BI-3406, a Potent and Selective SOS1–KRAS Interaction Inhibitor, Is Effective in KRAS-Driven Cancers through Combined MEK Inhibition

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ABSTRACT

KRAS is the most frequently mutated driver of pancreatic, colorectal, and non–small cell lung cancers. Direct KRAS blockade has proved challenging, and inhibition of a key downstream effector pathway, the RAF–MEK–ERK cascade, has shown limited success because of activation of feedback networks that keep the pathway in check. We hypothesized that inhibiting SOS1, a KRAS activator and important feedback node, represents an effective approach to treat KRAS-driven cancers. We report the discovery of a highly potent, selective, and orally bioavailable small-molecule SOS1 inhibitor, BI-3406, that binds to the catalytic domain of SOS1, thereby preventing the interaction with KRAS. BI-3406 reduces formation of GTP-loaded RAS and limits cellular proliferation of a broad range of KRAS-driven cancers. Importantly, BI-3406 attenuates feedback reactivation induced by MEK inhibitors and thereby enhances sensitivity of KRAS-dependent cancers to MEK inhibition. Combined SOS1 and MEK inhibition represents a novel and effective therapeutic concept to address KRAS-driven tumors.

SIGNIFICANCE: To date, there are no effective targeted pan-KRAS therapies. In-depth characterization of BI-3406 activity and identification of MEK inhibitors as effective combination partners provide an attractive therapeutic concept for the majority of KRAS-mutant cancers, including those fueled by the most prevalent mutant KRAS oncoproteins, G12D, G12V, G12C, and G13D.

See related commentary by Zhao et al., p. 17.

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INTRODUCTION

KRAS functions as a molecular switch, cycling between inactive (GDP-bound) and active (GTP-bound) states to transduce extracellular signals via cell-surface receptors. KRAS signaling occurs through engagement with effector proteins that orchestrate intracellular signaling cascades regulating tumor cell survival and proliferation. Aberrant activation of KRAS by deregulated upstream signaling (1), loss of GTPase-activating protein (GAP) function (2, 3), or oncogenic mutations results in increased GTP-bound KRAS and persistent downstream signaling (4, 5). Mutations in the KRAS gene occur in approximately one of seven of all human cancers, making it the most frequently mutated oncogene (6, 7). Up to 90% of pancreatic tumors bear activating KRAS mutations. Mutated KRAS is also observed at high frequency in other common tumors, including colorectal cancer (~44%) and non–small cell lung cancer (NSCLC; ~29%). Cancer-associated mutations in KRAS cluster in three hotspots (G12, G13, and Q61), with a majority (77%) of mutations causing single amino acid substitutions at G12. The KRAS missense mutation G12D is the most predominant variant in human malignancies (35%), followed by G12V (29%), G12C (21%), G12A (7%), G12R (5%), and G12S (3%). Besides G12, the hotspots G13 and Q61 show mutation rates of 10% and 6%, respectively (KRAS mutation frequencies were derived from American Association for Cancer Research GENIE v6.1 and The Cancer Genome Atlas; refs. 6, 7). In preclinical models, activated KRAS has been shown to drive both the initiation and the maintenance of a range of cancer types (8–11). Despite the compelling rationale to target KRAS, identification of potent direct inhibitors has been challenging. Promising early results from clinical trials with the two inhibitors, AMG 510 (sotorasib) and MRTX849 (adagrasib), both targeting the KRAS<sup>G12C</sup>-mutant allele covalently and specifically (12, 13), have been reported. These inhibitors demonstrated clinical activity primarily in NSCLC, where the KRAS<sup>G12C</sup> mutation frequency is highest (14, 15). Moreover, a nanomolar pan-RAS inhibitor binding to a second pocket on RAS has been described previously (16).

Despite this recent success, molecularly targeted therapies that effectively address the most prevalent KRAS mutant alleles beyond G12C, including G12D and G12V, are lacking. Attempts to indirectly target KRAS-driven tumors through
inhibition of downstream effectors of KRAS, such as members of the RAF–MEK–ERK cascade, have suffered limited clinical success (17), in part due to the capacity of cancer cells to adapt by rapidly increasing KRAS-GTP levels. The SHP2 protein-tyrosine phosphatase is an important mediator of cellular signaling through the RAS/MAP kinase pathway and is thought to act via activation of SOS1-regulated RAS-GTP loading. SHP2 inhibitors are being explored by several companies, with the most advanced inhibitors, RMC-4630 and TNO155, currently under study in Phase I clinical trials (18–21). Published data show particular sensitivity to SHP2 inhibitors in KRASG12D-mutant tumors (20).

Dynamic control of the extent and kinetics of the RAS–RAF–MEK–ERK signaling is governed by positive and negative feedback loops (22). SOS1 is a key guanine exchange factor (GEF) for KRAS that binds and activates GDP-bound RAS family proteins at its catalytic binding site and in this way promotes exchange of GDP for GTP. In addition to its catalytic site, SOS1 can also bind GTP-bound KRAS at the allosteric site that potentiates its GEF function, constituting a mechanism for positive feedback regulation (23). Depletion of SOS1 or specific genetic inactivation of its GEF function has been shown to decrease the survival of tumor cells harboring a KRAS mutation (24). This effect was not observed in wild-type cells that are not KRAS addicted (24). Pathway activation leads to ERK-mediated phosphorylation of SOS1, but not its paralog SOS2, thereby attenuating SOS1 GEF activity (25, 26). This suggests that SOS1 acts as an important node in the negative feedback regulation of the KRAS pathway (25, 26). On the basis of these lines of evidence, we hypothesized that a potent and selective SOS1 inhibitor would synergize with an MEK inhibitor, resulting in strong and sustained pathway blockade and a robust antitumor efficacy in KRAS-driven cancers.

In 2014, small molecules were described that bind to a lipophilic pocket of SOS1, in close proximity to the RAS-binding site (27). Binding of these ligands increased SOS1-mediated nucleotide exchange and consequently led to activation of RAS. Recently, SOS1 inhibitor tool compounds were reported (28), but these nonbioavailable compounds did not demonstrate the expected differential effect on KRAS-driven cancer cell lines versus wild-type cells.

In this article, we describe the discovery of BI-3406, a potent and selective SOS1–KRAS interaction inhibitor, and elucidate its mode of action both in vitro and in vivo. BI-3406 potently decreases the formation of GTP-loaded RAS and reduces cell proliferation of a large fraction of KRAS G12C-driven and non-G12C-driven cancers in vitro and in vivo. BI-3406 attenuates feedback reactivation by MEK inhibitors and enhances sensitivity of KRAS-dependent cancers to MEK inhibition, resulting in tumor regressions at well-tolerated doses in mouse models. Our data provide strong evidence that combined SOS1 and MEK inhibition represents an attractive therapeutic concept to address KRAS-driven human tumors.

