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Loss of Integrin-linked kinase sensitizes breast cancer to SRC inhibitors

Running title: Loss of ILK sensitizes breast cancer to SRC inhibitors

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Keywords

Breast cancer, synthetic lethality, bosutinib, Integrin-linked kinase, SRC.

Declaration of Conflicting Interests

The authors declared no potential conflicts of interest with respect to the research, authorship, and/or publication of this article.

Abstract

SRC is a non-receptor tyrosine kinase with key roles in breast cancer development and progression. Despite this, SRC tyrosine kinase inhibitors have so far failed to live up to their promise in clinical trials, with poor overall response rates. We aimed to identify possible synergistic gene-drug interactions to discover new rational combination therapies for SRC inhibitors. An unbiased genomewide CRISPR-Cas9 knockout screen in a model of triple-negative breast cancer revealed that loss of Integrin-linked kinase (ILK) and its binding partners α-Parvin and PINCH-1 sensitizes cells to bosutinib, a clinically approved SRC/ABL kinase inhibitor. Sensitivity to bosutinib did not correlate with ABL dependency; instead, bosutinib likely induces these effects by acting as a SRC tyrosine kinase inhibitor. Furthermore, in vitro and in vivo models showed that loss of ILK enhanced sensitivity to eCF506, a novel and highly selective inhibitor of SRC with a unique mode of action. Whole-genome RNA sequencing following bosutinib treatment in ILK knockout cells identified broad changes in the expression of genes regulating cell adhesion and cell-extracellular matrix. Increased sensitivity to SRC inhibition in ILK knockout cells was associated with defective adhesion, resulting in reduced cell number as well as increased G1 arrest and apoptosis. These findings support the potential of ILK loss as an exploitable therapeutic vulnerability in breast cancer, enhancing the effectiveness of clinical SRC inhibitors.

Statement of Significance

A CRISPR-Cas9 screen reveals that loss of integrin-linked kinase synergizes with SRC inhibition, providing a new opportunity for enhancing the clinical effectiveness of SRC inhibitors in breast cancer.

Introduction

SRC family kinases are group of nine non-receptor tyrosine kinases of which SRC is the prototype. SRC is a key regulator of cell-matrix and cell-cell adhesions and is known to play important roles in a number of cancer-associated phenotypes including proliferation, migration and invasion. Diseaseassociated upregulation of SRC kinase activity has been demonstrated in many malignancies, including breast, colon and gastric cancer. In breast cancer, elevated SRC expression is associated with poor prognosis (1). Despite the validated aetiological role of this target and several commercial drug discovery programs, there has been limited evidence of clinical efficacy in solid tumours, and no progress in understanding the molecular signatures that predict sensitivity to available inhibitors (2,3). Bosutinib (SKI-606) is a multi-kinase inhibitor and has activity against all nine SRC family members as well as ABL (4). Bosutinib is currently used as a second-line treatment for patients with chronic myeloid leukaemia that carry the BCR-ABL oncogene (5), and has shown activity in preclinical models of breast cancer (6,7). In a phase II clinical trial of bosutinib monotherapy in heavily pre-treated, advanced breast cancer patients, bosutinib presented poor overall response rates of 5.5% (8). However, there was evidence of disease stabilisation (37%), and together with the reported disease control rates following treatment with dasatinib, a multi-kinase inhibitor with activity against SRC family members (9), this suggests there is a subset of breast cancer patients that are responsive to dual SRC/ABL inhibition.

One of the reasons that SRC/ABL tyrosine kinase inhibitors (TKIs) such as bosutinib have so far failed to live up to their promise in clinical trials is due to limited information on the molecular determinants of SRC inhibitor sensitivity. To address this, we undertook an unbiased genome-wide CRISPR-Cas9 screen in a model of triple-negative breast cancer (TNBC) to identify genes whose depletion alters the cellular sensitivity to bosutinib treatment. We show that loss of the cell-matrix adhesion protein integrin-linked kinase (ILK) and its binding partners α -Parvin and PINCH-1 can potentiate the inhibitory effect of bosutinib.

Materials and Methods

Additional information on all Methods can be found in the Supplementary Information.

Cell Culture

All cell lines were grown in a humidified cell culture incubator at 37°C and 5% CO₂. MDA-MB-231, MDA-MB-468, MDA-MB-157, MDA-MB-134-VI, SKBR3 and MCF-7 were maintained in DMEM +10% FBS. ZR-75-1 and T47D were maintained in RPMI-1640 +10% FBS. All cell lines were passaged for no more than 20 passages. Automated cell counts for passage calculations were obtained from the CellCountess II automated cell counter (Thermo Fisher Scientific). Cells were routinely tested for mycoplasma and were authenticated by STR profiling.

2D cell viability assays

Unless stated otherwise, cells were seeded at 1,000 cells/well into flat-bottom 96-well plates (Greiner 655180). Edge wells were excluded and plates were left for 30 min RT before transferring to an incubator. Plates for real-time analysis were transferred to the IncuCyte FLR imaging system (Essen BioScience, USA) as previously described⁽¹⁰⁾. Unless stated otherwise, compounds were added (5 μ L/well) at two days post seeding, for five days. Compounds were reconstituted in dimethyl sulfoxide (DMSO). Plates were assayed for cell viability using the resazurin reduction assay and/or cell counts as described previously (10). The Spark 20M (TECAN) was used to quantify resazurin fluorescence following best practice procedures described previously (11). The ImageXpress system (Molecular Devices) was used to image Hoechst 33342 stained plates. CellProfiler (www.cellprofiler.org) was used to enumerate cell counts as previously described (12). Normalised cell viability was calculated as follows: sample / DMSO × 100. Relative half-inhibitory

concentrations (EC_{50}) were calculated in Prism from the nonlinear regression algorithm of the normalised cell viability plotted against an 8-point compound dilution.

3D spheroid cell viability assays

Cells were seeded at 2,000 cells/well into round-bottom 96-well plates (Corning #7007). Once spheroids had formed over two to three days, plates were imaged at $4\times$ magnification using the ImageXpress and subsequently compounds were added. On day seven post drugging, spheroids were stained with Calcein-AM (3.75 µM final) to give a 'live cell' readout, propidium iodide (1.0 µM final) or Draq7 (3.0 µM final) for a 'dead cell' readout and Hoechst 33342 (1.0 µM final) for a 'total cells' readout. The ImageXpress was used to create a single flattened maximum intensity projection from an eight-step Z stack with a 50 µM step size. A custom CellProfiler pipeline was used to quantify the spheroid area and intensity values for each spheroid. Data was normalised as follows: sample / DMSO × 100.

Cell adhesion challenge

Cells were seeded at 20,000 cells/well in serum-free media into 96-well plates (Greiner #655180). DMSO or bosutinib was added at the time of seeding. Plates were imaged every 10 min in the IncuCyte FLR for 2 h and then washed gently two times with 100 μ L PBS taking care to not disrupt adhered cells. Plates were then returned to the IncuCyte for a final post-wash scan to determine the adhered cells.

