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LYN-activating mutations mediate antiestrogen resistance in estrogen receptor–positive breast cancer

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Estrogen receptor–positive (ER+) breast cancers adapt to hormone deprivation and become resistant to antiestrogen therapy. Here, we performed deep sequencing on ER+ tumors that remained highly proliferative after treatment with the aromatase inhibitor letrozole and identified a D189Y mutation in the inhibitory SH2 domain of the SRC family kinase (SFK) LYN. Evaluation of 463 breast tumors in The Cancer Genome Atlas revealed four LYN mutations, two of which affected the SH2 domain. In addition, LYN was upregulated in multiple ER+ breast cancer lines resistant to long-term estrogen deprivation (LTED). An RNAi-based kinome screen revealed that LYN is required for growth of ER+ LTED breast cancer cells. Kinase assays and immunoblots analyses of SRC substrates in transfected cells indicated that LYN WT has higher catalytic activity than WT protein. Further, LYN D189Y exhibited reduced phosphorylation at the inhibitory Y507 site compared with LYN WT. Other SH2 domain LYN mutants, E159K and K209N, also exhibited higher catalytic activity and reduced inhibitory site phosphorylation. LYN D189Y overexpression abrogated growth inhibition by fulvestrant and/or the PI3K inhibitor BKM120 in 3 ER+ breast cancer cell lines. The SFK inhibitor dasatinib enhanced the antitumor effect of BKM120 and fulvestrant against estrogen-deprived ER+ xenografts but not LYN D189Y-expressing xenografts. These results suggest that LYN mutations mediate escape from antiestrogens in a subset of ER+ breast cancers.

Introduction

LYN is a member of the SRC family of protein tyrosine kinases (SFKs), key regulators of several cellular processes, including cancer cell growth, migration, invasion, and survival (1, 2). Overexpression of LYN, as measured by immunohistochemistry (IHC), is associated with an epithelial-to-mesenchymal transition and correlates with a shorter overall survival in breast cancer (3). SRC engages in bidirectional crosstalk with the estrogen receptor α (ERα) (4), in which its kinase phosphorylates ERα at Y537 (5), resulting in an enhancement of ER transcriptional activity (6).

Two-thirds of breast cancers express ER and/or progesterone receptor (PR), biomarkers indicative of hormone dependence (7). Therapies against ER+ breast cancers inhibit ER function by antagonizing ligand binding to ER (tamoxifen), downregulating ER (fulvestrant), or blocking estrogen biosynthesis and reducing circulating estrogen levels (8) (aromatase inhibitors [AIs]). Although AIs produce an objective tumor response rate of 30% to 40% in patients with metastatic ER+ breast cancer, a significant fraction of patients do not respond or progress quickly on this therapy (9). Thus, elucidating the molecular mechanisms underlying this resistance is critical for improving patient outcome. In addition, discovery of biomarkers predictive of clinical benefit from antiestrogens and prospective identification of patients who are resistant to these therapies are needed.

ER blockade with antiestrogens inhibits tumor cell proliferation in hormone-dependent ER+ breast cancers. This can be measured by IHC for the nuclear antigen Ki67, which identifies cells in the G1/S and M phases of the cell cycle (10). The Immediate Preoperative Anastrozole, Tamoxifen, or Combined with Tamoxifen (IMPACT) study showed that the high Ki67 score in tumors after 2 or 12 weeks of antiestrogen therapy predicts a shorter recurrence-free survival (11, 12). These data suggest that a high tumor cell proliferation (i.e., high Ki67) following treatment with an AI can be used to identify ER+ tumors that are resistant to endocrine therapy and as an unbiased approach to discover molecular effectors of such resistance.

The objective of this study is to identify kinase mutations associated with resistance to estrogen deprivation. We performed deep Kinome sequencing on 4 ER+/HER2 breast cancers that retained high Ki67 scores (14.8%–24.5%) following 2 weeks of treatment with the AI letrozole. We identified a novel D189Y somatic mutation in LYN in an endocrine-resistant tumor, as defined by the Ki67 score after treatment. Although stable overexpression of WT LYN (referred to herein as LYN WT) or the D189Y mutation in LYN (referred to herein as LYN D189Y) accelerated MCF-7 cell growth in estrogen-depleted media, the mutant was more potent than LYN WT at inducing this effect. LYN WT but not LYN D189Y exhibited reduced phosphorylation of the inhibitory Y507 residue, suggesting that this substitution limited the ability of LYN WT to inhibit ER activation.
Results

Deep kinome sequencing identifies a novel D189Y mutation in LYN. NCT00651976 is an IRB-approved clinical trial at Vanderbilt University, in which postmenopausal women with newly diagnosed operable breast cancer consented to treatment with letrozole (2.5 mg/d) for 10 to 21 days prior to surgery (Supplemental Figure IA; supplemental material available online with this article; doi:10.1172/JCI72573DS1). Tumor cell proliferation was assessed by Ki67 IHC in a core biopsy before treatment and in the surgical specimen (after treatment). From among the first 23 patients enrolled in this study, we selected 4 tumors that retained high Ki67 scores (14.8%–24.5%) after 2 weeks of letrozole treatment.