**RESULTS**

**Discovery of BI-3406, a Potent and Selective SOS1–KRAS Interaction Inhibitor**

To discover SOS1 inhibitors, we conducted a high-throughput screening of 1.7 million compounds using an alpha screen and a fluorescence resonance energy transfer assay as an orthogonal biochemical screen on SOS1 and KRASG12D. Several hits containing a quinazoline core were identified, best exemplified by BI-68BS (Supplementary Fig. S1A). A stoichiometric and saturable dissociation constant, using surface plasmon resonance on SOS1 (K_D = 470 nmol/L) and the corresponding activity in a GDP-dependent KRAS–SOS1 displacement assay (IC_50 = 1.3 μmol/L), indicated effective disruption of the SOS1–KRAS protein–protein interaction. Cocrystalization of BI-68BS and SOS1 confirmed binding to a pocket (27) next to the catalytic binding site on SOS1 (Supplementary Fig. S1B and S1C; Supplementary Table S1) with the quinazoline ring pi-stacking to His905SOS1. On the basis of the structural data, the interaction of the methoxy substituted of BI-68BS with Tyr884SOS1 most likely interfered with the competing Tyr884SOS1–Arg73K Ras interaction and consequently prevented KRAS from binding to SOS1 (Supplementary Fig. S1D). In an effort to optimize BI-68BS, several modifications were made, which led to the discovery of BI-3406 (Fig. 1A). As BI-68BS was originally synthesized as part of a project targeting EGFR, a methyl substituent was incorporated in the second position of the quinazoline core to effectively eliminate any interfering inhibition of kinase activity (tested in a panel of 324 kinases; Supplementary Tables S2 and S3). Introduction of a trifluoromethyl and an amino substituent at the phenethyl moiety filled the pocket more effectively and formed an H-bond...
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**Biochemical protein–protein interaction assays**

**A**

**B**

**C**

**D**

**E**

**F**

**G**
with M878SOSt, respectively, thereby significantly increasing potency. The tetrahydrofuryl substituent favorably balanced solubility and metabolic stability and improved interaction with Tyr884SOSt. Synthesis of BI-3406 is described in detail in the Supplementary Data section (for synthesis route see also Supplementary Fig. S1E and S1F). Crystallization data can be found in Supplementary Table S1.

A detailed biochemical characterization of BI-3406 was made possible through the analysis of a variety of interaction assays using SOS1 and SOS2 recombinant proteins, in combination with several mutant KRAS variants. BI-3406 was found to be a potent, single-digit nanomolar inhibitor binding to the catalytic site of SOS1 and thereby blocking the interaction with KRAS-GDP, as exemplified in the interaction assay with KRASG12D-mutant and KRASG12C-mutant oncoproteins (Fig. 1B).

A recently developed, covalent KRASG12C-specific inhibitor (ARS-1620) was able to interfere with the SOS1–KRASG12C protein–protein interaction, but, in contrast to BI-3406, had no effect on the protein–protein interaction of SOS1 with KRASG12D (Fig. 1C and D). Upon replacement of SOS1 with its paralog SOS2, BI-3406 lost its ability to interfere with KRAS binding, indicating that BI-3406 is a highly potent, SOS1-specific inhibitor that can address multiple KRAS-mutant oncoproteins (Fig. 1B). The SOS1 selectivity of BI-3406 can be explained by a potential clash of the compound with Val903 and the absence of pi-interaction in SOS2, which is revealed in an overlay of the published SOS2 apo structure [Protein Data Bank (PDB) code 6EIE] with our own SOS1 BI-3406 cocrystal structure (Supplementary Fig. S1G). In a biochemical protein–protein interaction assay, the introduction of the mutations Y884A and H905V in a recombinant SOS1 protein strongly impaired the ability of BI-3406 to disrupt the interaction with KRASG12D (Supplementary Fig. S1H).

Importantly, expression of FLAG–SOS1 transgenes in MIA PaCa-2 and HEK293 cells revealed that the SOS1 mutations H905V and H905I abrogated the ability of BI-3406 to inhibit phosphorylation of ERK (pERK) and cell proliferation, demonstrating selective SOS1 on-target activity of the compound in a cellular context (Fig. 1E and F; Supplementary Fig. S1I).

To further investigate whether BI-3406 was capable of cellular SOS1 inhibition, cells were treated with increasing concentrations of BI-3406. The compound inhibited RAS-GTP levels with an IC50 of 83–231 nmol/L in SOS1/KRAS-dependent NCI-H358 (KRASG12C) and A549 (KRASG12C) cells (Fig. 1G). Stimulation of starved NCI-H358 and MIA PaCa-2 cells with EGF resulted in an increase of RAS-GTP levels that was significantly more pronounced in NCI-H358 cells with EGF resulted in an increase of RAS-GTP levels that was significantly more pronounced in NCI-H358 (Fig. 1G). Stimulation of starved NCI-H358 and MIA PaCa-2 cells revealed that the SOS1 mutations H905V and H905I abrogated the ability of BI-3406 to inhibit phosphorylation of ERK (pERK) and cell proliferation, demonstrating selective SOS1 on-target activity of the compound in a cellular context (Fig. 1E and F; Supplementary Fig. S1I).

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**Association of KRAS Mutation Status with Sensitivity to SOS1 Inhibition**

The cellular activity of BI-3406 was further evaluated across a wider panel of cancer cell lines driven by different KRAS pathway activating mutations. As SOS1 is uniformly expressed across all tumor types, a SOS1 inhibitor could be broadly applicable in KRAS-driven indications (Supplementary Fig. S2A and S2B). Plotting the expression of SOS1 against SOS2 revealed that the cell lines used in our subsequent experiments harbored SOS1/SOS2 mRNA ratios representative of ratios observed in a large dataset of human tumors (Supplementary Fig. S2C). A dose-dependent partial reduction of pERK levels was observed in all RAS-mutated cell lines tested, with an IC50 value of 17 and 57 nmol/L (IC50 value was defined as the inflection point of the curve; Fig. 2A). No pERK modulation was observed in A375 melanoma cells that are KRAS wild-type and harbor an activating BRAFV600E mutation that likely renders them independent of KRAS signaling (Fig. 2A).

Cell lines expressing mutant KRAS have demonstrated variable dependencies upon KRAS for viability in two-dimensional (2-D) monolayer proliferation assays (29), whereas KRAS dependency is better modeled in anchorage-independent three-dimensional (3-D) growth assays. Consistent with this observation, we demonstrated that BI-3406 inhibited the 3-D growth of four KRAS-mutant cancer cells with an IC50 of 16–52 nmol/L, as half-maximum inhibitory concentration (Fig. 2B). In contrast, the 3-D growth of the two KRAS wild-type cancer cell lines, NCI-H520 and A375, was not appreciably affected (Fig. 2B), but responded to the broad antiproliferative agent panobinostat, a histone deacetylase inhibitor (Fig. 2C). Collectively, these data show a clear correlation between signaling pathway and growth inhibition by BI-3406 in KRAS-driven cancer cell lines.