Western blotting

Cell lysates were prepared in RIPA buffer (150 mM NaCl, 50 mM Tris pH 7.2, 0.1% SDS, 1.0% Triton X-100) with protease and phosphatase inhibitors (10 μ g/mL aprotinin, 125 mmol/L phenylmethylsulfonyl fluoride, 100 μ mol/L sodium orthovanadate, and 0.5 mmol/L sodium fluoride). For 3D spheroid protein extraction, spheroids were pooled and then solubilised in RIPA using sonication. Protein concentration was determined using the BCA protein assay kit. Whole-cell

lysates were fractionated by SDS-PAGE (20 µg per sample) and transferred onto nitrocellulose membranes. Primary antibodies used were: ABL (SCBT, #sc-23 and #sc-56887) and pY245 ABL (CST #2861); Arg (Ab134134); Cofillin (CST #5175); CrkL (CST #3182) and pY207 CrkL (CST #3181); FAK (CST #3285) and pY397 FAK (CST #3283); GAPDH (CST #5174); ILK (BD #611803 and Abcam #ab52480); mouse IgG1 (CST #5415); rabbit IgG (CST #2729); α-Parvin (CST #8190 and CST #4026); PINCH-1 (CST #11890); SRC (CST #2109) and pY416 SRC (CST #2101). Protein-antibody complexes were visualised by enhanced chemiluminescence detection (GE Healthcare) with horseradish peroxidase-conjugated secondary antibodies (CST #7076 and #7074) using the ChemiDoc XRS imaging system (Bio-Rad Laboratories). Experiments were repeated at least three times. Blots were cropped for clarity. The volume tool in Image Lab (v5.0) was used to quantify intensity. The same standard rectangle was used for each sample, as well as background correction for each lane.

Genome-wide CRISPR-Cas9 dropout screen in MDA-MB-231 cells

The CRSIPR screen was performed using the pooled lentiviral genome-wide CRISPR-SpCas9 TKOv3 library targeting 18,053 protein-coding genes (4 gRNAs/gene), a gift from Jason Moffat (Addgene #90294), that was amplified and packaged into lentiviral particles (Vector Core, University of Michigan) as described (13). A total of 9×10^7 MDA-MB-231 cells were transduced with the TKOv3 lentiviral library (71,090 gRNAs) at a low MOI (0.36) to ensure each cell receives only one sgRNA and to achieve >300-fold library coverage after selection. To select for transductants, 24 h after infection, cells were grown in the presence of 2.5 µg/mL puromycin for 48 h, which was considered the initial time point (T0). Cells were then passaged every 3-4 days and library coverage of \geq 300 cells per gRNA was maintained at every step. Seven days after puromycin selection (T7), two technical replicates containing 24×10^6 cells were established for DMSO and bosutinib (provided by Pfizer), respectively. An EC₂₀ concentration of bosutinib (0.9 µM) was added 24 h post seeding for 3 days and cells were treated with bosutinib for five rounds or for ~12

population doubling times over 20 days. In order to capture the full representation of the screen, $25 \times$ 10⁶ cells were collected at each passage point for genomic DNA extraction using the QIA amp Blood Maxi Kit (#51192). The representation of genome-integrated gRNA sequences was determined using a two-step PCR in which the first step 'PCR1' amplifies the lentiviral sequence containing the 20 bp gRNA cassette from genomic DNA and the second step 'PCR2' attaches unique barcodes for sample deconvolution and Illumina TruSeq adapters (i5 and i7) for next-generation sequencing (Supplementary Figure 1, Supplementary Tables 1-3). The final pooled library was quantified with the Qubit 2.0 (Invitrogen) and Agilent 2100 Bioanalyser platforms. The diluted and denatured NGS library was sequenced on a NextSeq® 500/550 High Output Kit v2 (75 cycles) with a custom 'dark cycle recipe', for dual-index, single-read reads. Sequencing was performed by the Edinburgh Welcome Trust Clinical Research Facility. DrugZ (14) was used to calculate a gene-level normalised Z-score for determining significant differences between the DMSO and bosutinib samples at both T19 (three rounds of drugging) and T27 (five rounds of drugging). Gene copy number data was incorporated into the analysis as double-stranded DNA breaks from CRISPR-Cas9 can generate false-positive hits in genes with high copy numbers due to cutting-related genotoxicity (15). There was no correlation between gene copy number and gene-level normalised Z-score (Supplementary Table 4). Supplementary Dataset 1 contains the DrugZ results file.

Interaction network analysis

Composite functional association networks were constructed for putative enhancer and suppressor genes using GeneMANIA (version 3.5.2; human interactions: https://genemania.org/) in Cytoscape (version 3.8.0: https://cytoscape.org/). Network edges were weighted according to evidence of co-functionality using GeneMANIA. Connected networks were clustered using the force-directed algorithm in the Prefuse toolkit (https://github.com/prefuse/Prefuse).

Patient survival analysis

cBioPortal was used to download the METABRIC data (16) and the cancer genome atlas (TCGA-PanCancer: http://cancergenome.nih.gov/) data. Data was analysed using the R packages *survival* (17), *survival* (18) and their dependencies. In the METABRIC dataset, the TNBC subset was created from the clinical data for estrogen receptor alpha (ER), progesterone receptor (PR) and human epidermal growth factor receptor 2 (HER2). The TNBC subset had a 75% overlap with the PAM50-basal METABRIC data. The TNBC subtype increased the total number of patients compared to the PAM50-basal subtype, thereby increasing the power of the statistical test for high vs low *ILK*. The optimal threshold for dichotomisation of the datasets into high and low *ILK* expression was identified using the *survivALL* R package and 'plotALL' function for multi-cut point analysis from the METABRIC dataset. The same dichotomisation threshold was then applied to the TCGA dataset in order to validate the cut point.

CRISPR knockout of ILK and ABL1 in mammalian cells

CRISPR-(e)SpCas9 target sites designed against *ILK* and *ABL1* were created using the Broad Institute's gRNA design tool (https://portals.broadinstitute.org/gpp/public/analysis-tools/sgrnadesign), which uses the optimal gRNA and off-target design principles of Doench *et al.* (19). The gRNAs target exons 1 and 8 of *ILK* (deposited with Addgene plasmid #163320 and 163321), within the N-terminal ankyrin repeat domain-1 and the C-terminal protein kinase domain, respectively (Supplementary Table 5). For *ABL1*, the two designed gRNAs target exon 4 (Deposited with Addgene plasmid #163322 and 163323) within the protein kinase domain of both *ABL1* isoforms 1a and 1b (Supplementary Table 6). The CRISPR plasmid constructs PX459v2 and eSpCas9(1.1) were purchased from Addgene (plasmid #62988 and #71814) and were deposited by Prof. Feng Zhang's laboratory (20). The CRISPR-(e)Cas9 constructs were created as described by Ran *et al.* (21). Individual transfected cells (Nucleofector system (Lonza)) were selected for clonal expansion using puromycin selection (PX459v2) and flow cytometry. Monoclonal colonies were screened for loss of the gene of interest using western blotting with the ILK monoclonal antibodies BD #611801 and Abcam #ab52480 or the ABL monoclonal antibodies 8E9 (sc-56887) and 24-11 (sc-23). The plasmids pDONR223-ABL1 (Plasmid #23939) and pLX301 (Plasmid #25895) were purchased from Addgene, and were deposited by William Hahn & David Root (22) and David Root (23), respectively. Using gateway cloning, pDONR223-*ABL1* (containing the *ABL1* isoform 1a (*ABL1-1A*)) was cut and inserted into pLX301. 293T cells were used for viral packaging.

Cell cycle analysis

Cells were seeded in 6-well plates and drugged at either 48 h post seeding (48 h drug time-point) or 72 h (24 h drug time-point). Cells were harvested using trypsin and fixed in cold 70% ethanol for 48 h. Pellets were treated with RNAse H (NEB, #M0297L) and stained with 20 μ g/ml propidium iodide. Cells were sorted on the BD LSRFortessa and analysed using BD FACSDiva software (v8.0.1).