Genomic DNA from 4 formalin-fixed, paraffin-embedded (FFPE) surgically excised tumors and matched normal adjacent breast tissue or histologically confirmed negative lymph nodes was deep sequenced using a capture approach, including 517 kinases and 46 cancer-related genes from the Agilent SureSelect Kinome Kit (Supplemental Table 2). Tumor cellularity ranged from 20% to 50%. Annotated variant call files were provided by Expression Analysis for the sequenced tumors. To prioritize variants for molecular analysis, we filtered all variants against the reference hg19 sequence that (a) were not in the capture-targeted region, (b) were known SNPs (based on the dbSNP database), or (c) were not exonic or were synonymous in their protein translation. These criteria resulted in 671 candidate variations among the 4 tumor/normal pairs. Next, somatic status was determined by comparing each tumor to its matched normal specimen. For these 4 tumors, 28, 15, 14, and 19 somatic mutations were identified, respectively. Resulting hits were manually screened for likely artifacts in the Integrative Genome Viewer for the presence of strand bias or variants lying only at read ends. After further manual filtering for likely artifacts, we identified a series of candidate somatic mutations in kinases (EPHA7, LYN, DYRKIA, DYRK2, MYLK3, PTK2, TSSK4, CSNK1G3, WNK2, and TEC; Table 1). All 4 tumors contained a mutation in PIK3CA, the gene encoding the p110α catalytic subunit of PI3K; 3 were hot-spot activating mutations (Q546E, H1047R); and 1 was T1025A. PIK3CA hot-spot mutations are associated with increased PI3K catalytic activity (13), cell transformation in vitro and in transgenic mouse models, and resistance to anoikis (14, 15). Related to the T1025A variant, PIK3CA with a substitution in the same base, T1025S, has been shown to transform chicken embryo fibroblasts (16).

One tumor, referred to as tumor C, showed a previously unreported D189Y somatic mutation in LYN (variant frequency of 8%; Table 1), which was not detected in adjacent normal breast tissue.

Table 1. Deep kinome sequencing of DNA from surgical specimens after treatment

<table>
<thead>
<tr>
<th>Tumor</th>
<th>Gene symbol</th>
<th>% Variation in tumor</th>
<th>Amino acid change</th>
<th>Domain</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>TEC</td>
<td>9%</td>
<td>V→I (V578I)</td>
<td>Kinase</td>
</tr>
<tr>
<td></td>
<td>CSNK1G3</td>
<td>9%</td>
<td>T→A (T1267A)</td>
<td>Kinase</td>
</tr>
<tr>
<td></td>
<td>TSSK4</td>
<td>3%</td>
<td>L→V (L295V)</td>
<td>Kinase</td>
</tr>
<tr>
<td></td>
<td>YP53</td>
<td>41%</td>
<td>R→G (R249G)</td>
<td>DNA binding</td>
</tr>
<tr>
<td></td>
<td>PIK3CA</td>
<td>16%</td>
<td>H→R (H1047R)</td>
<td>Kinase</td>
</tr>
<tr>
<td>B</td>
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<td>10%</td>
<td>Y→H (Y143H)</td>
<td>Ligand binding</td>
</tr>
<tr>
<td></td>
<td>PIK3CA</td>
<td>10%</td>
<td>Q→E (Q546E)</td>
<td>Helical</td>
</tr>
<tr>
<td>C</td>
<td>LYN</td>
<td>8%</td>
<td>D→Y (D189Y)</td>
<td>SH2</td>
</tr>
<tr>
<td></td>
<td>ATM</td>
<td>10%</td>
<td>D→H (D2987H)</td>
<td>None</td>
</tr>
<tr>
<td></td>
<td>DYRK2</td>
<td>18%</td>
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<td>Kinase</td>
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<td>Q→E (Q579E)</td>
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<td></td>
<td>PIK3CA</td>
<td>14%</td>
<td>T→A (T1025A)</td>
<td>Catalytic</td>
</tr>
<tr>
<td>D</td>
<td>PTK2</td>
<td>1%</td>
<td>A→V (A450V)</td>
<td>Unknown</td>
</tr>
<tr>
<td></td>
<td>WNK2</td>
<td>10%</td>
<td>G→R (G2199R)</td>
<td>None</td>
</tr>
<tr>
<td></td>
<td>DYRK2</td>
<td>3%</td>
<td>M→I (M226I)</td>
<td>Kinase</td>
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<tr>
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<td>DYRK1A</td>
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<td>None</td>
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<td></td>
<td>PIK3CA</td>
<td>2%</td>
<td>H→R (H1047R)</td>
<td>Kinase</td>
</tr>
</tbody>
</table>

Mutations identified by deep kinome sequencing using a capture approach (Agilent SureSelect Kinome Kit) and the Illumina Genome AnalyzerIIx in 4 ER+ HER2– primary breast tumors that retained high Ki67 scores (14.8%–24.5%) after 2 weeks of letrozole treatment.

was deep sequenced using a capture approach, including 517 kinases and 46 cancer-related genes from the Agilent SureSelect Kinome Kit (Supplemental Table 2). Tumor cellularity ranged from 20% to 50%. Annotated variant call files were provided by Expression Analysis for the sequenced tumors. To prioritize variants for molecular analysis, we filtered all variants against the reference hg19 sequence that (a) were not in the capture-targeted region, (b) were known SNPs (based on the dbSNP database), or (c) were not exonic or were synonymous in their protein translation. These criteria resulted in 671 candidate variations among the 4 tumor/normal pairs. Next, somatic status was determined by comparing each tumor to its matched normal specimen. For these 4 tumors, 28, 15, 14, and 19 somatic mutations were identified, respectively. Resulting hits were manually screened for likely artifacts in the Integrative Genome Viewer for the presence of strand bias or variants lying only at read ends. After further manual filtering for likely artifacts, we identified a series of candidate somatic mutations in kinases (EPHA7, LYN, DYRKIA, DYRK2, MYLK3, PTK2, TSSK4, CSNK1G3, WNK2, and TEC; Table 1). All 4 tumors contained a mutation in PIK3CA, the gene encoding the p110α catalytic subunit of PI3K; 3 were hot-spot activating mutations (Q546E, H1047R); and 1 was T1025A. PIK3CA hot-spot mutations are associated with increased PI3K catalytic activity (13), cell transformation in vitro and in transgenic mouse models, and resistance to anoikis (14, 15). Related to the T1025A variant, PIK3CA with a substitution in the same base, T1025S, has been shown to transform chicken embryo fibroblasts (16).