The growth-inhibitory effects of BI-3406 across different KRAS-mutated cell lines could be influenced by tumor lineage or comutations. Therefore, we evaluated the effect of SOS1 inhibition on a panel of isogenic cell lines, differing only in the status of their KRAS allele. We used NCI-H23 cells carrying a heterozygous KRASG12C allele and replaced the G12C codon by heterozygous G12D, G12V, G12R, and G13D or homozygous G12D, G13D, and Q61H mutations. BI-3406 showed comparable activity, independent of zygosity, with an approximate
Figure 2. Drug sensitivity profiling of cancer cell lines uncovers an association of KRAS mutation status with sensitivity to SOS1 inhibition. A, Inhibition of pERK activity by BI-3406 after 1 hour in 2-D assay conditions in a cancer cell line panel quantified by Western blotting (n = 2, means ± SD). A375 (KRAS wild-type (WT)), A549 (KRASG12D), DLD1 (KRASG13D), NCI-H23 (KRASG12C), and NCI-H520 (KRAS WT and BRAF WT). B, Inhibition of cell proliferation by BI-3406 in a cancer cell line panel in 3-D proliferation assays (n = 3, means ± SD). C, In vitro sensitivity of a panel of cell lines to the positive control, panobinostat (Sigma-Aldrich), in a 3-D proliferation assay (n = 3, means ± SD). D, Effect of BI-3406 on pERK levels in a panel of isogenic NCI-H23 cell lines. Values were normalized to total ERK protein (n = 2, means ± SD). E, In vitro sensitivity of a panel of isogenic cell lines treated with BI-3406 in a 3-D proliferation assay (n = 3, means ± SD). F, In vitro sensitivity of 40 cancer cell lines treated with BI-3406 in 3-D proliferation assays. Panels depict the proliferation data (n = 2), the respective cancer type, and the mutation status of selected genes. Cell lines are grouped on the basis of an IC50 cutoff of 100 nmol/L. The mutation status and zygosity are shown by a continuous color-coding scheme; blue boxes reflect wild-type (WT) and light blue boxes indicate an unknown status. Only recurring hotspot mutations are reported for KRAS, NRAS, HRAS, BRAF, EGFR, and HER2 (Supplementary Table S5). NCI-H2347 carries a KRASG12F mutation (asterisks); SCLC, small cell lung cancer.
to wild-type KRAS, pERK modulation was observed following treatment with BI-3406, but the wild-type cells were no longer able to grow in a 3-D proliferation assay.

We further profiled BI-3406 in a larger panel of 40 solid cancer cell lines with known oncogenic alterations in KRAS, NRAS, HRAS, EGFR, NFI, and BRAF (Fig. 2F; Supplementary Tables S4 and S5). Excitingly, BI-3406 sensitivity correlated with KRAS mutation status in this large cell line panel (Fisher exact test, \( P = 0.00337; \) Fig. 2F). Sensitive cell lines harbored a broad range of KRAS-mutant alleles (Supplementary Tables S4 and S5), including KRAS G12C, G12V, G12S, G12A, and G13D mutations. Although no difference in sensitivity could be observed on the basis of the zygosity of the KRAS mutation, it was notable that two of the three nonresponsive KRAS-mutant cell lines, as well as three of five nonresistant NRAS-mutant cell lines, were characterized by a Q61 mutation. NF1 is a tumor suppressor and a Ras GAP (2). Loss of NF1 function has been shown to increase Ras-GTP levels, hyperactivate Ras/MAPK signaling, and contribute to a variety of human cancers (32, 33). Therefore, we assessed whether NF1 aberrations were sensitive to BI-3406 treatment, irrespective of their KRAS status. No other driver mutations in components of the RTK/KRAS/MAPK pathway could be identified in several of these sensitive cell lines, suggesting NF1 aberrations are a key determinant for sensitivity to BI-3406 in these lines (Supplementary Table S4). Similarly, a fraction of NSCLC cell lines driven by EGFR mutations also responded to BI-3406 treatment, suggesting that oncogenic receptor tyrosine kinases (RTK) can confer sensitivity to SOS1 inhibition. As none of the six BRAF-mutant and five NRAS-mutant cell lines were sensitive to treatment with BI-3406 (Fig. 2F), we hypothesize that NRAS and BRAF mutations are associated with resistance to BI-3406 monotherapy (\( P < 0.001 \)). Collectively, our findings highlight the critical function of SOS1 in promoting KRAS/MAPK pathway activation in a large fraction of cancers driven by KRAS G12C and non-G12C alleles and NF1 aberrations, as well as EGFR mutations.

The pharmacodynamics of BI-3406 were further evaluated. In sensitive cell lines, treatment with BI-3406 resulted in sustained pathway modulation of ERK1/2 phosphorylation (Supplementary Fig. S2D and S2E), in contrast to insensitive cell lines, which exhibited weaker and more short-lived effects (NICI-H2170 and NICI-H1299; Supplementary Fig. S2E). Compared with pERK levels, levels of pAKT Ser473 and Thr308 were less strongly affected by BI-3406 (Supplementary Fig. S2D and S2E).

We subsequently tested BI-3406 side-by-side with the recently reported SOS1 inhibitor BAY-293 (28) and the SHP2 inhibitor SHP099 (18) in 2-D and 3-D proliferation assays across a panel of 24 cell lines, including 18 KRAS-mutated cell lines (Supplementary Table S6). The three compounds demonstrated no activity in 2-D proliferation assays. In 3-D proliferation assays, SHP099 showed the strongest antiproliferative effects, with an IC\(_{50}\) between 167 and 790 nmol/L in KRAS\(^{G12C}\), a subset of G12D cell lines, and in one G12S cell line it yielded modest effects in KRAS\(^{G12D}\) and KRAS\(^{G12V}\) cells (IC\(_{50}\) 1,180-4,411 nmol/L), whereas no effects were detectable in Q61L/H- and G12R KRAS-mutant tumor cells. BI-3406 caused cell-growth inhibition in all KRAS G12- and G13-mutant cell lines (IC\(_{50}\), 9-220 nmol/L) with the exception of G12R- and KRAS Q61L/H-mutant tumor cells. The previously published SOS1 inhibitor BAY-293 demonstrated only a very limited potency and, in contrast to BI-3406, no sizeable selectivity for KRAS-mutated cells as compared with KRAS wild-type cells (Supplementary Table S6). This suggests that BI-3406 and SHP099 possess a partially overlapping yet distinct profile across KRAS-mutated cell lines, with BI-3406 being more broadly active in 3-D proliferation assays.

To glean first insights regarding a potential therapeutic index of BI-3406, we tested the compound on primary cells and nontumorigenic cells in vitro. BI-3406 inhibited the proliferation of foreskin fibroblasts with an IC\(_{50}\) of 37 nmol/L, while two other cell types, primary smooth muscle cells and retinal pigment epithelial cells, were not affected (IC\(_{50}\) > 5 µmol/L; Supplementary Fig. S2F-S2H). The extremely potent and widely used MEK inhibitor trametinib affected proliferation of all three aforementioned cell types (retinal pigment epithelial cells IC\(_{50}\) of 12 nmol/L, primary smooth muscle cells IC\(_{50}\) of 843 nmol/L, and normal foreskin cells IC\(_{50}\) of 85 nmol/L).