Apoptosis and cell death assay

Cells were seeded in 6-well plates and drugged for 4 days. Medium was removed and spun down to include floating cells. Cells were then trypsinised and resuspended with 5% FBS in PBS. Cells were sorted on the BD LSRFortessa and analysed using BD FACSDiva software (v8.0.1). CellEvent Caspase 3/7 (apoptotic cells) and DAPI (dead cells) dyes were used.

Breast cancer cell line datasets

We generated a 16 BC cell line dataset for sensitivity to bosutinib using a PrestoBlue (Resazurin reduction) cell viability assay as described above (24). The cb5 cell line dataset is an integrated compendium of 5 Affymetrix datasets and was downloaded from Moleirinho *et al.* 2013 (25). Multiple cell line microarray datasets were combined for *ILK* expression levels in the 16 breast cancer cell lines. The Cancerxgene dataset was downloaded from: https://www.cancerrxgene.org/. The Cancerxgene dataset included 43 BC cell lines with data for bosutinib treatment. The Depmap interactive website is available at: https://depmap.org/portal/. The Depmap interactive portal of 20

BC cell lines used bosutinib cell viability data compared to *ILK* RNAseq expression levels and ILK relative protein expression.

RNA sequencing

Bosutinib EC₂₀ (0.9 µM) or DMSO were added to cells at 24 h post seeding in 10 cm dishes. Three independent experimental replicates were performed. Total RNA was extracted at 72 h post drugging using RNAeasy Miniprep Kit (Qiagen) according to the manufacturer's protocol. RNA quality and integrity were assessed on the Fragment Analyser Automated Capillary Electrophoresis System (Agilent Technologies). RNA yield and DNA contamination was quantified using the Qubit 2.0. Library preparation and sequencing was performed by the Edinburgh Welcome Trust Clinical Research Facility. The Lexogen QuantSeq 3'mRNA-Seq Library Prep Kit (FWD) for Illumina was used to generate libraries from 200 ng total RNA. Library quantity and quality were assessed using Qubit 2.0 and the Bioanalyser. Single-read sequencing was performed using the NextSeq 500/550 High-Output v2.5 (75 cycles) Kit on the NextSeq 550 platform (Illumina). Demultiplexed FASTQ files with md5 checksums were imported into the Lexogen Bluebee platform and the Quantseq 2.3.6 FWD pipeline was used. Trimming was performed using bbduk v35.92. STAR v2.5.2a (26) was used to align the read data to human genome build human GRCh38. The expected counts from HTSeqcount v0.6.0 were imported into the statistical software package R (www.r-project.org). Differential expression analysis was undertaken with the EdgeR (v3.26.8) and Limma (v3.40.6) packages. Significantly differentially expressed genes were determined using the FDR (Benjamini & Hochberg) p-value adjustment <0.05 and a log2 fold change of ± 1 . Gene ontology (GO) functional enrichment analysis was carried out using ToppGene (https://toppgene.cchmc.org/). Read count and differential expression analysis data is available in Supplementary Datasets 2-5.

Tumourigenicity assay

Immunodeficient Rag2-Il2rg double knockout mice (R2G2) were obtained from Envigo, UK. Female mice (2-4 months of age) were bilaterally injected subcutaneously with 2.5×10^6 MDA-MB-231

PX45v2 or ILK gRNA 2 cells (100 µL in PBS). Once tumours developed, mice were randomized (using the standard RAND function in Microsoft Excel) to daily oral gavage with either vehicle (50 mM lactic acid with 0.1M NaOH, pH 6), eCF506 (40 mg/kg body weight, 0.1 mL/20 g) or bosutinib (75 mg/kg body weight, 0.1 mL/20 g). Statisticians were blinded to treatment group allocation. Mice were monitored twice weekly and culled if they became sickly or tumours had reached a maximum size of 15 mm. Tumours were collected at the end of the study. Formalin-fixed tissues were processed into paraffin wax blocks. Sections were cut, dewaxed and rehydrated before heat mediated antigen retrieval in sodium citrate buffer. After blocking, sections were incubated with Ki67 antibody at 1:400 (Cell Signalling Technology #9027) followed by 3,3'-diaminobenzidine (DAB) Chromagen (Agilent #K3468) for visualisation. Sections were counterstained with hematoxylin and quPath used to quantify Ki67 positive nuclei (https://qupath.github.io/). All animal studies and procedures were approved by the University of Edinburgh Ethical review committee (Application #116-WGH-20) and conducted in accordance with United Kingdom Home Office regulations.

Results

Bosutinib demonstrates antiproliferative activity in several breast cancer cell lines

Due to limited published data on sensitivity to bosutinib in breast cancer cell lines (6), we tested a panel of eight cell lines in both 2D and 3D viability assays. Bosutinib sensitivity did not correlate with breast cancer subtype in our cell line panel (Figure 1A and Supplementary Table 7), however the triple-negative MDA-MB-231 and the luminal A MCF-7 cell lines were the most sensitive to bosutinib.

Both cell lines were more sensitive to bosutinib in a 3D spheroid growth assay (Figure 1A). However, although the half maximal effective concentration (EC₅₀) values for bosutinib were in the sub-micromolar range for both MDA-MB-231 and MCF-7 cells, no significant cell death was seen at these concentrations using a 3D spheroid live/dead assay (Figure 1B and Supplementary Figure 2A). In both cell lines, bosutinib treatment caused a dose-dependent decrease in spheroid area (EC₅₀=0.151 μ M and 0.194 μ M, respectively), but no corresponding decrease in Calcein-AM (viable cells) or increase in propidium iodide intensity (dead cells) until concentrations greater than 3 μ M.

We confirmed that bosutinib was inhibiting its designed targets SRC and ABL in the spheroid assay by western blot, where bosutinib treatment resulted in a dose-dependent reduction in pY419-SRC, with almost complete inhibition (~90% reduction) at 0.1 μ M (Figure 1C, and Supplementary Figure 2B). Total SRC was increased, as has been reported following treatment with SRC inhibitors (27). There was a marked reduction in ABL phosphorylation at Y245, a site required for its kinase activity, following treatment with bosutinib at 30 and 100 nM for the MDA-MB-231 and MCF-7 cells, respectively. This correlated with a 90% and 70% reduction in pY207-CRKL at 30 nM in the MDA-MB-231 and MCF-7 cell lines, respectively (Figure 1C and Supplementary Figure 2C). To provide mechanistic insight into the molecular pathways underlying bosutinib sensitivity, we performed a reverse-phase protein array (RPPA) analysis using 125 antibodies covering a panel of signalling proteins and phosphoproteins in MDA-MB-231 and MCF-7 cells following treatment with bosutinib (Supplementary Figure 3). This identified 17 and 7 significantly differentially regulated proteins and phosphoproteins in MDA-MB-231 and MCF-7 cells, respectively (p-value <0.05, log2 fold change ± 0.75). As expected, bosutinib treatment inhibited its designed targets SRC and ABL, with a significantly decreased signal intensity in SRC autophosphorylation at tyrosine (pY)419 (70% reduction at 1.25 μ M), as well as an 80% and 70% reduction in pY207-CRKL, a key substrate of ABL, at 1.25 μ M for the MDA-MB-231 and MCF-7 cells, respectively (Supplementary Figure 3). Other consistent changes that were seen between the cell lines included a decrease in pY15 CDK1 and total CDK1, and a decrease in phosphorylation of phospho-threonine (pThr)183-JNK (Supplementary Figure 3A).

