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Immune checkpoint blockade has achieved significant therapeutic success for a subset of cancer patients; however, a large portion of cancer patients do not respond. Unresponsive tumors are characterized as being immunologically “cold,” indicating that these tumors lack tumor antigen-specific primed cytotoxic T cells. Sitravatinib is a spectrum-selective tyrosine kinase inhibitor targeting TAM (TYRO3, AXL, MerTK) and split tyrosine-kinase domain–containing receptors (VEGFR and PDGFR families and KIT) plus RET and MET, targets that contribute to the immunosuppressive tumor microenvironment. We report that sitravatinib has potent antitumor activity by targeting the tumor microenvironment, resulting in innate and adaptive immune cell changes that augment immune checkpoint blockade. These results suggest that sitravatinib has the potential to combat resistance to immune checkpoint blockade and expand the number of cancer patients that are responsive to immune therapy.

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Sitravatinib potentiates immune checkpoint blockade in refractory cancer models

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Introduction

Immune checkpoint blockade has achieved significant therapeutic success for a subset of cancer patients; however, a large portion of cancer patients do not respond. Unresponsive tumors are characterized as being immunologically “cold,” indicating that these tumors lack tumor antigen-specific primed cytotoxic T cells. Sitravatinib is a spectrum-selective tyrosine kinase inhibitor targeting TAM (TYRO3, AXL, MerTK) and split tyrosine-kinase domain–containing receptors (VEGFR and PDGFR families and KIT) plus RET and MET, targets that contribute to the immunosuppressive tumor microenvironment. We report that sitravatinib has potent antitumor activity by targeting the tumor microenvironment, resulting in innate and adaptive immune cell changes that augment immune checkpoint blockade. These results suggest that sitravatinib has the potential to combat resistance to immune checkpoint blockade and expand the number of cancer patients that are responsive to immune therapy.
highlight the potential immune-activating effects of sitravatinib. We sought to investigate the induction of immune activity by sitravatinib and glesatinib in syngeneic murine cancer models. We found that inhibition of TAM receptor activation by the compounds reduces expression of markers associated with immune suppression in macrophages in a MerTK-dependent manner. Additionally, sitravatinib and glesatinib showed potent single-agent antitumor efficacy and enhanced the activity of PD-1 blockade through promoting an antitumor immune microenvironment.

**Results**

*MerTK inhibition with sitravatinib directly affects macrophage phenotype.* Since sitravatinib potently inhibits TAM receptors, which are important in mediating an immunosuppressive macrophage phenotype (5), we investigated the effect of sitravatinib on cytokine-stimulated polarization of primary bone marrow–derived macrophages (BMDMs). We found that sitravatinib inhibited the expression of *Arg1* stimulated by IL-4 and the expression of *Il-6* and *Il-12* induced by LPS treatment (Figure 1). To better mimic the cytokine milieu found in tumors, we stimulated BMDMs with LPS (Figure 1A) or IL-4 (Figure 1B) in the presence of conditioned media (CM) from murine lung cancer (KLN205) cells. Treatment with sitravatinib dose-dependently inhibited the IL-4 plus CM–mediated expression of *Arg1*, *Ym-1*, and *Fizz-1*, markers associated with an immunosuppressive macrophage phenotype, but did not affect the LPS plus CM–induced expression of *Tnfa*, *Il-6*, or *Il-12*, markers of an immunostimulatory macrophage phenotype (15). These data suggest that sitravatinib can prevent monocytes from polarizing into immunosuppressive macrophages in the presence of potent immune-suppressive signals. The data also demonstrate that sitravatinib did not abrogate the induction of an immunostimulatory macrophage phenotype. Similar results were obtained with
glesatinib, a small molecule that targets the TAM receptors and MET (Supplemental Figure 1, A and B; supplemental material available online with this article; https://doi.org/10.1172/jci.insight.124184DS1). To define whether MerTK activity was critical for the effect of the compounds on macrophage polarization, BMDMs from MerTK-deficient (MerTK–/–) animals were subjected to the experimental conditions described above. MerTK–/– BMDMs were unaffected by sitravatinib or glesatinib, indicating that MerTK is critical to the effect of the compounds on macrophage polarization (Figure 1 and Supplemental Figure 1, A and B). These data also suggest that the presence of Gas6 or protein S is important in regulation of macrophage phenotype.

