX, we obtained tissue from a fresh resection specimen from a patient with an FGFR2–KIAA1217 fusion ICC tumor, per our IRB-approved protocol (DFCI# 13-162). The tissue was rinsed in Hank’s Balanced Salt Solution and cut into 0.3–0.5 mm³ pieces with sterile razor blades. These tumor pieces were implanted subcutaneously into 6– to 8-week-old female NSG mice (NOD.Cg-Prkdcsid1Ifgrtg1Wj/Msfj, 00557, The Jackson Laboratory). Tumor size was measured with a digital caliper. Upon reaching approximately 500 mm³, mice were randomized to either vehicle control or 25 mg/kg TAS-120 (in hydroxypropyl methylcellulose) solution by oral gavage daily for 3 and 14 days prior to harvest. Tumor samples were collected for biochemical analysis and histology processing. For histology processing, tissue samples were fixed overnight in 4% buffered formaldehyde, embedded in paraffin, and then sectioned and stained with hematoxylin and eosin by the MGH Pathology Core. IHC was performed on paraffin-embedded sections (5-μm thickness). After deparaffinization and dehydration, slides were incubated for 10 minutes with 3% H₂O₂ at room temperature to block endogenous peroxidase activity. Specimens were brought to the boil in 10 mmol/L sodium citrate buffer (pH 6.0, 5 minutes, pressure cooker) for antigen retrieval. Slides were blocked for 1 hour in TBS-0.05% Tween 20 (Thermo Fisher Scientific). 1 drop per 1 mL of Protein Block (Dako X0909) and incubated with primary antibody for 1 hour at room temperature. Primary anti-Ki-67 antibody (Abcam, ab15580) was diluted with PBS-Protein Block (1 drop/mL) at a ratio of 1:400 and incubated with the tissue sections at 4°C. Specimens were reacted for 30 minutes with ImmPRESS HRP polymer reagent (Vector Laboratories) combined with secondary antibodies. Slides were then washed with PBS and stained for peroxidase for 1-2 minutes with the Betapoint DAB Chromogen reagent, washed with water, and counterstained with hematoxylin. Stained slides were photographed with an Olympus DP72 microscope.

**Population Growth Modeling with Clonal Pool System**

Cell pools containing 1% of each mutant cell line and 90% of FGFR2–PHGDH WT cells were seeded at low confluency in 6-well plates. We used 1% as an empirically chosen concentration that allowed growth of cells in the presence of different FGFR inhibitors for 2 weeks without individual mutant clones overtaking the entire population. This percentage was also sufficiently high to enable clone detection by ddPCR. Every experimental condition was performed in triplicate. Two independent experiments were used to generate data for Fig. 3B and C. Drug incubation (or DMSO-treated controls) started 24 hours after cell seeding, and drug treatment was refreshed every 3 to 4 days. After 1, 7, or 14 days in culture, the remaining cells were trypsinized and collected for genomic DNA extraction. Genomic DNA extracted from cells using the DNeasy Blood and Tissue Kit (Qiagen) was subjected to enzymatic fragmentation with either MsrI or HaeIII, and amplified using ddPCR Supermix for Probes (Bio-Rad) using FGFR2 assays (PrimePCR ddPCR Mutation Assay, Bio-Rad, and custom-designed). DNA template (20–40 ng) was added to 12.5 μL of ddPCR Supermix for Probes (Bio-Rad) and 1.25 μL of the primer/probe mixture. This reaction mix (final volume = 25 μL) was added to a DG8 cartridge together with 70 μL of Droplet Generation Oil for Probes (Bio-Rad) and used for droplet generation. Droplets were then transferred to a 96-well plate (Eppendorf) and then thermal cycled with the following conditions: 10 minutes at 95°C, 40 cycles of 94°C for 30 seconds, 55°C for 1 minute followed by 98°C for 10 minutes (ramp rate 2°C/second). Droplets were analyzed with the QX200 Droplet Reader (Bio-Rad) for fluorescent measurement of FAM and HEX probes. Gating was performed on the basis of positive and negative controls, and mutant populations were identified. The ddPCR data were analyzed with QuantaSoft analysis software (Bio-Rad) to obtain fractional abundance of the mutant DNA alleles in the WT/normal background. The quantification of the target molecule was presented as number of total copies (mutant plus WT) per sample in each reaction. Fractional abundance is calculated as follows: F.A. % = [Nmut/(Nmut + Nwt) × 100], where Nmut is number of mutant events and Nwt is number of WT events per reaction. The number of positive and negative droplets is used to calculate the concentration of the target and reference DNA sequences and their Poisson-based 95% confidence intervals, as described previously (45). Multiple replicates (minimum of three) were performed for each sample. ddPCR analysis of normal control genomic DNA from cell lines and no DNA template (water) controls was performed in parallel with all samples, including multiple replicates as contamination-free controls.

**In Silico Structural Modeling**

TAS-120 was docked into FGFR2 (PDBID: 1OEC). The loop structure (V488-V496) was modeled so that the S atom of C492 and the...
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Disclosure of Potential Conflicts of Interest

L. Goyal is a consultant/advisory board member for Taiho, Pieris Pharmaceuticals, and Agios Pharmaceuticals. J.K. Lennerz, N. Bardeesy, and A.J. Iafrate report receiving a commercial research grant from Bristol-Myers Squibb in association with Genentech. C.H. Benes reports receiving commercial research support from Genentech. D.T. Ting is a consultant/advisory board member for Genentech. S. Reyes reports receiving commercial grants from Genentech and, in his personal capacity, from Taiho and Merck. A. Bardelli reports receiving commercial research grants from Biocartis and Takeda, is a consultant/advisory board member for Cytometryx, and is a recipient of institutional research support from NCI SPORE P50 CA127003. A. Bardelli and N. Bardeesy have ownership interest (including stock, patents, etc.) in Neophere, Phoremost, and Kither. The authors are not aware of any other potential conflicts of interest.

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Authors’ Contributions


Study supervision: L. Goyal, J.K. Lennerz, W.C. Hahn, A.X. Zhu, R.B. Corcoran, N. Bardeesy

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