Loss of ABL does not alter bosutinib effects in MDA-MB-231 and MCF-7 cells

As bosutinib is a dual SRC/ABL inhibitor, which effectively inhibited ABL activity in both the MDA-MB-231 and MCF-7 cell lines, we asked whether the antiproliferative activity of bosutinib was due to ABL inhibition. We created *ABL1* knockout isogenic cell lines using CRISPR-eCas9 (Supplementary Figure 4A). Using 2D cell viability assays there were no differences between bosutinib EC₅₀ values for the *ABL1* knockout MCF-7 or MDA-MB-231 cell lines compared to the parental cells or *ABL1* knockout cell lines in which *ABL1* was re-expressed (Figure 1D). Similar results were observed with three other MDA-MB-231 *ABL1* gRNA cell lines (Supplementary Figure 4B). Loss of ABL1 also did not alter the sensitivity to bosutinib in the 3D spheroid assay (Supplementary Figure 4C). Thus, loss of ABL1 does not alter bosutinib activity in MDA-MB-231 and MCF-7 cells in both 2D and 3D assays, indicating that bosutinib is unlikely to act via ABL1 inhibition.

A genome-wide CRISPR-Cas9 knockout screen identifies the ILK-Parvin-PINCH complex as a key determinant of bosutinib sensitivity

To identify genes that can alter bosutinib sensitivity we performed an unbiased genome-wide CRISPR-Cas9 negative selection screen in the MDA-MB-231 cell line (Figure 2A). We chose the Toronto knockout version 3 (TKOv3) library (13) that contains sequence-optimised guide RNA (gRNA) following empirically determined on- and off-target prediction enabling increased accuracy and scalability of the CRISPR screen. Cells were transduced with the lentiviral TKOv3 library at a low multiplicity of infection (MOI=0.36) and selected with puromycin. Since almost complete modification of genomic loci occurs after seven days of CRISPR treatment (28), cells were passaged for seven days after puromycin selection (T7), after which either an EC_{20} concentration of bosutinib (0.9 µM) or DMSO as a control were added 24 h post seeding for 3 days. A 24 h time point for bosutinib addition was chosen in order for cell attachment to occur, but before log-phase growth began. Cells were treated in this way for around 12 population doubling times (PDs) over 20 days. The screen was performed in technical duplicates and genomic DNA was extracted and genomeintegrated gRNA sequences were amplified and labelled with barcodes via a 2-step polymerase chain reaction (PCR). Next-generation sequencing was used to quantify the library representation of the gRNA sequences and DrugZ (14) was used to calculate gene-level normalised Z-scores for determining differences between the DMSO and bosutinib samples at the T19 (~7 PDs) and T27 (~12 PDs) time points (Figure 2A). This allowed the identification of enhancer genes whose loss of function enhance sensitivity to bosutinib and suppressor gene interactions whose loss of function reduce bosutinib efficacy.

Hierarchical cluster analysis of the individual gRNAs from the top enhancer and suppressor genes revealed groupings of replicates and treatments across time points, emphasising consistency in the results between the T19 and T27 time points at the gRNA level (Supplementary Figure 5A). DrugZ analysis of the T27 time point identified 19 enhancer genes and 23 suppressor genes with a significant gene-level normalised Z-score (FDR<0.05) with 12 enhancer genes and 15 suppressor genes also identified at the T19 time point (Figure 2B and Supplementary Figure 5B). Integrin-linked kinase (*ILK*) and α -Parvin (*PARVA*) were the highest ranked enhancer genes at both the T27 and T19 time points. PINCH-1 (*LIMS1*) also had a significantly decreased gene-level normalised Z-score upon bosutinib treatment at both time points (Supplementary Table 4). These three proteins form the ILK-Parvin-PINCH (IPP) complex. The IPP complex tethers cells to their surrounding environment via integrins and is involved in cytoskeleton remodelling, angiogenesis, proliferation, survival and differentiation (29). Hence, within the pool of CRISPR knockout cells, those cells with loss of ILK or its binding partners α -Parvin and PINCH-1 significantly enhanced the inhibitory effect of bosutinib on MDA-MB-231 cell growth.

Functional network analysis revealed connectivity between many of the top enhancer genes (Figure 2C and Supplementary Figure 6A). Interestingly, both the expected targets of bosutinib, SRC and ABL, and many of the enhancer genes, such as those involved in the IPP complex, *AMBRA1, EED* and *TYMS*, converge on the focal adhesion kinase (FAK) network (Supplementary Figure 6B) and are linked to cytoskeletal organisation. Consistent with this, gene ontology (GO) analysis of the top enhancer genes centred around the interactions of the IPP complex. The key enriched molecular pathways were adherens junctions, cell-extracellular matrix (ECM) interactions, focal adhesions and the regulation of cytoskeletal remodelling and cell spreading (Figure 2D). There were also a number of interesting suppressor genes such as *CBFB*, *RUNX1*, *EP300* and *AHR*, but no GO terms were significantly enriched for the top suppressor genes (Supplementary Figure 6C and Supplementary Table 8).

Loss of the ILK-Parvin-PINCH complex enhances sensitivity to bosutinib

We focused on *ILK* as the top ranked enhancer gene for validation studies. Analysis of the METABRIC and TCGA-PanCancer breast cancer datasets showed that high *ILK* expression was significantly associated with poor breast cancer-specific survival for the TNBC and basal subtypes (Kaplan Meier log-rank test, p<0.05), but not for the remaining breast cancer patients (Figure 2E-F and Supplementary Figure 7), suggesting a functional role for ILK in these subtypes. To validate that loss of ILK, α -Parvin or PINCH-1 can potentiate the inhibitory effect of bosutinib, we generated clonal *ILK* knockout cell lines. We created MDA-MB-231 *ILK* knockout cell lines using two gRNAs targeting *ILK* and a transiently expressed CRISPR-Cas9 system (PX459v2) to generate single-cell-derived cell lines lacking ILK. We generated three cell lines for ILK gRNA 1 and three for ILK gRNA 2 (Supplementary Figure 8A). The IPP complex members are interdependent for their protein stability (30). Accordingly, our western blot analysis showed that *ILK* knockout in MDA-MB-231 cells resulted in the downregulation of PINCH-1 and α -Parvin in addition to loss of ILK (Figure 3A and Supplementary Figure 8A). Therefore, we did not endeavour to create PINCH-1 or α -Parvin CRISPR knockouts in addition to the *ILK* knockout cell lines.

We next assessed the impact of bosutinib treatment on cell viability in cells with or without ILK. Loss of ILK resulted in a ~4-fold decrease (p-value <0.0001) in the EC₅₀ of bosutinib compared to the empty-gRNA CRISPR control (PX459v2) (Figure 3B). ILK loss alone did not impact the growth rate of the cells, as assessed by the change in cell confluence over time (Figure 3C). At the EC₂₀ concentration of 0.9 μ M, bosutinib significantly decreased the real-time growth of all MDA-MB-231 *ILK* knockout cell lines compared to the PX459v2 control (Figure 3C and Supplementary Figure 8B). This resulted in a ~3-fold change decrease in cell confluence between cells with ILK loss and the PX459v2 control at the end of the assay (Figure 3D). Bosutinib treatment also resulted in a significant decrease in cell number in all six MDA-MB-231 *ILK* knockout cell lines compared to the PX459v2 control (Figure 3E, Supplementary Figure 8C). A similar enhancement of bosutinib activity was seen in the T47D breast cancer cell line following knockdown of *ILK* (Supplementary Figure 8D, E). Cell cycle analysis revealed no changes in cell cycle distribution upon ILK loss or bosutinib treatment (EC_{20} , 0.9 μ M). However, there was an increased G1 phase arrest (p-value <0.05) when *ILK* knockout cells were treated with bosutinib (Supplementary Figure 8F). Notably, there was also a 4-fold increase (p-value <0.0001) in the percent of apoptotic cells when *ILK* knockout cells were treated with bosutinib compared to PX459v2 control cells (Figure 3F). Taken together, these data confirm that loss of ILK in MDA-MB-231 cells enhances sensitivity to bosutinib through an enhanced G1 arrest and increased apoptosis.