Sitravatinib has potent antitumor activity in vivo. To determine the single-agent antitumor efficacy of sitravatinib, we administered the compound to immunocompetent mice bearing KLN205, CT1B-A5, or E0771 tumors (Figure 2, A–C). In each model, sitravatinib significantly inhibited tumor progression and induced tumor regression. Glesatinib also showed single-agent activity in each tumor model (Supplemental Figure 2, A–C). We observed no adverse effects of the compounds but noted that treatment with sitravatinib or glesatinib reduced tumor-induced splenomegaly, suggestive of immune modulatory activity.

To demonstrate the effect of the compounds on tumor cell viability, we performed in vitro colony-forming and MTS viability assays. Each compound reduced colony formation in a dose-dependent manner (Figure 2D and Supplemental Figure 2D) and inhibited tumor cell viability with an IC₅₀ of approximately 1 μM (Figure 2E and Supplemental Figure 2E), a concentration considerably higher than the predicted plasma concentration of sitravatinib (10 nM) after dosing at 20 mg/kg (Supplemental Table 1). These data suggest that the potent antitumor activity observed was unlikely due solely to direct tumor cell killing but related to microenvironmental changes induced by sitravatinib.

To determine the effect of the compounds on MerTK activity in vivo, we probed lysates from treated KLN205 tumors for active MerTK and found that both compounds suppressed MerTK phosphorylation (Figure 3A; see complete unedited blots in the supplemental material), with sitravatinib showing a stronger effect. Histologic analysis of KLN205 tumors demonstrated that sitravatinib-treated tumors...
showed increased necrosis, elevated cleaved caspase-3 (CC3), decreased Ki67 expression, and reduced microvessel density (CD31), while tumor cell death induced by glesatinib was less substantial and it did not reduce microvessel density (Figure 3B). These results suggest both compounds target critical pathways in the tumor microenvironment that support tumor progression.

*Sitravatinib alters the immune landscape of KLN205 tumors to favor immune checkpoint blockade.* Given the potent antitumor activity of sitravatinib and the fact that TAM receptors contribute to the immune microenvironment in tumors, we performed gene expression analysis (NanoString Technologies) using tumor tissue collected from mice bearing KLN205 tumors treated with control, glesatinib, or sitravatinib. Compared with those treated with glesatinib, we found that sitravatinib-treated tumors had more significant gene set changes. Multiple genes associated with angiogenesis were among the most downregulated in sitravatinib-treated tumors, consistent with its blockade of VEGFR family RTKs, while inflammatory genes were among the highest upregulated in treated tumors (Figure 4, A and B). Interestingly, the expression of Pd-l1 (Cd274) was elevated by glesatinib and sitravatinib, suggesting that the compounds may enhance response to immune checkpoint blockade (16). This is supported by analysis of gene program/pathways that were altered by sitravatinib, which included multiple pathways associated with innate and adaptive immune response (Figure 4C).

To further investigate the effect of sitravatinib on the immune landscape of KLN205 tumors, flow cytometry was performed on tumors and spleens from mice treated for 6 days (Figure 5). Sitravatinib significantly
reduced the number of tumor-associated immunosuppressive myeloid cells, including monocytic MDSCs and M2 macrophages. Sitravatinib also increased the number of CD4+ T cells and exhausted CD8+ T cells characterized by PD-1 and CTLA-4 expression. In comparison, we found that spleens from sitravatinib-treated mice had an increase in CD4+ T cells and proliferating CD8+ T cells. The immune cell changes in the tumors and the spleen suggested that sitravatinib might enhance the effect of PD-1 blockade. Although glesatinib had similar trends (Supplemental Figure 3, A–C), the immune cell changes were not as significant as seen with sitravatinib. These results might be explained by the less potent activity of glesatinib against MerTK or through contribution of other distinct sitravatinib RTK targets.