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Results

Deep kinome sequencing identifies a novel D189Y mutation in LYN. NCT00651976 is an IRB-approved clinical trial at Vanderbilt University, in which postmenopausal women with newly diagnosed ER+ HER2– operable breast cancer consented to treatment with letrozole (2.5 mg/d) for 10 to 21 days prior to surgery (Supplemental Figure IA; supplemental material available online with this article; doi:10.1172/JCI72573DS1). Tumor cell proliferation was assessed by Ki67 IHC in a core biopsy before treatment and in the surgical specimen (after treatment). From among the first 23 patients enrolled in this study, we selected 4 tumors that retained the highest Ki67 level in the biopsy after treatment (14.8%–24.5%; Supplemental Table 1). Consistent with a previous report (12), we considered an after treatment Ki67 score of >10% as a surrogate of endocrine resistance.

Genomic DNA from 4 formalin-fixed, paraffin-embedded (FFPE) surgically excised tumors and matched normal adjacent breast tissue or histologically confirmed negative lymph nodes was deep sequenced using a capture approach, including 517 kinases and 46 cancer-related genes from the Agilent SureSelect Kinome Kit (Supplemental Table 2). Tumor cellularity ranged from 20% to 50%. Annotated variant call files were provided by Expression Analysis for the sequenced tumors. To prioritize variants for molecular analysis, we filtered all variants against the reference hg19 sequence that (a) were not in the capture-targeted region, (b) were known SNPs (based on the dbSNP database), or (c) were not exonic or were synonymous in their protein translation. These criteria resulted in 671 candidate variations among the 4 tumor/normal pairs. Next, somatic status was determined by comparing each tumor to its matched normal specimen. For these 4 tumors, 28, 15, 14, and 19 somatic mutations were identified, respectively. Resulting hits were manually screened for likely artifacts in the Integrative Genome Viewer for the presence of strand bias or variants lying only at read ends. After further manual filtering for likely artifacts, we identified a series of candidate somatic mutations in kinases (EPHA7, LYN, DYRKIA, DYRK2, MYLK3, PTK2, TSSK4, CSNK1G3, WNK2, and TEC; Table 1). All 4 tumors contained a mutation in PIK3CA, the gene encoding the p110α catalytic subunit of PI3K; 3 were hot-spot activating mutations (Q546E, H1047R); and 1 was T1025A. PIK3CA hot-spot mutations are associated with increased PI3K catalytic activity (13), cell transformation in vitro and in transgenic mouse models, and resistance to anoikis (14, 15). Related to the T1025A variant, PIK3CA with a substitution in the same base, T1025S, has been shown to transform chicken embryo fibroblasts (16).

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Figure 1. Deep kinome sequencing identifies a D189Y mutation in LYN. (A) Sanger sequencing of tumor C verified the presence of the D189Y mutation in LYN; the point mutation (G to T) is shown by the arrow. (B) Lysates from 10 ER+ HER2– tumors from patients treated 10–21 days with letrozole prior to surgery were analyzed by RPPA. Antibody signal intensity was log normalized and correlated to Ki67 score after letrozole treatment (POST).

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cancer correlation ($P = 0.006$) between Y416 phosphorylated SRC (P-SRC) (which detects all activated SFKs) and a high Ki67 score after letrozole (Figure 1B).

RNAi screening identifies LYN as required for hormone-independent ER+ breast cancer growth. To complement this study, we screened a siRNA library targeting 779 kinases in MCF-7 breast cancer cells resistant to long-term estrogen deprivation (LTED) (20, 21). This screen identified LYN as one of the top hits whose knockdown markedly reduced growth of MCF-7 /LTED cells (Figure 2A). Knockdown of LYN with an alternative siRNA (not in the screening library) reduced LYN but not SRC protein levels, as measured by immunoblot analysis (Figure 2B), and significantly inhibited growth of MCF-7 /LTED, MDA-361/LTED, HCC-1428/LTED, and ZR75-1/LTED cells (Figure 2C), suggesting that LYN plays a role in hormone-independent ER+ breast cancer growth.

LYN expression was upregulated in 3 of 4 LTED cell lines compared with their parental counterparts (Figure 3A). These changes were paralleled by LYN mRNA expression in MCF-7/LTED and MDA-361/LTED cells (3.6-fold and 1.5-fold, respectively) compared with their respective parental cells, whereas the other SFKs showed variable changes (Figure 3B). YES1 mRNA was upregulated 1.4-fold in MCF-7/LTED cells, and the other SFKs were downregulated or unchanged.Only LYN mRNA expression was significantly upregulated in MDA-361/LTED cells (Figure 3B).

LYND189Y accelerates hormone-independent growth and blocks the antitumor effect of ER and PI3K antagonists. We next investigated the role of LYND189Y in estrogen-independent growth using ER+/from the same patient (Supplemental Figure 1B). We next confirmed mRNA expression of DI89Y LYN by cDNA sequencing in both directions (Supplemental Figure 1C). The allele frequency of DI89Y, which is comparable to that of mutant PIK3CA (14%) in the same tumor, probably reflects the clonal nature of this alteration as well as dilution with WT reads in DNA from adjacent normal tissue that could not be macrodissected out. DI89Y (565G>T) is located in the LYN SH2 domain, which binds phosphorylated tyrosines (2, 17). The DI89Y substitution was predicted to have a high functional affect on LYN, as assessed by mutationassessor.org (http://mutationassessor.org), which estimates the potential consequences of amino acid substitutions in protein function (18). Sanger sequencing confirmed the presence of DI89Y LYN in DNA isolated from the tumor before treatment (Figure 1A).