SRC but not ABL is responsible for the increased activity of bosutinib in combination with ILK loss

The other designed target of bosutinib is SRC. However, as there are nine SRC family members it is difficult to create suitable knockouts. In this respect, the SRC-specific inhibitor eCF506 is an effective tool compound as it inhibits all nine SRC family members and has been shown to have a 1,000-fold selectivity for SRC over ABL and potent growth inhibitory effects in both MDA-MB-231 and MCF-7 cell lines (24,27), supporting the importance of SRC activity in driving their proliferation. 2D cell viability assays confirmed that there were no differences between eCF506 EC₅₀ values for the *ABL1* knockout MDA-MB-231 cell lines compared to the parental cells (Supplementary Figure 9A). However, in the *ILK* knockout cells there was a ~4-fold decrease (p-value <0.0001) in the EC₅₀ for eCF506 compared to the PX459v2 CRISPR control cells (Figure 4A). Similar to bosutinib, analysis of apoptotic cells revealed a 6-fold increase (p-value <0.0001) in Caspase 3/7 positive cells upon eCF506 inhibition (40 nM; EC₂₀) in ILK depleted cells compared to the PX459v2 control cells (Figure 4B). In addition, cell cycle analysis showed a small but consistent increase in the G1 peak for eCF506 in ILK depleted cells (Supplementary Figure 8F). Treatment of the MDA-MB-231 *ILK* knockout isogenic cell lines with the ABL allosteric inhibitor GNF-2 (31) did

not change cell viabilities (Supplementary Figure 9B). As expected, addition of eCF506 to the PX459v2 MDA-MB-231 cells resulted in a significant reduction in cell growth as measured in realtime in the IncuCyte and end-point cell counts (Figure 4C). Similar to bosutinib treatment, the combined effect of ILK loss and eCF506 treatment (60 nM; EC₃₀) resulted in a further significant decrease in cell growth as well as cell counts compared to the PX459v2 control (Figure 4D-E). Thus, loss of ILK in the MDA-MB-231 cell lines enhanced the effects of SRC inhibition in the absence of ABL inhibition. These findings taken together confirm that SRC inhibition is required for the decreased viability in cells with ILK loss following treatment with bosutinib or eCF506.

Low ILK expression moderately correlates with sensitivity to bosutinib

To establish whether *ILK* expression was associated with bosutinib sensitivity in a larger panel of breast cancer cell lines, Pearson's rank correlation coefficients were used as a measure of the strength of association between sensitivity to bosutinib and *ILK* mRNA expression levels in three different cell line datasets, two publicly available and one in-house (Supplementary Table 9, Supplementary Figure 10). We generated a dataset from 16 breast cancer cell lines using EC₅₀ cell viability measurements to determine sensitivity to bosutinib (24). For *ILK* mRNA expression, we used an integrated compendium of 5 Affymetrix cell line datasets (cb5) (25). Decreased *ILK* expression correlated with a lower bosutinib EC₅₀, although with only a moderate effect size (Pearson's: r(61) = 0.358, p = 0.004). The publicly available Cancerxgene dataset allowed us to include 43 breast cancer cell lines with data for bosutinib treatment queried against the cb5 dataset. This also resulted in a weak but positively correlated relationship (Pearson's: r(119) = 0.283, p = 0.002). Similar correlations were seen between bosutinib sensitivity and *LIMS1* and *PARVA* in these two datasets (Supplementary Table 9). Additionally, we utilised the Depmap interactive portal of 20 breast cancer cell lines with bosutinib cell viability data compared to ILK RNAseq and protein expression levels. Although this dataset had a positive correlation for both RNAseq and protein

expression it was not significant (Pearson's: r(18) = 0.395, p = 0.085 and r(18) = 0.299, p = 0.199, respectively), while the correlation with *LIMS1* and *PARVA* was significant (Supplementary Table 9). The small number of cell lines in these datasets which have low *ILK* expression limits the predictivity of these analyses, but taken together suggest that *ILK* gene expression or protein alone is not sufficient to predict sensitivity to bosutinib.

In order to look at gene expression of *ILK* in human breast tumours, we utilised an integrated compendium of 17 Affymetrix datasets (cb17) representing 2,996 breast cancers (25). This dataset has detection calls calculated from perfect and mis-matched probes, which effectively indicate whether genes are detectably 'present' or 'absent' in each sample. The basal subtype had significantly lower *ILK* expression (Tukey one-way ANOVA, p<0.001) compared to all other subtypes (Supplementary Figure 11A). Interestingly, *ILK* was significantly called 'absent' in 15% of basal breast tumours (Chi-square test, p<0.001) compared to around 6% for other subtypes (Supplementary Figure 11B, C). Although METABRIC and TCGA-PanCancer samples do not have 'absent' or 'present' detection calls, *ILK* expression was significantly lower in the basal subtype (Tukey HSD, p< 0.05) compared to all other subtypes except HER2 (Supplementary Figure 11D). A similar pattern was seen for PARVA but not *LIMS1*, other components of the IPP complex. Interestingly, in the same datasets *SRC* was higher in the basal subtype (Supplementary Figure 11D). It remains to be established whether there is a small subset of patients that would benefit from bosutinib treatment due to the absence of *ILK*, and the sensitisation of the luminal A T47D cell line to SRC inhibitor treatment following loss of *ILK*, indicates that this will not be subtype specific.

ILK loss in combination with bosutinib regulates extracellular matrix and adhesion protein transcripts

We undertook an unbiased genome-wide RNA sequencing approach to understand and exploit the mechanism(s) at the transciptional level that enabled ILK loss to potentiate SRC inhibition in our

TNBC model. Principal component analysis identified good clustering of similar samples and separation between different groups. Hierarchical clustering of log2 counts per million for the 500 most variable genes across all 12 samples also correctly clustered samples (Supplementary Figure 12A). The largest variation (dimension 1) was seen between the presence or absence of ILK which resulted in 444 DEGs (Figure 5A, B). This identified genes associated with the ECM, actin cytoskeleton, membrane structures, integrin and growth factor binding (Supplementary Figure 12B) consistent with ILKs known role in cell-matrix adhesions. For the MDA-MB-231 PX459v2 CRISPR control, bosutinib treatment resulted in 392 DEGs (DEGs; log2 fold change ± 1 and p-value < 0.05) (Figure 5B). Importantly, there were 504 DEGs unique to the combination of ILK loss and bosutinib addition (Figure 5B). To determine the magnitude of change in the 504 uniquely DEGs we compared the normalised read counts for bosutinib treatment with or without ILK. This resulted in 134 genes with significant (log2 fold change ±1 and p-value <0.05) changes upon bosutinib addition and ILK loss (Figure 5C). These 134 DEGs are shown across all samples, confirming that the changes are unique to bosutinib treatment of ILK knockout cells (Figure 5D). These uniquely DEGs were involved in ECM and cell adhesion processes as shown using ToppGene (Figure 5E). Network analysis of the 134 unique DEGs revealed a number of interactions with the most prominent being a collagen-integrin network of associated genes (Figure 5F), but also adhesion complex associated, and ECM-related gene expression interactions (Supplementary Figure 12C). Clearly, cell adhesion and cell-ECM interactions were the core effected processes in the context of ILK knockout-mediated bosutinib sensitivity, suggesting they may be responsible for the increased sensitivity to bosutinib observed in the ILK depleted cells.