**SOS1 Inhibition Suppresses Tumor Growth in Xenograft Models of KRAS-Driven Cancers**

BI-3406 is an orally bioavailable compound (Supplementary Fig. S3A), and single administration was sufficient to reduce Ras-GTP and pERK levels in A549 xenograft tumors over a period of 24 and 7 hours, respectively (Supplementary Fig. S3B and S2C). At a dose of 50 mg/kg twice a day, relevant levels of unbound exposures were achieved for the first 12 hours when compared with unbound IC\(_{50}\) levels in A549 cells (Supplementary Fig. S3A). In MIA PaCa-2 tumor-bearing mice, twice-daily compound treatment with 50 mg/kg BI-3406 resulted in pathway modulation over a period of up to 10 hours (Fig. 3A; Supplementary Fig. S3D). At the 24-hour time-point, the compound was cleared (Supplementary Fig. S3A and S3D) and pERK levels returned to baseline in both A549 and MIA PaCa-2 tumors (Fig. 3A; Supplementary Fig. S3B). In the same experiment, a reduction of pERK levels was observed by IHC in surrogate tissue (murine skin) over a similar period (Fig. 3B; Supplementary Fig. S3E). The use of phosphorylation markers can be challenging in a clinical setting, effects on RAS-dependent gene-expression signatures were analyzed in the MIA PaCa-2 xenograft model. Prolonged suppression of known pathway-related genes, such as SPRY4 and DUSP6, and transcriptional regulators, such as FOSL1, EGR1, ET51, ETV4, and ETV5, was observed (Fig. 3C; Supplementary Fig. S3F; Supplementary Table S7), in line with published data on gene-expression responses to other specific MAPK pathway inhibitors (34, 35). Of note, no effects on SOS2 mRNA expression were observed upon treatment with BI-3406 during the period of observation (Supplementary Fig. S3G and S3H), suggesting no compensatory upregulation.

On the basis of its potent cellular activity and favorable pharmacokinetic properties, the efficacy of BI-3406 was evaluated in established, subcutaneous KRAS\(^{G12D}\)-mutated MIA PaCa-2 xenografts. Twice-daily treatment with either 12 or 50 mg/kg of BI-3406 was well tolerated and resulted in prolonged dose-dependent tumor growth inhibition (\( P < 0.005 \) as compared with vehicle control; Fig. 3D and E).
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A375 (*, P < 0.008) and LoVo (**, P < 0.001) xenograft models (Fig. 3F; Supplementary Fig. S3I and S3J). Similar tumor growth inhibitory effects were observed in tested xenograft models: SW620 colorectal (KRAS$^{G12C}$, BRAF WT), LoVo colorectal (KRAS$^{G12D}$, BRAF WT), MIA PaCa-2 pancreas (KRAS$^{G12C}$, BRAF WT), and A549 NSCLC (KRAS$^{G12S}$, BRAF WT). Significant TGI was achieved in all tested KRAS-mutant xenograft models, with the exception of the KRAS WT model A375 (*, P < 0.05; **, P < 0.01; ***, P < 0.001, one tailed t test).

**Figure 3.** SOS1 inhibition suppresses tumor growth and KRAS/MAPK signaling in xenograft models of KRAS-driven cancers. A, pERK levels analyzed by a multiplexed immunoassay in explanted MIA PaCa-2 tumors treated with 50 mg/kg BI-3406 twice daily at the timepoint 0 and 6 hours (n = 5 animals/group, means ± SEM, two-tailed t test). B, pERK levels in mouse skin treated with BI-3406 (4 h), BI-3406 (24 h), BI-3406 (10 h), BI-3406 (4 h), Vehicle control (4 h), Vehicle control (24 h), Vehicle control (0 h), Vehicle control (10 h) (t test). C, Gene-expression profiling of pharmacodynamics biomarkers in a MIA PaCa-2 in vivo biomarker experiment (n = 4–5 animals/group, medians of normalized gene expression). A subset of nine genes shows time-dependent modulation after BI-3406 (50 mg/kg) treatment, visualized as a color-coded expression heat map. D, Antitumor effect of BI-3406 in the MIA PaCa-2 xenograft model (n = 7 animals/group, means ± SEM, one-tailed t test), bid, twice a day. E, Median body-weight change of mice bearing subcutaneous MIA PaCa-2 xenografts administered as described in D (n = 7 animals/group, medians). F, Responses of different xenograft models after treatment with BI-3406 (50 mg/kg bid) or vehicle (control). Tumor growth inhibition (TGI) was determined on the basis of tumor size after 20–23 days of continuous treatment (n = 7–9 animals/group, means ± SD). Genotypes of tested xenograft models: SW620 colorectal (KRAS$^{G12V}$, BRAF WT), LoVo colorectal (KRAS$^{G12D}$, BRAF WT), MIA PaCa-2 pancreas (KRAS$^{G12C}$, BRAF WT), and A549 NSCLC (KRAS$^{G12S}$, BRAF WT). Significant TGI was achieved in all tested KRAS-mutant xenograft models, with the exception of the KRAS WT model A375 (*, P < 0.05; **, P < 0.01; ***, P < 0.001, one tailed t test).

Similar tumor growth inhibitory effects were observed in SW620 (KRAS$^{G12X}$), LoVo (KRAS$^{G12X}$), and A549 (KRAS$^{G12S}$) xenograft models (Fig. 3F; Supplementary Fig. S3I and S3J). No antitumor response was observed in the BRAF-mutant A375 xenograft model (Supplementary Fig. S3K), consistent with the lack of effect on cell proliferation in this cell line in vitro. Thus, oral administration of BI-3406 monotherapy inhibits the growth of KRAS G12C, G12V, G13D, and G12S xenograft models.

**Dual SOS1 and MEK Inhibition as Effective Strategy to Treat KRAS-Mutant Tumors**

Previous work showed that many cancer models develop adaptive resistance to MEK inhibitors, often due to the reactivation of SOS1 (17). Therefore, we reasoned that dual SOS1 and MEK inhibition could constitute an effective strategy to treat KRAS-mutant tumors. Consistent with this hypothesis, the combination of BI-3406 with the MEK inhibitor trametinib yielded strong synergistic antiproliferative effects in MIA PaCa-2 (KRAS$^{G12C}$) and DLD1 (KRAS$^{G13D}$) cells in vitro (Supplementary Fig. S4A). On the basis of these promising cellular data, we tested BI-3406 plus trametinib in both the pancreatic cancer MIA PaCa-2 and the colorectal cancer LoVo (KRAS$^{G13D}$) xenograft mouse models. The MEK inhibitor trametinib was primarily used because of its favorable mouse pharmacokinetic properties ($t_{1/2} = 33$ hours; ref. 36). The combination of 50 mg/kg BI-3406 twice daily with the clinically relevant dose of trametinib (0.1–0.125 mg/kg, twice a day; for calculation details please see description in Supplementary Data) was well tolerated (Supplementary Fig. S4B and S4C) and caused substantial regressions in the entire cohort of MIA PaCa-2 tumor–bearing mice (Fig. 4A and B). Furthermore, following combination treatment, slow regrowth of tumors was detectable only 22 days after drug withdrawal (Fig. 4A). Similar results were observed in LoVo xenografts, with the effect of the BI-3406 and trametinib combination being significantly stronger compared with both monotherapies, with sustained tumor inhibition for 7 days following drug withdrawal (Fig. 4C and D). We tested two KRAS$^{G12C}$ colorectal cancer patient-derived xenograft (PDX) models and one KRAS$^{G12D}$ and one KRAS$^{G12S}$ pancreatic cancer PDX model and observed improved antitumor activity using a combination of BI-3406 with trametinib (Fig. 4E and F; Supplementary Fig. S4D–S4G). As expected on the
basis of proliferation assays using KRAS Q61− mutant cells, monotherapy of BI-3406 resulted in only weak efficacy in the KRASQ61K-mutant PDX model, yet the SOS1 and MEK inhibitor combination significantly improved antitumor activity as compared with both monotherapies (P = 0.0026; Supplementary Fig. S4G). The combination treatment was very well tolerated (Supplementary Fig. S4B and S4C and S4H–S4K).