We next used the cBio Cancer Genomics Portal (19) to explore LYN mutations in 463 invasive breast carcinomas included in The Cancer Genome Atlas (TCGA; accessed September 9, 2014). There were 4 LYN mutations, E159K, K209N, G418R, and A503D, with the first 2 located in the SH2 domain and the last 2 located in the kinase domain. Despite its low frequency, LYN is the most commonly mutated SFK family member in invasive breast cancers in TCGA (4, 2, 1, and 1 case for LYN, YES1, HCK, and BLK, respectively). LYN mutations have been also reported in low frequency in melanoma and endometrial, lung, and colorectal carcinoma (Supplemental Table 3). Reverse-phase protein array (RPPA) analysis in lysates from 10 tumors of patients enrolled in the NCT00651976 trial (Supplemental Table 4) revealed a significant correlation ($P = 0.006$) between Y416 phosphorylated SRC (P-SRC) (which detects all activated SFKs) and a high Ki67 score after letrozole (Figure 1B).

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**LYND189Y accelerates hormone-independent growth and blocks the antitumor effect of ER and PI3K antagonists.** We next investigated the role of LYND189Y in estrogen-independent growth using ER+/
PI3KCA mutant MCF-7 cells stably transduced with GFP (control), WT LYN, or D189Y LYN V5-tagged vectors. The D189Y mutation was verified by Sanger sequencing (Supplemental Figure 2). Expression of LYN\textsuperscript{WT} and LYN\textsuperscript{D189Y} in MCF-7 cells was equivalent, as assessed by immunoblot with a V5 antibody (Figure 4A). Treatment with fulvestrant downregulated levels of ER and its target gene product IGF-I receptor (IGF-IR), but this effect was partially abrogated in MCF-7/LYN\textsuperscript{WT} and MCF-7/LYN\textsuperscript{D189Y} cells, particularly in cells expressing the mutant. Overexpression of LYN\textsuperscript{D189Y} also prevented the complete inhibition of S473 phosphorylated AKT (P-AKT) and phosphorylated PRAS40 (P-PRAS40) levels mediated by BKM120 (Figure 4A), suggesting that mutant LYN contributes to PI3K activation.

Although stable transduction of either LYN\textsuperscript{WT} or LYN\textsuperscript{D189Y} accelerated MCF-7 cell growth in estrogen-depleted media, the mutant was significantly more potent than LYN\textsuperscript{WT} at inducing this effect (P < 0.001). Further, LYN\textsuperscript{D189Y} overexpression dampened the growth inhibitory effect of the ER downregulator fulvestrant or the pan-PISK inhibitor BKM120 (ref. 22 and Figure 4B). More pronounced effects were seen in 3D Matrigel in estrogen-free media: MCF-7/LYN\textsuperscript{D189Y} cells formed larger colonies than MCF-7/LYN\textsuperscript{WT} cells (P < 0.0001) and were less sensitive to fulvestrant and BKM120 (P < 0.05; Figure 4C). To extend these results to other cell lines, we stably transduced HCC1428 and ZR75-1 cells with GFP, LYN\textsuperscript{WT}, and LYN\textsuperscript{D189Y} vectors (Supplemental Figure 3). Stably transduced HCC1428 and ZR75-1 cells showed similar results in 3D Matrigel in estrogen-free media upon treatment with fulvestrant and BKM120 (Figure 4C). In Supplemental Figure 4, we show these 3D growth data normalized to untreated controls.

We finally examined the effects of LYN\textsuperscript{WT} or LYN\textsuperscript{D189Y} on caspase-dependent apoptosis in MCF-7 cells. Treatment with BKM120 significantly increased caspase-7 activity in MCF-7 cells, whereas treatment with fulvestrant did not (Figure 4D). Overexpression of LYN\textsuperscript{D189Y} but not LYN\textsuperscript{WT} completely suppressed BKM120-induced caspase-7 activity (Figure 4D), suggesting that LYN\textsuperscript{D189Y} can rescue cells from the apoptosis induced by inhibition of PI3K.

SRC inhibitors block growth of MCF-7 cells expressing LYN\textsuperscript{WT} or LYN\textsuperscript{D189Y}. We next investigated whether the gain of function induced by overexpression of WT or mutant LYN is reversed by the small molecule SFK inhibitors dasatinib (LYN IC\textsubscript{50}, 8.5 nM) (23, 24) and AZD0530 (LYN IC\textsubscript{50} 5 nM) (25). Dasatinib but not AZD0530 markedly suppressed Y416 P-SRC levels in MCF-7/LYN\textsuperscript{D189Y} cells (Supplemental Figure 4), suggesting that the mutant stabilizes ER and/or regulates ER transcription. Consistent with previous results, dasatinib and AZD0530 inhibited MCF-7/LYN\textsuperscript{D189Y} cell growth by 1.8-fold and 2.2-fold, respectively (Figure 5C). MCF-7/LYN\textsuperscript{WT} and MCF-7/LYN\textsuperscript{D189Y} cells were still sensitive to both SFK inhibitors, albeit less so to AZD0530 (Figure 5C). In Supplemental Figure 6, we show the data shown in Figure 5C normalized to untreated controls.