Sitravatinib enhances the efficacy of PD-1 blockade. To test the combinatorial effect of sitravatinib and PD-1 blockade, we treated cohorts of KLN205 or E0771 tumor-bearing mice with the compound with or without anti–PD-1. Mice were treated for 2.5 weeks in each model. Sitravatinib significantly enhanced the efficacy of PD-1 blockade (Figure 6, A and B). Glesatinib treatment also enhanced the efficacy of anti–PD-1 therapy (Supplemental Figure 4, A and B) but was less potent that sitravatinib. In fact, 2 of 14 mice bearing E0771 tumors treated with anti–PD-1 plus sitravatinib showed complete remission and stayed tumor free for 50 days. To test whether these 2 animals developed an adaptive antitumor response with immune memory, we rechallenged the animals by implanting E0771 cells in the mammary fat pad (on the opposite side from the original injection). Control mice were also implanted with E0771 cells and showed robust and predictable tumor formation and growth. However, the 2 animals that showed complete response to sitravatinib and anti–PD-1 failed to grow E0771 tumors upon rechallenge (Figure 6C).

Discussion

Our study has found that sitravatinib and glesatinib have potent single-agent antitumor activity; moreover, sitravatinib can alter the immune landscape of tumors to favor immune checkpoint blockade. In 3 different preclinical tumor models, including KLN205, an aggressive lung cancer model that is resistant to checkpoint blockade, the tumor response to sitravatinib and glesatinib was rapid and consistent (Figure 2, A–C, and Supplemental Figure 2, A–C). Consistently, sitravatinib was also found to have potent antitumor activity in preclinical models of sarcoma (17).
Our in vitro studies demonstrate that each compound has direct antitumor cell effects, albeit at markedly higher concentrations than required for in vivo efficacy. The major targets of each compound are pathways associated with microenvironmental features of tumors, and the predicted in vivo concentration of the compounds is substantially below the observed in vitro IC50 values. We therefore propose that the potent antitumor activity of each compound is largely due to the effect of sitravatinib and glesatinib on the tumor microenvironment. Therapies against some of these targets, for example, angiogenesis, have been explored for decades with limited success in solid tumors (18). A major unmet challenge with the use of targeted therapeutic agents is the propensity of tumor cells to activate alternative pathways that lead to therapeutic resistance. Sitravatinib and glesatinib have a relatively broad spectrum of targets that are inhibited at pharmacologically achievable doses. It is possible that inhibition of multiple aspects of the tumor microenvironment reduces the chance of resistance-derived inhibition of a single but important biologic process.

What may distinguish sitravatinib and glesatinib relative to other multireceptor tyrosine kinases is their activity against the TAM receptors. Of the multireceptor tyrosine kinase inhibitors in active clinical development, sitravatinib and glesatinib are arguably the most effective TAM receptor inhibitors currently available. We propose that the inhibition of TAM receptors enables the compounds to alter the immune landscape. This is the first report to our knowledge of the effect of sitravatinib and glesatinib on the immunosuppressive tumor microenvironment, as evidenced in vitro by observations using primary BMDMs, in vivo through gene expression and flow cytometry, and functionally through the potentiation of immune checkpoint blockade. Since resistance to immune therapy remains an unmet challenge in the treatment of many cancers, these compounds may offer therapeutic benefit that improves response to immune checkpoint blockade.
Methods

Cell lines. KLN205, a murine lung cancer cell line, was obtained from ATCC (CRL-1453), and E0771, a murine breast cancer cell line, was a gift from Philip Thorpe (University of Texas Southwestern Medical Center). CT1B-A5, an isogenic pancreatic cancer cell line, was derived from \( KPfC \) \((Kras^{LSL-G12D}; Trp53fl/fl; PDX\ Cre/+)) mice. Cells were cultured in DMEM (Invitrogen) or RPMI (Invitrogen) containing 5% FBS and maintained at 37°C in a humidified incubator with 5% CO2 and 95% air and confirmed to be pathogen free before use.

Animal studies. KLN205 cells \((0.5 \times 10^6)) were injected subcutaneously into 6-week-old DBA/2 mice, CT1B-A5 cells \((1 \times 10^6)) were injected subcutaneously into 6-week-old C57BL/6 mice, and E0771 cells \((0.5 \times 10^6)) were injected orthotopically into the mammary fat pad of 6-week-old female C57BL/6 mice. Therapy was initiated in mice with a tumor volume of 300 mm\(^3\) (KLN205) or 500 mm\(^3\) (E0771) and included control (Ctrl, vehicle, once per day), anti–PD-1 (PD-1, i.p. 10 mg/kg, every 3 days), sitravatinib (sitra, p.o. 20 mg/kg, once per day), or anti–PD-1 in combination with sitravatinib at the indicated dose. Mice were treated for 2.5 weeks. \(^{**}P < 0.01\) anti–PD-1 in combination with sitravatinib vs. sitravatinib alone by t test. Two of fourteen mice bearing E0771 tumors treated with the combination therapy showed complete remission and stayed tumor free for 50 days. (C) Rechallenge growth curve of the 2 tumor-free animals from B. We injected \(0.5 \times 10^6\) E0771 cells orthotopically into the mammary fat pads of 6-week-old female C57BL/6 mice (Ctrl, \(n = 4\)) and on the contralateral side from the original injection of the 2 tumor-free animals.