ILK knockout in combination with bosutinib causes cell adhesion defects

The IPP complex members and SRC are involved in regulating cell matrix adhesion. Taken together with the RNAseq data, this suggested that defective adhesion may be involved in the increased

sensitivity to bosutinib eCF506 treatment in ILK knockout cells. To characterise the adhesive defects the ILK knockout cells, we first analysed their focal adhesions by carrying out in immunofluorescence using a number of antibodies against adhesion proteins (Supplementary Figure 13A, B). Using paxillin as a marker of focal adhesions, we showed that adhesion length was significantly reduced in the ILK knockout cells. Furthermore, the distribution of adhesions, was also altered: ILK knockout cells lacked fibrillar adhesions located towards the centre of a cell. These fibrillar adhesions are distinguished from focal adhesions by their low or absent phosphotyrosine content as was also seen in the ILK knockout cells. Fibrillar adhesions evolve from mature focal adhesions indicating that the ILK knockout cells have a defect in focal adhesion maturation. This has been described previously in ILK knockout fibroblasts (30). Integrin clustering in focal adhesions results in the activation of FAK at the tyrosine (Y)397 autophosphorylation site. Consistent with decreased integrin activation, ILK loss decreased pY397-FAK by 2.4-fold (p-value= 0.047; Figure 6A; quantification Supplementary Figure 13C). We then measured cell-substrate adhesion following an adhesion 'challenge' with a gentle PBS wash. In the ILK knockout cells, there was a complete loss of adherent cells following the adhesion challenge (Figure 6B; data for additional ILK gRNA lines shown in Supplementary Figure 13D). Interestingly, bosutinib, but not eCF506, caused an increase in cell adhesion after the adhesion challenge in the control cells, but both drugs were not able to rescue the adhesion defect in the ILK knockout cells (Figure 6B and Supplementary 13D). It is not clear why bosutinib increases adhesion in the control cells as the reduction in adhesion observed with eCF506 is more consistent with the well documented role of SRC family kinases in cell-matrix adhesion (32). eCF506 reduced pY397-FAK levels in both control and ILK knockout cells. This can be attributed to the ability of eCF506 to bind SRC in an inactive conformation leading to destabilisation of the FAK-SRC complex and a loss of FAK phosphorylation. In contrast, bosutinib binds SRC in an active conformation, thus stabilising FAK-SRC binding and Y397-FAK phosphorylation (24). The loss of ILK did not alter the ability of SRC and FAK to form a complex,

although the levels of pY397-FAK complexed with SRC was reduced reflecting the reduced phosphorylation of FAK on Y397 in the *ILK* knockout cells (Supplementary Figure 13E).

To address whether inhibition of FAK, could also synergise with ILK loss, ILK knockout cells were treated with the FAK inhibitor PND-1186. There was no difference in EC₅₀ values for PND-1186 between the PX459v2 and ILK knockout cells, although pY397-FAK was effectively inhibited in both (Figure 6C and Supplementary Figure 14A). pY419-SRC was unaltered following PND-1186 treatment (Supplementary Figure 14A, B). Previous studies have shown that inhibition of FAK is not sufficient to reduce phosphorylation of SRC and its substrates in integrin adhesion complexes suggesting that either the residual FAK activity is sufficient to maintain SRC-dependent phosphorylation events, or that SRC is activated in a FAK-independent manner (33). To determine whether downstream signalling pathways were differentially regulated following FAK and SRC inhibitor treatment in the ILK knockout cells, a RPPA was carried out. This identified 24 and 10 uniquely differentially regulated proteins and phosphoproteins following bosutinib and PND-1186 respectively (p-value <0.05, log2 fold change), while 13 were common to both drugs (Supplementary Figure 14C). Taken together these data suggest that although SRC and FAK have overlapping functions the reduction in Y397-FAK phosphorylation following FAK inhibitor treatment is not sufficient to prevent downstream SRC signalling and/or that they have independent signalling roles.

Initial microscopy observations showed that bosutinib treatment (EC₂₀; 0.9 μ M) led to a rapid "rounding up" of MDA-MB-231 *ILK* knockout cells within one hour of addition (Supplementary Figure 15A, B). We investigated this change by quantifying the number of total and 'rounded cells' following bosutinib treatment using the IncuCyte S3 (Figure 6D). Within the first hour of bosutinib addition, there was an increase in the number of rounded cells (cells with an area <600 μ m² and eccentricity <0.65): in the *ILK* knockout cells there was a 2.6-fold increase in rounded cells (p-value <0.0001) following bosutinib treatment compared to the ILK gRNA 2 DMSO-treated cells (Figure

6D). This cell rounding was accompanied by an immediate decrease in total cell number in the MDA-MB-231 *ILK* knockout cells, but not PX459v2 cells (Figure 6E), and a 3.0-fold reduction (p-value <0.0001) in cell number after 72 h in the *ILK* knockout cells as well as a 1.3-fold reduction (p-value =0.045) in the PX459v2 cells. Tracking of apoptotic cells using a fluorescent reporter of caspase 3/7 activity showed that apoptotic cell numbers correlated with the rounded phenotype (Supplementary Figure 15C, D). Taken together with the increased apoptosis observed by FACS in the *ILK* knockout cells following treatment with bosutinib over 4 days (Figure 3F), this indicates that the cell rounding and loss of adhesion contributes to the cell death and decreased cell numbers observed. Thus, the increased sensitivity to bosutinib following ILK loss can be attributed to reduced adhesiveness when SRC inhibition is combined with ILK loss. Interestingly, siRNA targeting of *ILK* in MDA-MB-231 cells did not result in enhanced sensitivity to bosutinib or eCF506 despite reducing ILK by 90% (Supplementary Figure 16), suggesting that complete loss of ILK is required to disrupt adhesion complexes in the MDA-MB-231 cells. In support of this, quantification of focal adhesions in the siRNA *ILK* treated cells showed that there was no difference in focal adhesions in contrast to the disruption of adhesions seen in the sgRNA *ILK* knockout cells (Supplementary Figure 13B).