LYN\textsuperscript{D189Y} exhibits increased catalytic activity and signaling output compared with that of LYN\textsuperscript{WT}. The differences in the cellular effects of LYN\textsuperscript{WT} and LYN\textsuperscript{D189Y} (Figure 4) suggested the mutation conferred LYN a gain of function. Consistent with this notion, overexpression of LYN\textsuperscript{WT} or LYN\textsuperscript{D189Y} resulted in increased phosphorylation of FAK, IGF-IR/InsR, HER3, EGFR, and MAPK in MCF-7 cells maintained in FBS-containing media and following serum starvation (Figure 6, A and B). Activated SFK, as measured by Y416 P-SRC levels, was similar in LYN\textsuperscript{WT} and LYN\textsuperscript{D189Y} cells. However, overexpression of LYN\textsuperscript{D189Y} increased phosphorylation of FAK (a downstream target of SFKs) at Y576/577 and Y925, EGFR at the SRC site Y845, HER3, and IRS-1 above levels induced by ectopic expression of LYN\textsuperscript{WT} (Figure 6, A and B), suggesting that LYN\textsuperscript{D189Y} increases the catalytic activity of LYN. On the other hand, HCC1428 and ZR75-1 cells stably transduced with LYN\textsuperscript{D189Y} but not LYN\textsuperscript{WT} contained markedly higher levels of Y416 P-SRC compared with their respective parental cells (Figure 6C).

To interrogate a biochemical difference between WT and mutant LYN, we performed in vitro kinase assays. V5-tagged LYN\textsuperscript{WT} and LYN\textsuperscript{D189Y} were precipitated from stably transduced MCF-7 cells.
Figure 4. LYN<sup>D189Y</sup> accelerates hormone-independent growth and counteracts the effect of PI3K and ER inhibitors. (A) Cells were treated with 10% DCC-FBS with or without 1 μM BKM120 or 1 μM fulvestrant for 24 hours. Protein lysates were prepared and analyzed by immunoblot with the indicated antibodies. (B and C) MCF-7 cells stably transduced with GFP, LYN<sup>WT</sup>, or LYN<sup>D189Y</sup> were treated with 10% DCC-FBS with or without 1 μM BKM120 or 1 μM fulvestrant for 12 days. Media and drugs were replenished every 2 to 3 days. (B) Cells were fixed and stained with crystal violet. Representative images and quantitation of integrated intensity (percentage of control) are shown (n = 2; *P < 0.05 vs. respective GFP, #P < 0.01 vs. respective LYN<sup>WT</sup>). (C) Cells were cultured in Matrigel with or without 1 μM BKM120 or 1 μM fulvestrant and photographed on day 15. The average size of each cellular structure was quantitated using ImageJ. Data are presented as percentage of control (n = 2; *P < 0.05 vs. respective GFP, #P < 0.05 vs. respective LYN<sup>WT</sup>). (D) Cells were treated overnight in serum-free media with or without 1 μM BKM120 or 1 μM fulvestrant and then analyzed using the Caspase-Glo 3/7 assay. Data are presented as percentage of control (n = 2; *P < 0.05 vs. Con, *P < 0.05 vs. GFP BKM120).
LYN<sup>D189Y</sup> showed a 2.2-fold increase in kinase activity compared with LYN<sup>WT</sup> in the absence of added ATP (Figure 6D). However, based on the location of the mutation in the SH2 domain of LYN, we speculated that D189Y would mainly prevent an inactive conformation of the kinase. SFKs are phosphorylated by C-terminal SRC kinase (CSK) in a C-terminal regulatory tyrosine, which keeps the SFKs in an inactive conformation involving an intramolecular interaction between this residue and the SH2 domain (17). Upon dephosphorylation of the C-terminal tyrosine (Y507 in LYN), SFKs adopt an open conformation in which Y416 (Y396 in LYN) in the activation loop is phosphorylated and the kinase is fully activated. We speculated that D189Y would reduce the interaction between the regulatory tyrosine and the SH2 domain, thus preventing the inactive conformation of LYN. Indeed, LYN<sup>D189Y</sup> overexpression markedly decreased phosphorylation of LYN at the Y507 C-terminal regulatory site compared with LYN<sup>WT</sup> in MCF-7, HCC-1428, and ZR75-1 cells (Figure 6, B and C). To corroborate these data, we precipitated V5-tagged LYN<sup>WT</sup> and LYN<sup>D189Y</sup> from MCF-7 cell lysates and performed immunoblot analysis with an antibody that recognizes LYN phosphorylated in Y507 (P-LYN). Y507 P-LYN content in LYN<sup>WT</sup> and LYN<sup>D189</sup> was corrected for V5 expression in the V5 pulldowns, as measured by densitometry. Compared with LYN<sup>WT</sup>, LYN<sup>D189Y</sup> contained 71% lower levels of Y507 P-LYN in 10% FBS and 14% lower levels of Y507 P-LYN in serum-free media (Figure 6E).

We next extended these studies to other LYN mutants identified in TCGA. V5-tagged lentiviral vectors encoding LYN<sup>E159K</sup>, LYN<sup>K209N</sup>, LYN<sup>A370T</sup>, LYN<sup>G418R</sup>, and LYN<sup>A503D</sup> were prepared and stably transduced into MCF-7 cells (Figure 7A). We were unable to express the A503D variant. Similar to LYN<sup>D189Y</sup>, the 2 other SH2 domain mutants (LYN<sup>E159K</sup> and LYN<sup>K209N</sup>) expressed lower levels of Y507 P-LYN (Figure 7, A and B). Compared with MCF-7/LYN<sup>WT</sup> cells, LYN<sup>E159K</sup> and LYN<sup>K209N</sup> cells exhibited enhanced growth in estrogen-free media and were less sensitive to the inhibitory effect of fulvestrant and BKM120 (Figure 7C). In Supplemental Figure 7, we show these growth data normalized to untreated controls.

**Pharmacological inhibition of the SFKs enhances the antitumor effect of BKM120 and fulvestrant in vivo.** The results of the RNAi screen in LTED ER<sup>+</sup> breast cancer cells (Figure 2), the higher levels of LYN in 3 of 4 LTED cell lines (Figure 3), and the growth advantage conferred by LYN<sup>WT</sup> (Figure 4) all suggested that overexpression of SFKs, such as LYN, can mediate escape from endocrine therapy. If so, addition of a SFK inhibitor should improve the effect of this therapy and/or delay acquired drug resistance. We examined this possibility in ER<sup>+</sup>/PIK3CA mutant MCF-7 xenografts. We recently reported that growth of these xenografts established in ovariectomized mice and in the absence of added estrogens is synergistically inhibited by a combination of fulvestrant and BKM120 (21). Inhibition of the SFKs with dasatinib modestly increased the growth inhibitory effect of BKM120 in MCF-7/LYN<sup>D189Y</sup> cells, while it significantly increased the effect of fulvestrant (Figure 8A). In Supplemental Figure 8, we show these growth data normalized to untreated controls.