Real-time quantitative PCR. BMDMs were harvested from WT C57BL/6 or \(MerTK^{-/-}\) mice using standard techniques (19). To determine if glesatinib and sitravatinib alter the expression of markers of macrophage polarization, BMDMs were stimulated for 2 hours with 20 ng/ml LPS or 18 hours with 40 ng/ml IL-4, immune stimulatory or immune-suppressive signals, respectively, and different concentrations of
sitravatinib or glesatinib in the presence or absence of CM collected from KLN205 cells. RNA was extracted using Qiagen RNeasy Mini Kit (74106), cDNA was synthesized using a Bio-Rad iScript cDNA synthesis kit. The expression of TNF-α, IL-6, IL-12, arginase 1, YM-1, and Fizz-1 was measured by SYBR-Green dependent q-PCR (Supplemental Table 2). Three independent experiments were performed. Duplicates were run in each experiment.

**MTS assay.** The MTS colorimetric assay (Promega) was performed as per the manufacturer’s instructions. This assay is based on the conversion of MTS into soluble formazan by endogenous dehydrogenase enzymes found in metabolically active cells. Cells were plated at 2,000 cells/well in tissue culture–treated 96-well plates. The following day, sitravatinib or glesatinib was added to each plate in a dilution series across the plate, such that 8 different concentrations of the drug were tested. On day 5, 20 μl MTS was added, followed by a 2-hour incubation at 37°C, and then the absorbance was read at 490 nm on a plate reader. The 96-well plate data were imported into an in-house database of in vitro drug sensitivity assays (DIVSA); IC₅₀ values were calculated using the DRC model (R package “drc”; http://www.bioassay.dk).

**Liquid colony-forming assay.** Cell lines were cultured in 6-well tissue culture plates at low density (200 cells/well for E0771 and 500 cells/well for KLN205) in 2 ml media. Cells were plated in respective culture conditions and allowed to settle for 1 week or until marked colony formation. Cells were then fixed with 10% formalin and stained with crystal violet. Images were analyzed with ImageJ (NIH) or NIS Elements.

**Western blot analysis.** KLN205 tumor tissues from mice treated for 9 days were lysed using RIPA buffer (Cell Signaling, 9806), supernatants were recovered by centrifugation, protein concentration was measured using a Pierce BCA Protein Assay Kit (Thermo Fisher Scientific, 23225), and equal amounts of total protein were separated by SDS-PAGE. Proteins were transferred to nitrocellulose membranes (Bio-Rad), followed by a blocking in 5% BSA in TBST. The membranes were incubated overnight at 4°C with primary antibody, p-MerTK (Novus Biologicals, NB300-690), MerTK (R&D Systems, AF591), Actin (MilliporeSigma, A2066), Axl (Santa Cruz, sc-1096), and PDGF-Rα (Cell Signaling, 3174), followed by corresponding horseradish peroxidase–conjugated secondary antibody (Jackson ImmunoResearch). Specific bands were detected by using WesternSure PREMIUM chemiluminescent substrate (Li-Cor) on a Li-Cor imaging system (Odyssey-Fc).

**Histology and tissue analysis.** Formalin-fixed tissues were embedded in paraffin and cut into 5-μm sections. Sections were evaluated by H&E and immunohistochemical analysis using antibodies specific for CD3 (Cell Signaling, 77699); Ki67 (Abcam, 15580), and cleaved caspase-3 (Cell Signaling, 9664). Negative controls included omission of primary antibody. Color images were obtained with a Nikon Eclipse E600 microscope using a Nikon Digital Dx1200me camera and ACT1 software (Universal Imaging Corporation). Pictures were analyzed using NIS Elements (Nikon).