ILK loss enhances SRC inhibition in vivo

Finally, we characterised the effect of combined SRC inhibition and ILK loss on tumour growth *in vivo*. Immunodeficient R2G2 mice were subcutaneously injected with MDA-MB-231 cells with or without ILK, and once tumours had established, were treated with either bosutinib or eCF506 at doses previously shown to inhibit SRC activity in tumours (24)(Figure 7A). The rapid growth of the control tumours resulted in all animals being sacrificed by day 24 of treatment which was extended to day 28 in the bosutinib treated animals. In contrast eCF506 reduced the growth of the control tumours and all animals were sacrificed at day 35 at the end of the experiment. There was a delay in tumour growth in the ILK gRNA tumours, but once established the tumours also grew rapidly and

animals were sacrificed after 35 days of treatment. Treatment with eCF506 completely blocked the growth of the ILK gRNA tumours (Figure 7A) and analysis of the final tumours showed a significant reduction in tumour volumes (Figure 7B). Western blot analysis confirmed effective inhibition of SRC in tumours from both the bosutinib and eCF506 treated animals (Figure 7C, D). Ki67 positivity in the tumours was analysed as a measure of proliferation and showed that there was no difference in the proliferation of the ILK deficient tumours compared to the controls at the end point of the study when the tumours were collected (Figure 7E). Taken together with the lack of effect of ILK loss on proliferation in 2D (Figure 3C), this indicates that ILK is not required for the intrinsic proliferative capacity of the cells. Bosutinib treatment prevented the rapid growth of the ILK gRNA tumours at the later time points, however final tumour volumes and Ki67 positivity were not significantly different between the ILK gRNA vehicle and bosutinib treated animals (Figure 7). In contrast treatment with eCF506 completely blocked the growth of the ILK knockout tumours. The difference in activity of bosutinib and eCF506 is not fully understood but may reflect the reported ability of eCF506 to inhibit SRC in an inactive conformation leading to disruption of protein-protein interactions in addition to inhibition of SRC kinase activity (24), as pY418 SRC was reduced in both bosutinib and eCF506 treated tumours (Figure 7C, D).

Discussion

SRC/ABL TKIs such as bosutinib and dasatinib have been investigated for treating breast cancer for some time (6,8,9,34,35). Yet so far these drugs have had limited clinical efficacy in solid tumours, despite attempts to validate potential gene-signatures of sensitivity (36,37). We aimed to identify possible synergistic gene-drug interactions in an attempt to understand the molecular determinants of bosutinib sensitivity and discover new rational combination therapies for SRC inhibitors. Using an unbiased genome-wide CRISPR-Cas9 knockout screen, we identified loss of the ILK-Parvin-PINCH complex as the key enhancer of SRC inhibition in *in vitro* and *in vivo* breast cancer models. ILK serves as a key scaffold protein in cell–matrix interactions and focal adhesions, tethering cells to

their surrounding environment and interacting with multiple signaling pathways through interactions with its binding partners Parvin and PINCH (29). Furthermore, Horton and co-workers identified a consensus integrin adhesome comprising 60 proteins representing the core cell adhesion machinery which is centred around four functionally interconnected axes, with the ILK-PINCH-PARVIN complex segregating into one of these (38). SRC is also a key regulator of integrin adhesions and ILK is known to interact with the SH3 domain of SRC demonstrating interconnected signalling pathways (39). However, they also have distinct biological activities and our data support a model in which targeting either SRC or ILK alone is not sufficient to inhibit proliferation and induce apoptosis, but that taking out both of these integrin complex components has a synergistic effect. The RNASeq analysis identified a number of integrins and ECM proteins to be significantly differentially expressed in the ILK knockout cells, with further changes in adhesion related genes seen following treatment with bosutinib. This supports defective adhesion signalling as being the critical pathway involved in the enhanced sensitivity to bosutinib. Interestingly there was no synergism between loss of ILK and treatment with the FAK inhibitor PND-1186 suggesting that the decreased activity of FAK in the ILK knockout cells is already sufficient to disrupt adhesion signalling and prime sensitivity to SRC inhibition. Previous studies have shown that SRC and FAK inhibitors can reduce the phosphorylation of different proteins within integrin adhesion complexes, and that FAK inhibition is not sufficient to reduce phosphorylation of SRC substrates (33). Taken together with our data, this suggests that although SRC and FAK have overlapping functions, the reduction in pY397-FAK following FAK inhibitor treatment is not sufficient to prevent downstream SRC signalling and/or that they have independent signalling roles. Indeed, analysis of signalling pathways that were altered following treatment with either bosutinib or PND-1186 revealed a number of differences in the ILK knockout cells. A greater understanding of the interplay between ILK, SRC and other adhesion signalling pathways in tumours will be required to fully exploit this vulnerability.

Upregulation of ILK is frequently observed in cancer tissues such as breast, colorectal and prostate (40). In addition, in prostate cancer, ILK expression is a promising prognostic marker and is correlated with tumour grade, progression and 5-year survival (41). In breast cancer there is limited data, but one study has reported that ILK overexpression is associated with poor overall survival using immunohistochemistry methods (42), and that mRNA levels are higher in tumour compared to adjacent non-cancerous tissues. We utilised the two largest publicly available breast cancer gene expression datasets (METABRIC and TCGA) to show that high *ILK* expression is associated with poor outcomes for the TNBC and basal subtypes. This is consistent with a number of *in vitro* and *in vivo* studies that have reported ILK-dependent effects on breast cancer proliferation, migration, invasion and tumour growth and initiation (43). Whether there is a subtype specific role for ILK in breast cancer remains to be established.

The potential of ILK as an anticancer target has been recognised for a number of years, with a number of putative inhibitors showing significant suppressive effect on cancer development and progression and hence a potential to target ILK for cancer treatment (40). However, although it was originally thought that ILK exerts its oncogenic effects via its kinase domain, mammalian ILK lacks kinase activity (44,45) and thus off-target effects are anticipated for small molecules designed to target ILK kinase activity (e.g. Cpd 22 (46), QLT0267 (47) and QLT0254 (48)). These putative 'ILK inhibitors' in our hands and others (49) do not act via ILK kinase inhibition and do not mimic *ILK* knockdown (49). Notably, a recent study in a model of chronic myeloid leukemia has found QLT-0267 synergises with dasatinib treatment (50); however, they did not demonstrate that it was working via inhibition of ILK kinase activity, and further investigation into the potential mechanism is required. There is therefore a need to create new bona fide ILK inhibitors that could be used in combination with SRC kinase inhibitors such as bosutinib and eCF506. One possibility is the development of drugs that can disrupt the IPP complex, which has shown some promise (51). This

work provides the first step towards new rational drug combinations of ILK and SRC inhibitors in tumours expressing ILK.

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Author Contributions

Conceptualisation, VB and HB; Methodology, HB, OM and DP; Software, DP; Formal Analysis, HB, BG, DP, AB, AL and AS; Investigation, HB, BG, AL, CT, MM and JC; Writing – Original Draft, HB; Writing – Review & Editing, HB, VB, OM, AL, AU and AB; Supervision, VB; Funding Acquisition, VB.

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Data Availability

The datasets generated during and/or analysed during the current study are available as supplementary files. Additional data is available from the corresponding author on reasonable request.

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Figure Legends

Figure 1. Effect of bosutinib treatment on MDA-MB-231 and MCF-7 cell lines. A) Bosutinib 2D and 3D EC₅₀ values in a panel of breast cancer cell lines. MDA-MB-157 and SKBR3 are unable to form spheroids. B) 3D spheroid cell viability. Spheroid area, Calcein-AM intensity and propidium iodide intensity were quantified using a custom CellProfiler pipeline. Staurosporine (1 μ M) was used as a positive control to induce cell death. C) Western blot analysis of cell lines grown as 3D spheroids. Data are representative of two independent experimental replicates. D) Loss of *ABL1* does not alter bosutinib sensitivity. 2D bosutinib 8-point dilution for half-inhibitory concentrations (EC₅₀). Additional 2D and 3D data are shown in Supplementary Figure 4. All data shown are the mean of at least three independent experimental replicates except MDA-MB-134-VI and MDA-MB-468 which were two independent experimental replicates. ***, p-value < 0.0002, ****, p-value < 0.0001, as determined by a two-way ANOVA with Bonferroni multiple comparisons correction. All error bars are S.E.M.