We next assessed the effects of combined inhibition of ER, PI3K, and SFKs on tumor growth in vivo. MCF-7 xenografts were established in ovariectomized athymic mice. Treatment with BKM120 and fulvestrant or BKM120, fulvestrant, and dasatinib,
but not dasatinib alone, inhibited growth of established tumors compared with vehicle (P < 0.0001; Figure 8B). Treatment with triple combination induced near-complete tumor regression and was significantly more effective than BKM120 and fulvestrant or dasatinib alone (P < 0.01; Figure 8B). Treatment with BKM120 and fulvestrant as well as BKM120, fulvestrant, and dasatinib decreased ER, PR, IGF-IR, P-AKT, P-PRAS40, phosphorylated S6 (P-S6), and phosphorylated MAPK (P-MAPK) tumor levels. Treatment with dasatinib alone reduced P-AKT, P-S6, and P-MAPK levels as well as those of ER, PR, and IGF-IR (Figure 8C), suggesting that, in untreated tumors, activated SFKs stabilize ER and its transcriptional output in vivo. Consistent with drug-mediated target inhibition, dasatinib-treated tumors exhibited lower levels of Y416 P-SRC, as measured by IHC (Figure 8D). Overall, the drug combinations were well-tolerated, with <10% weight loss (Supplemental Figure 9). These results suggest that pharmacological inhibition of SFKs enhances the antitumor effect of BKM120 and fulvestrant against estrogen-deprived MCF-7 xenografts in vivo.

Discussion

Although endocrine therapies are effective against ER+ breast cancers, de novo and acquired drug resistance remain a significant clinical problem. The factors mediating resistance to antiestrogens have been the subject of considerable investigation and include alterations in pathways such as E2F/cyclin D/CDKs (21), PI3K/mTOR (20), and FGFR (26). To further explore genetic alterations that may lead to intrinsic endocrine resistance, we performed deep kinome sequencing on 4 ER+/HER2– human breast cancers that retained high Ki67 levels (a cellular marker of endocrine resistance) following short-term treatment with the AI letrozole. In one of these tumors, we identified a novel D189Y mutation in the SH2 domain of LYN.
EGFR, HER3, STAT3, AKT, and MAPK. However, LYN D189Y expression resulted in higher phosphorylation of FAK, EGFR, HER3, and IRS-1 compared with LYN WT. Although stable transduction of either LYN WT or LYN D189Y accelerated MCF-7, HCC1428, and ZR75-1 cell growth in estrogen-depleted media, the mutant was substantially more potent than LYN WT at inducing this effect.

We also showed a concurrent decrease in phosphorylation of LYN at the C-terminal regulatory site Y507 upon expression of the mutant. RNAi screening identified LYN as being required for hormone-independent ER+ breast cancer growth, and knockdown of LYN inhibited growth of 4 of 4 LTED cell lines. We next studied the functional role of LYN mutations in several ER+ human breast cancer cell lines. Overexpression of LYN WT or LYN D189Y in MCF-7 cells increased phosphorylation of SFKs (at Y416) and known targets of SFKs, including FAK, IGF-IR/InsR, EGFR, HER3, STAT3, AKT, and MAPK. However, LYN D189Y expression resulted in higher phosphorylation of FAK, EGFR, HER3, and IRS-1 compared with LYN WT. Although stable transduction of either LYN WT or LYN D189Y accelerated MCF-7, HCC1428, and ZR75-1 cell growth in estrogen-depleted media, the mutant was substantially more potent than LYN WT at inducing this effect.

We also showed a concurrent decrease in phosphorylation of LYN at the C-terminal regulatory site Y507 upon expression of the mutant.

Analysis of TCGA breast cancer data revealed 4 LYN mutations, 2 in the SH2 domain of LYN. Further, RNAi screening identified LYN as being required for hormone-independent ER+ breast cancer growth, and knockdown of LYN inhibited growth of 4 of 4 LTED cell lines. We next studied the functional role of LYN mutations in several ER+ human breast cancer cell lines. Overexpression of LYN WT or LYN D189Y in MCF-7 cells increased phosphorylation of SFKs (at Y416) and known targets of SFKs, including FAK, IGF-IR/InsR, EGFR, HER3, STAT3, AKT, and MAPK. However, LYN D189Y expression resulted in higher phosphorylation of FAK, EGFR, HER3, and IRS-1 compared with LYN WT. Although stable transduction of either LYN WT or LYN D189Y accelerated MCF-7, HCC1428, and ZR75-1 cell growth in estrogen-depleted media, the mutant was substantially more potent than LYN WT at inducing this effect.