As SOS2 may promote resistance to SOS1 over time, we analyzed the colorectal cancer PDX model B8032, but found no compensatory upregulation of SOS2 mRNA levels upon 21 days of treatment with both SOS1 and MEK inhibitor (Supplementary Fig. S4L and S4M).
The SOS1 Inhibitor BI-3406 Prevents Adaptive Resistance to MEK Inhibition

The mechanism underlying the SOS1/MEK inhibitor combination efficacy was evaluated with regard to the impact on modulation of the KRAS–RAF–MEK–ERK cascade. In vitro MEK inhibitor treatment at clinically relevant doses, in the low nmol/L range (see description in Supplementary Data), resulted in a progressive increase of MEK1/2 Ser217/221 phosphorylation, an effect termed adaptive resistance or negative feedback relief (Fig. 5A; Supplementary Fig. S5A; Supplementary Table S8). In addition, the combination of BI-3406 and trametinib elicited a reduction of pERK and blockade of adaptive resistance, measured by pERK and S6 phosphorylation as assessed by reverse phase protein array (RPFA) analysis in two colorectal cancer PDX models (Supplementary Fig. S3I; Supplementary Table S8). In addition, the combination led to enhanced reduction of DUSP6 mRNA in MIA PaCa-2 tumors (Supplementary Fig. S5J) and augmented induction of apoptosis as shown in the KRAS-driven cell line DLD1 (Supplementary Fig. S5K).

Finally, we investigated whether the beneficial effect of BI-3406 described above could be extended to a direct KRAS
inhibitor, the clinical KRAS\textsuperscript{G12C} inhibitor AMG 510. Strikingly, the combination of AMG 510 with BI-3406 resulted in stronger and more prolonged suppression of pERK as compared with AMG 510 monotherapy in NCI-H358 (KRAS\textsuperscript{G12C}) cells in vitro (Supplementary Fig. S5L). The addition of the SOS1 inhibitor to AMG 510 largely prevented the rebound of pERK at the 72-hour timepoint. A similar effect was observed at 72 hours upon combination of AMG 510 with the SHP2 inhibitor SHP099 (Supplementary Fig. S5L).

In summary, the SOS1 inhibitor BI-3406 enhances the extent and duration of MAPK pathway inhibition upon combination with a MEK or KRAS G12C inhibitor, suggesting it is able to counteract adaptive resistance. This highlights SOS1 inhibition as a promising combination option for MAPK pathway and direct KRAS inhibitors. In line with this, we show that a SOS1–MEK inhibitor combination enables long-term pathway inhibition, resulting in tumor regressions in multiple KRAS-driven cancer models at well-tolerated doses (Fig. 5D).

**DISCUSSION**

KRAS mutations are the most frequent gain-of-function alterations found in patients with cancer, yet KRAS-driven tumors are largely refractory to anticancer therapies. Despite more than two and a half decades of research describing the central role of SOS1 in developmental and oncogenic signaling pathways, most notably in the direct activation of RAS oncoproteins (37–40), no SOS1 inhibitor has progressed to the clinic. The previously described catalytic SOS1 modulator BAY-293 (28) inhibited cancer cell proliferation with weak potency and irrespective of KRAS status. Here, we describe a highly potent and selective small-molecule inhibitor, BI-3406, that binds to SOS1 and thereby blocks protein–protein interaction with RAS-GDP. BI-3406 is the first example of an orally bioavailable SOS1–KRAS interaction inhibitor that reduces RAS-GTP levels and curtails MAPK pathway signaling in vitro and in vivo. BI-3406 limits the growth of the majority of tumor cells driven by KRAS variants at positions G12 and G13, as shown in 3-D proliferation assays. As tumors bearing these KRAS mutations are most prevalent in colorectal cancer, pancreatic cancer, and NSCLC, these results provide compelling evidence that the SOS1–KRAS interface is a druggable target of potential clinical importance, and highlight BI-3406 as a first-runner of a new generation of GDP-KRAS–directed inhibitors with promising therapeutic potential. In contrast to covalent KRAS\textsuperscript{G12C}–specific inhibitors (12, 14), this novel approach holds promise for impact across the majority of mutant KRAS alleles, including the two most prevalent variants, G12D and G12V. Interestingly, our data suggest that tumors harboring codon 61 mutations (such as Q61H) appear to be less sensitive to SOS1 inhibition, possibly because these mutant isoforms have the lowest intrinsic GTPase activity and may require less upstream signaling to remain GTP bound (41). The KRAS\textsuperscript{G12R} variant, which is relatively common in pancreatic cancers (~20% prevalence), showed no modulation of pERK following treatment with BI-3406. This finding is in line with a recent publication describing an inability of the catalytic domain of SOS1 to interact with this KRAS\textsuperscript{G12R}, mutant oncoprotein (31). The sensitivity spectrum we have observed toward SOS1 inhibition further supports the concept of oncogenic KRAS G12 and G13 variants functioning in a semiautonomous manner (42) and remaining susceptible to regulation by SOS1 for optimal GTP loading. Collectively, our data suggest that BI-3406 will be able to affect about 80% to 90% of all KRAS-driven cancers.

We have carried out a comprehensive screening for effective combination partners. Synergy was observed upon combination of SOS1 with MEK inhibitors, leading to tumor regressions in multiple mutant KRAS-driven cancer models at well-tolerated doses. Of the two SOS isoforms, SOS1 and SOS2, only SOS1 was phosphorylated by ERK, resulting in the reduction of its GEF activity (26). Treatment with a MEK inhibitor reduces the activity of ERK1/2, resulting in release of a negative feedback loop, thus increasing the activity of SOS1-mediated formation of GTP-loaded KRAS (25, 26). Combination of MEK inhibitor with BI-3406 thus blocks the negative feedback release by reducing pMEK1/2 and pERK1/2 levels, supporting sustained pathway inhibition and tumor regressions (Fig. 5D). Tumor stasis was observed with a SOS1/MEK inhibitor combination in colorectal and pancreatic cancer PDX models. This may indicate that, in these tumor types, additional feedback and bypass mechanisms are effective, and triple combinations are needed to shut down KRAS signaling and achieve tumor regressions. Because of the favorable tolerability of the SOS1/MEK treatment, combinations with standard-of-care treatments will be further evaluated with the aim to achieve tumor regressions in colorectal and pancreatic cancer models. We demonstrated that, in addition to the combination of BI-3406 with trametinib (MEK inhibitor), the combination of BI-3406 with the clinical KRAS\textsuperscript{G12C} inhibitor AMG 510 results in enhanced and prolonged MAPK pathway suppression. Our study highlights SOS1 inhibitors as promising combination partners for inhibitors directly targeting KRAS, the GDP-bound form of KRAS, or downstream MAPK pathway intermediates. This finding is also in line with a recent report describing a marked synergy in NSCLC cell lines combining SOS1 inhibition with vertical EGFR inhibition (43).

BI-3406 is a selective inhibitor of SOS1 and does not target the paralog GEF SOS2. Simultaneous genetic inactivation of both SOS proteins leads to rapid death in mouse models, in contrast to single-gene perturbations (44). Thus, although the SOS1 selectivity may reduce the monotherapy impact of BI-3406 on KRAS and the MAPK pathway, it can facilitate combination therapies because of the expected superior tolerability of a SOS1-specific inhibitor compared with a pan-SOS1/SOS2 inhibitor (44, 45). Furthermore, targeting SOS1 can selectively exploit its key function in adaptive feedback control that is not shared with its paralog SOS2 (25, 26). No upregulation of SOS2 expression was observed in our biomarker and efficacy experiments. It remains to be determined whether patients with cancer treated with a SOS1 inhibitor will exhibit induction of SOS2 levels.