**Nanostring analysis.** KLN205 tumor tissues from mice after 6 days of treatment were lysed in RLT lysis buffer and purified according to the manufacturer’s instructions (QIAGEN). RNA was sent to the Genomic and RNA Profiling Core in Baylor College of Medicine and analyzed using a preassembled nCounter Pan-Cancer Immune Profiling Panel (mouse) and the nCounter system (NanoString Technologies) according to the manufacturer’s instructions. Samples were then normalized based on the geometric means of the supplied positive controls and the panel of housekeeping genes, as recommended by the manufacturer. Only genes that were significantly different (P < 0.05; t test, false discover rate adjusted) and at least 1.5-fold differentially expressed between groups were considered.

**Flow cytometry analysis.** Tumors were digested with a cocktail containing collagenase I (45 μg/ml; Worthington), collagenase II (15 μg/ml; Worthington), collagenase III (45 μg/ml; Worthington), collagenase IV (45 μg/ml; Worthington), elastase (0.075 μg/ml; Worthington), hyaluronidase (30 μg/ml; MilliporeSigma), and DNase type I (25 μg/ml; MilliporeSigma) for 40 minutes at 37°C and passed through a 70-μm cell strainer (Falcon). Splenocytes were isolated from spleens and passed through a 70-μm cell strainer (Falcon). Suspensions were washed twice with PBS and stained with Fixable Viability Dye (Thermo Fisher) for 1 hour. The cell suspensions were then washed and stained with antibodies detecting CD11b (BD Bioscience, 557657), Ly-6C (BD Bioscience, 562728), Ly-6G (BD Bioscience, 740953), F4/80 (Biolegend, 123132), CD274 (PD-L1, BD Bioscience, 563369), CD11c (BD Bioscience, 564079), I-A/E (BD Bioscience, 562009), CD3 (BD Bioscience, 553061), CD4 (BD Bioscience, 562891), CD8 (BD Bioscience, 563332), CD279 (PD-1, BD Bioscience, 563059), CD152 (CTLA-4, BD Bioscience, 565778), and CD25 (IL-2 receptor α, BD Bioscience, 562694) for 1 hour at 4°C. Surface-stained cells were fixed, permeabilized, and stained for intracellular markers arginase 1 (R&D Systems, ICS868P), iNOS (Thermo Fisher, 17-5920-82), FoxP3 (BD Bioscience, 560401), and Ki67
Cells were analyzed using FACS LSRLFortessa SORP, and analysis was performed using FlowJo, with the help of the Moody Foundation flow cytometry facility at University of Texas Southwestern Medical Center.

**Pharmacokinetic analysis.** Athymic mice were dosed with sitravatinib at 10 or 20 mg/kg by oral gavage once per day for 4 or 14 consecutive days. Blood samples were collected before dosing and at 1, 4, 6, 10, 16, and 24 hours after dosing in Na citrate tubes and centrifuged and plasma was harvested and stored at −80 degrees. Plasma samples were analyzed for levels of sitravatinib using an Agilent 1200 HPLC system coupled with an MDS Sciex API3200 triple quadrupole mass spectrometer at MethylGene. The processed sample (5–10 μl) was injected onto a Thermoacquisil C18 (4 mm × 50 mm, 3 μm). An isocratic elution with the mobile phase A/B (0.5% formic acid, 10 mM ammonium formate in water, 0.5% formic acid, 10 mM ammonium in 10:90 MeOH/acetonitrile) at 40:60 was used. The flow rate was set at 1 ml/min. Sitravatinib was eluted at 1.26 minutes. The total run time was 5 minutes. Positive electrospray ionization mode was used. Sitravatinib was monitored at the MRM transition of m/z 629.9 → m/z 554.9. The HPLC-MS/MS system control and quantitation were carried out using the Analyste(R) 1.4 software.

**Statistics.** Data are reported as mean ± SD. Statistical analysis was performed with a 2-tailed t test or ANOVA using GraphPad Prism software (version 4.00 for Windows). For all analyses, P < 0.05 was considered statistically significant.

**Study approval.** Animal experiments in this study were approved by and performed in accordance with the Institutional Animal Care and Use Committee at the University of Texas Southwestern Medical Center at Dallas.