We also showed a concurrent decrease in phosphorylation of LYN at the C-terminal regulatory site Y507 upon expression of the mutant.
ing signaling duration and/or amplitude. However, molecular modeling of autoinhibited SRC did not indicate how the mutation might alter the Y507-mediated conformation of the protein due to its distal location. Additionally, modeling of an activating peptide substrate did not suggest a direct role for D189 in the interaction between the SH2 domain and the activating peptide. Since no active SFK structures that include the SH2 domain are available, determining whether a tyrosine substitution at D189 might alter intermolecular or intramolecular interactions in the active state of LYN D189Y mutant in MCF-7, HCC1428, and ZR75-1 cells (Figure 6, B and C). Phosphorylation of Y507 by c-SRC tyrosine kinase (CSK) maintains LYN in an inactive conformation, which requires an intramolecular interaction between Y507 and the SH2 domain. Upon dephosphorylation, LYN adopts an open conformation in which the activation loop Y396 is phosphorylated and the kinase is activated (17). Thus, we speculated that the LYN D189Y mutation may prevent/hinder the interaction between Y507 and the SH2 domain, thus abrogating LYN inactivation and potentially increasing signaling duration and/or amplitude. However, molecular modeling of autoinhibited SRC did not indicate how the mutation might alter the Y507-mediated conformation of the protein due to its distal location. Additionally, modeling of an activating peptide substrate did not suggest a direct role for D189 in the interaction between the SH2 domain and the activating peptide. Since no active SFK structures that include the SH2 domain are available, determining whether a tyrosine substitution at D189 might alter intermolecular or intramolecular interactions in the active state of LYN D189Y mutant in MCF-7, HCC1428, and ZR75-1 cells (Figure 6, B and C). Phosphorylation of Y507 by c-SRC tyrosine kinase (CSK) maintains LYN in an inactive conformation, which requires an intramolecular interaction between Y507 and the SH2 domain. Upon dephosphorylation, LYN adopts an open conformation in which the activation loop Y396 is phosphorylated and the kinase is activated (17). Thus, we speculated that the LYN D189Y mutation may prevent/hinder the interaction between Y507 and the SH2 domain, thus abrogating LYN inactivation and potentially increasing signaling duration and/or amplitude. However, molecular modeling of autoinhibited SRC did not indicate how the mutation might alter the Y507-mediated conformation of the protein due to its distal location. Additionally, modeling of an activating peptide substrate did not suggest a direct role for D189 in the interaction between the SH2 domain and the activating peptide. Since no active SFK structures that include the SH2 domain are available, determining whether a tyrosine substitution at D189 might alter intermolecular or intramolecular interactions in the active state of
but was not associated with clinical or Ki67 response (32). These investigators also identified a LYNA370T mutation in an AI-resistant tumor and suggested the SFK FYN as a hub of connectivity highly associated with Ki67 status in breast tumors treated with AIs (32). The LYND189Y mutation identified in this study was present in a tumor also harboring an activating PIK3CA T1025A mutation, suggesting a potential link between aberrant LYN and PI3K signaling in endocrine resistance. Consistent with the notion of LYN limiting the full effect of ER and PI3K inhibitors, pharmacological inhibition of the SFKs with dasatinib enhanced the antitumor effects of BKM120 plus fulvestrant against estrogen-deprived parental MCF-7 xenografts and MCF-7 xenografts overexpressing LYNWT. The prototype SFK SRC has previously been shown to play a role in endocrine resistance. Elevated SRC kinase activity promotes invasion and motility in tamoxifen-resistant MCF-7 cells (33), and combined inhibition of SRC and ER impairs breast can-

Figure 9. Pharmacological inhibition of SFKs enhances the effect of BKM120 and fulvestrant against MCF-7/LYNWT xenografts but not MCF-7/LYND189Y xenografts. (A) MCF-7 cells stably transduced with LYNWT or LYND189Y were injected s.c. into athymic ovariectomized mice, as described in the Methods. Mice bearing tumors ≥150 mm³ were randomized and treated as in Figure 8B. Data are presented as log₂ of mean tumor volume (*P < 0.0001 vs. vehicle, +P < 0.05 vs. BKM plus Fulv). (B) Xenografts from A were harvested 4 hours after the last dose of BKM120 or dasatinib and 24 hours after the last dose of fulvestrant. Tumor lysates were analyzed by immunoblot using the indicated antibodies. (C) IHC for Y416 P-SRC. Representative images of tumors from A and quantitative comparison of membrane histoscores, as described in the Methods (H-score; *P < 0.05). Scale bars: 200 μm.
cer growth in vitro and in vivo (33–37). Although LYN has been shown to be important in triple-negative breast cancer (3), to our knowledge, we are the first to identify a possible role for LYN in ER+ breast cancers resistant to estrogen deprivation.

Knockdown of CSK, a negative regulator of SRC, has been shown to suppress fulvestrant-induced proteasomal degradation of ER (38). Since phosphorylation by CSK inhibits SFK activity (17), these data suggest the converse, that is, that derepressed SFKs, as a result of loss of CSK or activating mutations in LYN, block fulvestrant-induced downregulation of ER and growth inhibition (Figure 4). Consistent with this hypothesis, treatment with dasatinib restored the effect of fulvestrant on ER levels (Supplemental Figure 5) and increased the growth inhibitory effects of fulvestrant (Figure 8A). We have shown previously that treatment with BKM120 and fulvestrant induces regression of estrogen-deprived MCF-7 xenografts (21). We expanded upon this by showing that treatment with the combination of dasatinib, BKM120, and fulvestrant was more effective at suppressing ER+/PIK3CA mutant MCF-7 xenograft growth compared with BKM120 and fulvestrant (Figure 8B). LYNΔ^DYQ shows hyperactivate LYN (Figure 6D), diminishing the effectiveness of dasatinib in vivo (Figure 9A). We postulate that resistance to dasatinib in LYN mutant xenografts is due to an inability of LYNΔ^DYQ to achieve an inactive conformation (Figure 6, B and E, and Figure 7A), without changing the conformation and/or affinity of the ATP-binding site. Thus, higher doses may restore the “competitiveness” of dasatinib for the ATP-binding site. Given the lack of detectable inhibition of Y416 P-SRC by dasatinib in MCF-7/LYNΔ^DYQ tumors, we speculate that higher doses would be needed to achieve adequate target inhibition LYN in vivo. These data also suggest that SFKs compensate for the combined effect of PI3K and ER inhibitors against ER+ breast cancers. Phase I and II clinical studies are currently investigating dasatinib in combination with letrozole or fulvestrant (plus an IGF-IR antibody) in patients with ER+ breast cancer (39).