Figure 2. The ILK-Parvin-PINCH complex is the top enhancer of bosutinib induced cell growth inhibition. A) Overview of the genome-wide CRISPR-Cas9 knockout screen in MDA-MB-231 cells. T0, initial timepoint immediately after puromycin selection; T7, 7 days post puromycin selection; T19, three rounds of drug treatment; T27, five rounds of drug treatment. B) DrugZ analysis of the genome-wide bosutinib CRISPR-Cas9 screen. The gene-level normalised Z-score from the T27 time point is shown. Genes are ranked according to the enhancer genes. The Benjamini and Hochberg

(B&H) false-discovery rate (FDR) was used to calculate significance. Green data points are enhancer genes with FDR <0.05 and red are suppressor genes with FDR <0.05. C) Network map for the enhancer genes identified at T27. Network edges were weighted according to evidence of co-functionality using GeneMANIA in Cytoscape. D) ToppGene gene ontology analysis for the combined T19 and T27 enhancer genes. Blue, cellular component category; Yellow, pathway category. Black dots represent q-values. E) High *ILK* expression correlates with worse survival. Kaplan Meier plot for the METABRIC TNBC subtype. The optimal threshold for dichotomisation of the datasets into high and low *ILK* expression was identified using the *survivALL* R package and 'plotALL' function for multi-cut point analysis using the METABRIC dataset. F) Kaplan Meier plot for the TCGA-PanCancer basal subtype used as a validation dataset. Breast cancer-specific survival was used.

Figure 3. Loss of the ILK-Parvin-PINCH complex potentiates the effects of bosutinib in MDA-MB-231 cells. A) CRISPR-Cas9 targeting *ILK* results in ILK loss in the MDA-MB-231 cell line as assessed by Western blot. B) Bosutinib 8-point dose response curves using normalised cell counts form Hoechst-stained images. Significance test refers to EC_{50} values calculated in Prism. C-G) Cells were treated with the EC_{20} for bosutinib (0.9 μ M). C) ILK loss potentiates bosutinib inhibition in real-time assays. Cells were seeded in 96-well plates at 8,000 cells/well and transferred to the IncuCyte Zoom. Arrow shows drug addition at 24 h post seeding. D) End-point quantification using normalised cell confluence from the IncuCyte software. E) End-point quantification using normalised cell counts form Hoechst-stained images. F) Apoptosis assay using CellEvent Caspase 3/7 and fluorescence-activated cell sorting. All data shown are the mean of at least three independent experimental replicates. *, p-value < 0.05, ***, p-value < 0.001, ****, p-value < 0.0001, as determined by a two-way ANOVA with Bonferroni multiple comparisons correction. All error bars are S.E.M. PX459v2, empty-gRNA CRISPR control cells; gRNA 1A, gRNA 1B and gRNA 2, *ILK* knockout clones.

Figure 4. Loss of the ILK-Parvin-PINCH complex potentiates the effects of eCF506 in MDA-MB-231 cells. A) eCF506 8-point dilution for half-inhibitory concentrations (EC_{50}). B) Apoptosis assay using CellEvent Caspase 3/7 and fluorescence-activated cell sorting. eCF506 treatment (40 nM; EC_{20}). C) ILK loss potentiates eCF506 inhibition in real-time. MDA-MB-231 cells were seeded in 96-well plates at 8,000 cells/well and transferred to the IncuCyte Zoom. Arrow shows drug addition. D) End-point quantification using normalised cell confluence from the IncuCyte software. E) End-point quantification using normalised cell confluence from the IncuCyte software. E) End-point quantification using normalised cell confluences. C-E) An EC_{30} concentration for eCF506 of 60 nM is shown. C-E, Data shown is the mean of two independent experimental replicates. * p-value < 0.05, **** p-value < 0.0001, as determined by a two-way Anova with Bonferroni multiple comparisons correction. All error bars are S.E.M. PX459v2, empty-gRNA CRISPR control cells; gRNA 1A, gRNA 1B and gRNA 2, *ILK* knockout clones.

Figure 5. ILK loss in combination with bosutinib treatment results in differentially expressed genes. A) Principal component analysis using the Limma R package. B) Venn Diagram of differentially expressed genes. A cut-off of FDR (Benjamini & Hochberg) p-value adjustment <0.05 and a log2 fold change of ± 1 was used for the differential expression analysis (DEA). Bosutinib EC₂₀ (0.9 μ M) used. Bosutinib DEA: PX459v2-DMSO vs PX459v2-bosutinib; ILK loss DEA: PX459v2-DMSO vs ILK gRNA 2-DMSO; ILK loss + bosutinib DEA: ILK gRNA 2-DMSO vs ILK gRNA 2-bosutinib. Venn diagram created using the VennDiagram R package. C) Volcano plot of the 504 differentially expressed genes unique to ILK gRNA 2-bosutinib samples shown in B. The 504 unique genes were subject to a further cut-off of p-value <0.05 (horizontal dotted blue line) and a log2 fold change of ± 1 (vertical dotted blue lines) for ILK gRNA 2-bosutinib. D) Heatmap of the 134 unique and significantly changed genes shown in C, across all 12 samples. Heatmap created using the ggplots R package. Euclidean distance and complete linkage clustering were used. E) ToppGene gene ontology analysis for the 134 unique DEGs. The top hits from the biological process category are shown. Black dots represent q-values. F) Integrin- collagen network map from the 134 unique DEGs created using GeneMANIA in Cytoscape. Associated genes were determined in GeneMANIA based on published databases and specific omics publications.

Figure 6. *ILK* knockout in combination with bosutinib treatment causes cell adhesion defects. A) Western blot analysis of cell lines grown in 2D. Data are representative of three independent experimental replicates. B) Adhesion challenge: IncuCyte cell confluence was normalised to DMSO PX459v2. DMSO as solid fill, bosutinib as clear fill. C) 2D 8-point dilution for half-inhibitory concentrations (EC₅₀) for PND-1186. End-point quantification using normalised cell counts form Hoechst-stained images. D) Cell-by-cell mask and a 'rounded cells' classifier whereby area < 600 μ m² and eccentricity <0.65, one hour post drug addition. E) IncuCyte S3 cell-by-cell quantification for MDA-MB-231 cells plated at 8,000 cells per well. Arrow shows drug addition. A bosutinib EC₂₀ (0.9 μ M) concentration was used in all experiments.

Figure 7. ILK loss enhances bosutinib and eCF506 treatment *in vivo*. A) Tumour growth rates for PX459v2 and ILK gRNA 2 tumours growing in immunodeficient Rag2-Il2rg double knockout mice (5 mice per group, 2 tumours per mouse) following treatment with vehicle, bosutinib (75 mg/kg) or eCF506 (40 mg/kg) given once daily via oral gavage. B) Final tumour volumes at day 35. C) Bosutinib and eCF506 inhibit pSrc *in vivo* as shown via western blot. D) Quantification of westerns shown in C. E) Ki67 analysis using QuPath. ***, p-value < 0.0002, ****, p-value < 0.0001, as determined by a two-way ANOVA with Bonferroni multiple comparisons correction. All error bars are S.E.M. Author Manuscript Published OnlineFirst on December 17, 2021; DOI: 10.1158/0008-5472.CAN-21-0373 Author manuscripts have been peer reviewed and accepted for publication but have not yet been edited.





Research.

Figure 2



Research.

Figure 3





Figure 5







Research.





Loss of Integrin-linked kinase sensitizes breast cancer to SRC inhibitors

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