Recently, inhibitors targeting the protein-tyrosine phosphatase SHP2 (encoded by the gene \textit{PTPN11}), a common node downstream of RTKs that is required for RAS activation, have been reported (18, 19). Interestingly, these reports also suggest that inhibition of SHP2 can attenuate adaptive MEK inhibitor resistance in KRAS-dependent cancers (46–49). Our comparative analysis of the SHP2 inhibitor tool compound SHP099 suggested activity in cell lines harboring G12C, a
subset of G12D, and possibly G12S KRAS variant–driven cell lines, while the SOS1 inhibitor BI-3406 demonstrated activity in all KRAS G12- and G13-mutant cell lines tested, with the exception of cell lines driven by the G12R oncoprotein. While only BI-3406 was active in the KRAS G13-driven context, both inhibitors lacked single-agent activity in KRAS Q61-mutant cell lines, suggesting an overall broader impact on KRAS-mutant cancers by the SOS1 inhibitor. Future studies will be required to compare and contrast the capabilities of SOS1 and SHP2 inhibitors to overcome adaptive resistance to KRAS/RAF/MEK/ERK-targeted agents across KRAS-driven cancers.

Although the precise mechanism by which SHP2 contributes to KRAS activation is yet to be determined, SHP2 is not a direct activator of KRAS and may in part act via SOS1 (50, 51). Ongoing clinical evaluations will show whether SOS1 inhibitor–MEK inhibitor and SHP2 inhibitor–MEK inhibitor combinations will differ in terms of safety and response rates across tumors with different KRAS alterations.

Collectively, our study provides a new chemical probe for further dissection of the cellular functions of SOS1 in tumorigenesis and MEK inhibitor–driven drug resistance. Importantly, the pharmacologic properties of BI-3406 and close analogues hold the promise of developing clinical SOS1 compounds that, in combination with MEK inhibitors and potentially other RTK/MAPK pathway inhibitors, could provide significant clinical benefit across a broad patient population currently lacking molecularly targeted, precision medicine options. A phase I clinical trial has been initiated (NCT04111458) for patients with advanced KRAS-mutated cancers to evaluate safety, tolerability, pharmacokinetics (NCT04111458) for patients with advanced KRAS-mutated cancers to evaluate safety, tolerability, pharmacokinetics, and pharmacodynamics properties, and preliminary efficacy of BI 1701963, a SOS1–KRAS inhibitor closely related to BI-3406, alone and in combination with the MEK inhibitor trametinib.

**METHODS**

Additional descriptions of methods can be found in the Supplementary Data.

**Cell Culture**

Tumor cell lines were obtained from the ATCC or the German Collection of Microorganisms and Cell Culture (DSMZ). All cell lines used in this study were cultured according to the manufacturer’s instructions and authenticated by short tandem repeat analysis at Boehringer Ingelheim (Supplementary Table S9). With regard to the 2-D proliferation assays, cells were seeded in their respective medium supplemented with 2% FCS. For the 3-D proliferation assay, the cells were embedded in soft agar, which required three separate layers within a well; a bottom layer formed of 1% agar solution, a cell layer formed of 0.3% agar solution, and a medium layer (described in detail in the Supplementary Data). BI-3406, trametinib, or a positive control (e.g., panobinostat) was added with increasing concentrations. Briefly, 6 × 10⁵ cells were seeded in six wells and grown to 70% confluence. Cells were washed with ice-cold PBS and lysed in 80 μL of an lysis cocktail. Lysates were quickly frozen in liquid nitrogen and stored at −80°C until further usage. After normalizing protein concentrations, 40 μg of protein was added in duplicates to wells of the RADY Bi-Lisa plate coated with RADY-GTP-binding protein, and incubated at 4°C for 30 minutes while shaking at 400 rpm. After washing, antigen-presenting buffer was added for 2 minutes. To measure bound RAS-GTP levels, wells were subsequently incubated with the provided protease inhibitor cocktail. Lysates were quickly frozen in liquid nitrogen and stored at −80°C until further usage. After normalizing protein concentration, 40 μg of protein was added in duplicates to wells of the RADY Bi-Lisa plate coated with RADY-GTP-binding protein, and incubated at 4°C for 30 minutes while shaking at 400 rpm. After washing, antigen-presenting buffer was added for 2 minutes. To measure bound RAS-GTP levels, wells were subsequently incubated with an anti-RAS primary antibody (1:50) followed by a horseradish peroxidase (HRP)–labeled secondary antibody (1:500), and finally by adding an HRP detection reagent. Absorbance was measured at 490 nm using an EnSpire Multimode Reader (PerkinElmer). Background was determined by a negative control well and subtracted from all samples. The same assay was used to determine amount of RAS-GTP levels in tumor lysate.

**Measurement of KRAS-GTP Levels**

RAS-GTP levels were analyzed using a RAS G-LISA Assay Kit (Cytoskeleton Inc., #BK131) according to the manufacturer’s instructions. Briefly, 6 × 10⁵ cells were seeded in six wells and grown to 70% confluence. Cells were washed with ice-cold PBS and lysed in 80 μL of an lysis buffer supplemented with the provided protease inhibitor cocktail. Lysates were quickly frozen in liquid nitrogen and stored at −80°C until further usage. After normalizing protein concentration, 40 μg of protein was added in duplicates to wells of the RADY Bi-Lisa plate coated with RADY-GTP-binding protein, and incubated at 4°C for 30 minutes while shaking at 400 rpm. After washing, antigen-presenting buffer was added for 2 minutes. To measure RAS-GTP levels, wells were subsequently incubated with an anti-RAS primary antibody (1:50) followed by a horseradish peroxidase (HRP)–labeled secondary antibody (1:500), and finally by adding an HRP detection reagent. Absorbance was measured at 490 nm using an EnSpire Multimode Reader (PerkinElmer). Background was determined by a negative control well and subtracted from all samples. The same assay was used to determine RAS-GTP levels in tumor lysate.

**Synthesis of BI-3406**

Synthesis conditions are described in the Supplementary Data. A schematic representation of the synthesis can be found in Supplementary Fig. S1E and S1F.