In summary, ER+ breast cancers harbor multiple molecular alterations capable of mediating hormone-independent growth. We have identified LYN mutations as a potential mechanism of escape from estrogen deprivation in ER+ breast cancers. These results support (a) further sequencing studies of LYN in ER+ tumors refractory to endocrine therapy and (b) the need to develop potent SFK inhibitors, which, in combination with PI3K and ER inhibitors, may be an effective treatment for endocrine-resistant breast cancer.

Methods

**Deep kinome sequencing.** Sequencing was performed on DNA extracted from 4 FFPE surgically removed breast tumors that retained a high Ki67 score after 2 weeks of therapy with letrozole (clinical trial NCT00651976). Patient-matched normal adjacent FFPE tissue was also sequenced. Genomic DNA isolated from these tumors and from adjacent normal breast tissue was fragmented into 200-bp sections using a Covaris E210, followed by purification using AMPure XP beads (Agencourt). Library preparation and capture was performed via specific methods set by the Agilent SureSelect Human Kinome Kit (Agilent); libraries were normalized to 147 ng/μl prior to capture. Enrichment of kinase regions was performed by adding 500 ng of prepared library along with hybridization buffer, blocking mix, and capture library provided with the kit. Post-capture samples were normalized to 2 nM and then pooled in a 6-plex; clustering was performed on a cBot (Illumina) using the TruSeq PE Cluster Kit v2 (Illumina) and samples were loaded onto an Illumina GAIIx for sequencing, using the TruSeq SBS Kit v4 GA (Illumina) for 72-bp PE sequencing.

**RPPAs.** Lysates were prepared from frozen core biopsies of the surgical specimens and analyzed by RPPA as described previously (20, 40, 41). FFPE tumor sections before and after treatment were used for Ki67 IHC (DAKO no. M7240) and scored as described previously (12). siRNA screen. MCF-7/LTED cells were transfected with the Dramacon RTF Protein Kinase siRNA library as described previously (21, 42).

**Cell growth assays.** Cells with or without dasatinib (LC Laboratories), AZD0530 (Selleck Chemicals), BKM120 (Selleck Chemicals), or fulvestrant (ICI 182780, Tocris Biosciences) were seeded in 10% dextran-coated charcoal-treated FBS (DCC-FBS) for proliferation in 2D growth assay and counted or fixed/stained with crystal violet as described previously (20). For siRNA experiments, cells in 100-mm dishes were transfected with a siRNA targeting LYN (Hs LYN 13, Qiagen) or a nonsilencing control using HiPerfect Transfection Reagent according to the manufacturer’s protocol (Qiagen). The next day, cells were reseeded in 10% DCC-FBS for proliferation assays and immunoblot analyses (20). 3D growth assays were conducted in growth factor–reduced Matrigel (BD Biosciences) as described previously (43) with 10% DCC-FBS–containing media. Phase-contrast pictures were taken using an Olympus CK40 microscope and Gelcount scanning.

**Immunoblot analysis and immunoprecipitation.** Cell lysates were subjected to immunoblot analysis as described previously (20). Immunoprecipitation was performed using Dynal protein G beads (Invitrogen) and V5 (Invitrogen) or 4G10 pTyr (Millipore) antibodies (44). In vitro kinase assays were performed in immune complexes precipitated with a V5 antibody as described previously (45).

**Apoptosis assay.** Cells (2 × 10^5 cells per well) were plated in white 96-well plates, treated overnight in serum-free media with or without BKM120 or fulvestrant, and then subjected to the Caspase-Glo 3/7 Assay according to the manufacturer’s protocol (Promega).

**Xenograft studies.** Female ovariectomized athymic mice were implanted s.c. with a 14-day-release 17β-estradiol pellet (0.17 mg) and 10^7 MCF-7 parental cells or stably transfected with LYNWT or LYN^E159K. After 4 weeks, mice bearing tumors ≥150 mm^3 were randomly assigned to treatment with vehicle (80 mM sodium citrate buffer, pH 3), dasatinib (15 mg/kg/d, per os [p.o.]), or BKM120 (30 mg/kg/d, p.o.) and fulvestrant (5 mg/wk, s.c.), or BKM120, fulvestrant, and dasatinib. Tumor diameters were measured using calipers twice per week, and volume in mm^3 was calculated with the formula: volume = width^2 × length/2. Tumors were harvested and snap frozen in liquid N_2 or fixed in 10% formalin prior to paraffin embedding for IHC.

**Statistics.** In cell proliferation assays, significant differences were determined by repeated-measures ANOVA with Bonferroni post-hoc test, except as noted below. Paired 2-tailed t tests were used to determine significant differences in siRNA proliferation assays, Caspase-Glo 3/7 assay, real-time qPCR assays, and IHC histoscores. A P value of less than 0.05 was considered statistically significant. Bar graphs show mean ± SD, unless otherwise stated in the figure legends.

**Study approval.** Breast tumor blocks were from a cohort of patients who had consented to the use of any deidentified tissues for research purposes under the auspices of an IRB-approved protocol (Vanderbilt IRB no. NCT00651976). Inclusion criteria for this study were...
postmenopausal women with newly diagnosed ER'/HER2 operable breast cancer treated with letrozole (2.5 mg/d) for 10 to 21 days prior to surgery. Tumor cell proliferation was assessed by Ki67 IHC in a core biopsy before treatment and in the surgical specimen (after treatment). Experiments with mice were approved by the Vanderbilt Institutional Animal Care and Use Committee, protocol no. M/10/111.

Additional details are provided in the Supplemental Methods.

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