**Protein–Protein Interaction Assays**

Details on protein expression and purification can be found in Supplementary Data. Measurements of various protein–protein interactions were performed using the AlphaScreen Technology developed by PerkinElmer. Recombinant KRAS proteins, based on KRAS isoform 4B (UniProt ID, P01116–2), were: KRASG12D (1–169, N-terminal 6His-tag, C-terminal avi-tag) from Xtal BioStructures, Inc. and KRASG12C (1–169, N-terminal avi-tag, biotylated, mutations: C515S, C80L, and C118S). Biotinylation was performed in vitro with recombinant BirA biotin-protein ligase as recommended by the manufacturer (Avidity LLC). Interacting proteins such as SOS1 (564–1,049, N-terminal GST-tag, TEV cleavage site), and SOS2 (562–1,047, N-terminal GST-tag, TEV cleavage site) were expressed as glutathione S transferase (GST) fusions. Accordingly, the alpha screen beads were glutathione-coated Alpha Blue Acceptor Beads (PerkinElmer, AL 109 R) and Alpha Screen Streptavidin-conjugated Donor Beads (PerkinElmer, 6760002L). Nucleotide was purchased from Sigma (GDP #G7127) and Tween-20 from Bio-Rad ( #161-0781). All interaction assays were carried out in PBS, containing 0.1% BSA, 0.05% Tween-20, and 10 μmol/L GDP. Assays were carried out in white ProxiPlate-384 Plus Plates (PerkinElmer, #6008280) in a final volume of 20 μL. In brief, biotinylated KRAS proteins (10 nmol/L final concentration) and GST-SOS1 or GST-SOS2 (10 nmol/L final concentration) were mixed with glutathione acceptor beads (5 μg/mL final concentration) in buffer containing GDP and were incubated for 30 minutes at room temperature. After addition of streptavidin donor beads (5 μg/mL final concentration) under green light, the mixture was further incubated for 60 minutes in the dark at room temperature. Single oxygen-induced fluorescence was measured at an EnSpire Multimode Plate Reader (PerkinElmer) according to the manufacturer’s recommendations. Data were analyzed using the GraphPad Prism–based data software.
Biomarker and Pharmacokinetics/Pharmacodynamics Analysis

pERK and pAkt modulation in tumors was determined using the Phospho/Total ERK1/2, Phospho(Ser473)/Total Akt, and Phospho-AKT (Thr308) Whole Cell Lysate Kits (Meso Scale Diagnostics, K15107D, K15100D, and K151D1D). Tumors were homogenized using Ready Prep Mini Grinders (#163-2146, Bio-Rad) and lysed in MSD TRIS lysis buffer plus inhibitors (as provided in the kit). Protein concentration was determined by Bradford analysis. 0.8 μg/μl protein lysate was used for pERK measurements (biological replicates) according to the recommendations of the manufacturer. Signal intensities were measured using a Meso SECTOR S 600 reader. The pERK to total ERK ratio was calculated and the data were plotted in GraphPad Prism. This assay was also used for measurement of pharmacodynamics modulation in several tumor cell lines (Supplementary Fig. S2E).

pERK levels were determined in mouse skin based on IHC staining (H-scores). IHC was performed on formalin-fixed, paraffin-embedded tissue, 3 μm sections using anti-Phospho-p44/42 MAPK (ERK1/2) (Thr202/Tyr204) (1:40, Cell Signaling Technology). Antibody incubation and detection were carried out at 37°C. Antigen retrieval was performed using Thermo PT Module with buffer, pH 6 (Dako, #K8005), and visualized using the EnVision Kit (Dako). Appropriate positive and negative controls were included with the study sections. Digital images of whole-tissue sections were acquired using a Aperio AT2 Histology Scanner (Leica Microsystems). Images were evaluated by a pathologist with 20× objective. IHC scores were generated using HALO Software 3.0, (F. Trapani) and H-score was generated using HALO Software 3.0, Scanner (Leica Microsystems). Images were evaluated by a pathologist.

Cell Line–Derived Efficacy Studies and Biomarker Studies in Mice

All animal studies were approved by the internal ethics committee and the local governmental committee. Group sizes in efficacy studies were selected after performing power analysis. Female BomTac:NMRI-Fem1 × 10⁶ mice were used in all xenograft studies. For biomarker and efficacy experiments with MIA PaCa-2 tumors, female mice were engrafted subcutaneously with 10 × 10⁶ cells suspended in Matrigel. In the case of biomarker studies with MIA PaCa-2 tumors, mice were randomized by tumor size in groups of 5 mice once tumors reached a size of 170–500 mm³. Mice were treated once at timepoint 0 and 6 hours. Tumors were explanted and snap-frozen to analyze biomarker modulation. Details of bioanalysis of mouse blood samples can be found in the Supplementary Data.

In the case of efficacy experiments, mice were randomized in groups (n = 7 mice/treatment group) by tumor size by the automated data storage system Sepia on day 7 (Fig. 3D) or 12 (Fig. 4A) once tumors reached a size between 95 and 180 mm³. Compound treatment was initiated after randomization based on body weight. Tumor size was measured by an electronic caliper, and body weight was monitored daily. The analysis largely followed the procedures described in refs. 53 and 54. Number of subcutaneous cells injected and size of tumor for randomization was as follows with a group size of 7 to 10 mice: A549 (10 × 10⁶ cells; 62–150 mm³; n = 7, Fig. 3F; Supplementary Data; Supplementary Fig. S3G and S3H), LoVo (10 × 10⁶ cells; 123–173 mm³; n = 7, Figs. 3G and 4F), SW620 (5 × 10⁶ cells, 80–125 mm³, Fig. 3F), and A375 (5 × 10⁶ cells, 64–149 mm³, n = 6–7, Fig. 3F). In the case of the biomarker studies with A549 cells, mice were randomized once tumors reached a size between 209 and 320 mm³ (Fig. 3B and C; Supplementary Data). BI-3406 was dissolved in 0.5%Natrosol. Trametinib was dissolved in 0.5% DMSO and 0.5% Natrosol. The control group was treated with 0.5% of Natrosol orally in the same frequency as the treatment groups (twice daily). All compounds were administered intragastrically by gavage (10 mL/kg). Details on formulation of compounds can be found in the Supplementary Data.

PDX Studies

PDX model characterization and profiling are described in detail in the Supplementary Data and Supplementary Table S13. PDX tumor fragments (4 × 4 × 4 mm³) were implanted on the right hind flanks of NSG female mice purchased from the Jackson Laboratory and allowed to grow to an average volume of 100–250 mm³ as monitored by caliper measurements. At enrollment, animals were randomized and treated orally on a 5 days on/2 days off schedule for convenience, to avoid weekend treatments, with vehicle (0.5% Natrosol) twice a day (6 hours apart), BI-3406 (SOS1 inhibitor) at 50 mg/kg twice a day (6 hours apart), trametinib at 0.1 mg/kg twice a day (6 hours apart), or the combination thereof. Mice were 11 weeks old and treatment group sizes included at least 5 to 7 mice per group. All animals received LabDiet 5053 chow ad libitum. Trametinib was purchased from Chemietek. In the PDX studies, tumor growth was monitored twice a time with calipers and the tumor volume (TV) was calculated as TV = (D × d^2)/2, where “D” is the largest and “d” is the smallest superficial visible diameter of the tumor mass. All measurements were documented as mm³. Body weights were measured twice weekly and used to adjust dosing volume and monitor animal health. RPPA analysis of explanted tumor material is described in detail in the Supplementary Data (Supplementary Fig. S5I; Supplementary Table S8).

RNA Isolation and Sequencing Library Preparation for Expression Profiling

Cells were lysed in TRI Lysis Reagent (Qiagen, #79306) according to the manufacturer’s instructions. Instead of chloroform, 10% vol/vol 1-bromo-2-chloropropane (Sigma-Aldrich, #B9673) was added. Total RNA was isolated with RNAeasy Mini Kit (Qiagen, #79306). Quant-seq libraries were prepared using the QuantSeq 3′ mRNA-Seq Library Prep Kit FWD for Illumina from Lexogen (#015.96) according to the manufacturer’s instructions. Samples were subsequently sequenced on an Illumina NextSeq 500 System with a single-end 76 bp protocol. Single-end sequencing reads from grafted samples were filtered into human and mouse reads using DisAmbiguate (52), based on mapping to hg38 and mm10. The filtered reads were then processed with a pipeline building upon and extending the implementation of the ENCODE “Long RNA-seq” pipeline. Additional details on the methods are outlined in the Supplementary Data.

Whole-Exome Sequencing

In-house DNA libraries were prepared using the Agilent SureSelectXT Human AllExon 50 Mb Enrichment Kit and subsequently sequenced on an Illumina HiSeq 2000 with a 100 bp paired-end protocol. Sequencing data from in-house cell lines were completed with data retrieved from the Cancer Cell Line Encyclopedia (CCLE) and Catalogue of Somatic Mutations in Cancer (COSMIC).

Analysis of Gene Expression by QuantiGene Single Plex Technology (Affymetrix)

RNA was isolated from tumors as described above. The following probes were used: DUSP6 (SA-11958) and GAPDH (SA-10001). The analysis was performed according to the manufacturer’s recommendations. The DUSP6 levels of the individual tumors were normalized to their respective GAPDH levels.

Variant Calling from Whole-Exome Sequencing Data (DNA Sequencing)

Paired-end sequencing reads were mapped against the human genome hg38 using bwa. We used strelka2 and the Ensemble Vari-
Statistical Analysis
Statistical analyses and bioinformatics analysis were performed with R version 3.5.0 and Bioconductor 3.7 or GraphPad Prism. A Fisher exact test was used for computing the associations of gene mutations with the sensitivity status of cell lines and for comparison of tumor volumes from the control group with one treatment group. For calculations of tumor volume, absolute values were used for statistical analysis. Because of the observed variation, nonparametric methods were applied. In case several treatment groups were compared, one-sided nonparametric Mann-Whitney–Wilcoxon U tests were applied to compare treatment groups with the control, as reduced tumor growth was expected following treatment. The P values for the tumor volume (efficacy parameter) were adjusted for multiple comparisons according to Bonferroni-Holm within each subtopic (comparisons vs. control, comparisons monotherapies vs. combination therapy), whereas the P values of the body weight (tolerability parameter) remained unadjusted in order not to overlook a possible adverse effect. The level of significance was fixed at α = 5%. An (adjusted) P value of less than 0.05 was considered to show a statistically significant difference between the groups, and differences were seen as indicative whenever 0.05 ≤ P < 0.10. Data are represented as dot plots with bar graphs for mean±SD or SEM, as indicated. In the case of the PDX experiments, statistical significance was determined using an unpaired t test per row and the Holm–Sidak method to correct for multiple comparisons (Fig. 4E and F; Supplementary Fig. S4D–S4F).

Data Availability
Atomic coordinates and structure factors for the cocrystal X-ray structures of BI-68BS and BI-3406 and SOS1 have been deposited at the PDB under accession nos. 6SER (BI-68BS) and 6SCM (BI-3406). Data are available in Supplementary Table S1. Expression data generated and analyzed in this study have been deposited in the Gene Expression Omnibus database under the accession no. GSE128385. Processed data are available in Supplementary Table S7.

Disclosure of Potential Conflicts of Interest
M.H. Hofmann reports grants from FFF (the work was supported by the Austrian Research Promotion Agency) and personal fees from Boehringer Ingelheim RCV GmbH & Co KG (full time employee) during the conduct of the study, as well as has been listed as inventor on patent applications for SOS1 inhibitors. M. Gmachl reports personal fees from Boehringer Ingelheim RCV GmbH & Co KG (full time employee) during the conduct of the study, as well as has been listed as inventor on patent applications for SOS1 inhibitors. J. Ramharter reports grants from FFF (the work was supported by the Austrian Research Promotion Agency) and personal fees from Boehringer Ingelheim RCV GmbH & Co KG (full time employee) during the conduct of the study. C. Kofink reports grants from FFF (the work was supported by the Austrian Research Promotion Agency) and personal fees from Boehringer Ingelheim RCV GmbH & Co KG (full time employee) during the conduct of the study. D.-A. Botesteanu reports grants from FFF (the work was supported by the Austrian Research Promotion Agency) and personal fees from Boehringer Ingelheim RCV GmbH & Co KG (full time employee) during the conduct of the study. D. Gerlach reports grants from FFG (the work was supported by the Austrian Research Promotion Agency) and personal fees from Boehringer Ingelheim RCV GmbH & Co KG (full time employee) during the conduct of the study. N. Pototschnig reports personal fees from Boehringer Ingelheim RCV GmbH & Co KG (full time employee) during the conduct of the study. F. Trapani reports grants from FFG (the work was supported by the Austrian Research Promotion Agency) and personal fees from Boehringer Ingelheim RCV GmbH & Co KG (full time employee) during the conduct of the study. N. Rumpel reports personal fees from Boehringer Ingelheim and he has been listed as inventor on patents for SOS1 inhibitors. J. Böttcher reports personal fees from Boehringer Ingelheim RCV GmbH & Co KG (full time employee) during the conduct of the study and Forma Therapeutics at the time the work was performed. A. Zoephel reports grants from FFG (the work was supported by the Austrian Research Promotion Agency) and personal fees from Boehringer Ingelheim RCV GmbH & Co KG (full time employee) during the conduct of the study. S.-C. Fu reports other from Boehringer Ingelheim (sponsored research) during the conduct of the study. J. Bottcher reports personal fees from Boehringer Ingelheim RCV GmbH & Co KG (full time employee) during the conduct of the study. S. Lieb reports grants from FFF (the work was supported by the Austrian Research Promotion Agency) and personal fees from Boehringer Ingelheim RCV GmbH & Co KG (full time employee) during the conduct of the study. C.P. Vellano reports other from Boehringer Ingelheim (sponsored research) during the conduct of the study and outside the submitted work, as well as part of this work was performed under a sponsored research collaboration between MD Anderson and Boehringer Ingelheim, for which the latter provided funding support. J.C. O’Connell reports other from Forma Therapeutics (reports employment with Forma Therapeutics at the time the work was performed) during the conduct of the study and Forma Therapeutics (Forma Therapeutics collaboration with Boehringer Ingelheim) outside the submitted work. J. Moll reports full time employment with Boehringer-Ingelheim RCV GmbH & Co KG. M. Petronzelli reports other from Boehringer Ingelheim (full time employee) during the conduct of the study. T.P. Heffernan reports other from Boehringer Ingelheim (sponsored research) during the conduct of the study and personal fees from Culligen Inc. outside the submitted work. D.B. McConnell reports grants from FFF (the work was supported by the Austrian Research Promotion Agency) and personal fees from Boehringer Ingelheim RCV GmbH & Co KG (full time employee) during the conduct of the study. N. Kraut reports grants from Boehringer Ingelheim (the work was supported by the Austrian Research Promotion Agency) during the conduct of the study, as well as reports full time employment with Boehringer Ingelheim RCV GmbH & Co KG. No potential conflicts of interest were disclosed by the other authors.
Authors’ Contributions

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Pan-KRAS–SOS1 Protein–Protein Interaction Inhibitor BI-3406


BI-3406, a Potent and Selective SOS1–KRAS Interaction Inhibitor, Is Effective in KRAS-Driven Cancers through Combined MEK Inhibition

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