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Chapter 6

ERa Cofactor phosphorylation

SRC3 phosphorylation at Serine 543 is a positive independent prognostic factor in ER positive breast cancer

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Abstract

Purpose: The steroid receptor coactivator SRC3 is essential for the transcriptional activity of estrogen receptor α (ER α). SRC3 is sufficient to cause mammary tumorigenesis, and has also been implicated in endocrine resistance. SRC3 is posttranslationally modified by phosphorylation, but these events have not been investigated with regard to functionality or disease association. Here, we investigate the spatial selectivity of SRC3-pS543/DNA binding over the human genome and its expression in primary human breast cancer in relation with outcome.

Experimental Design: Chromatin immunoprecipitation, coupled with sequencing, was used to determine the chromatin binding patterns of SRC3-pS543 in the breast cancer cell line MCF7 and two untreated primary breast cancers. IHC was used to assess the expression of SRC3 and SRC3-pS543 in 1,650 primary breast cancers. The relationship between the expression of SRC3 and SRC3-pS543, disease-free survival (DFS), and breast cancer specific survival (BCSS) was assessed.

Results: Although total SRC3 is selectively found at enhancer regions, SRC3pS543 is recruited to promoters of ER α responsive genes, both in the MCF7 cell line and primary breast tumor specimens. SRC3-pS543 was associated with both improved DFS (P = 0.003) and BCSS (P = 0.001) in tamoxifen untreated high-risk patients, such a correlation was not seen in tamoxifen-treated cases, the interaction was statistically significant (P = 0.001). Multivariate analysis showed SRC3-pS543 to be an independent prognostic factor.

Conclusions: Phosphorylation of SRC3 at S543 affects its genomic interactions on a genome-wide level, where SRC3-pS543 is selectively recruited to promoters of ER α -responsive genes. SRC3-pS543 is a prognostic marker, and a predictive marker of response to endocrine therapy. Clin Cancer Res; 22(2); 479–91. ©2015 AACR.

Translational Relevance: SRC3 is an essential coactivator for the transcriptional activity of ER α and is required for estrogen-dependent cell proliferation. Analogous to ER α , SRC3 is typically found at enhancer regions involved in long-range regulation of hormonal-responsive genes. In this study, we establish the genome-wide chromatin binding preferences for activated SRC3 (SRC3-pS543) in ER α -positive breast cancer, as well as the association

between SRC3-pS543 expression and clinicopathologic features and clinical outcomes in 1,650 primary breast cancers. Our findings reveal that while ER α and total SRC3 are predominately found in distal enhancers and introns, SRC3-pS543 is predominately located at promoters of genes. SRC3-pS543 was associated with favorable clinicopathologic features and was found to be an independent prognostic factor as well as a predictive marker with regard to tamoxifen treatment. These observations illustrate that high expression of SRC3-pS543 can potentially identify a population of early ER-positive breast cancer with a good clinical outcome without receiving adjuvant therapy.

Introduction

Estrogen receptor α (ER α) is a key transcription factor in normal breast development and plays a central role in the pathogenesis of approximately 70% of all breast cancers. Consequently, ERa is recognized as the key therapeutic target in this group of patients (1). The p160 steroid receptor coactivator (SRC) family member, SRC3 (also known as amplified in breast cancer 1; AIB1) is an essential coactivator for the transcriptional activity of ERa, which recruits SRC3 to chromatin (2). SRC3 is required for estrogen-driven proliferation of MCF7 breast cancer cells (3) and is critical for normal mammary development (3). Overexpression of SRC3 can induce mammary tumorigenesis in mice (4), while its absence protects against breast cancer development (5, 6). SRC3 was originally identified as a gene mapping to a region of chromosome 20 (20q12–13), which is frequently amplified in breast cancer and was shown to be amplified in 10% of breast cancers (7). SRC3 mRNA levels are significantly higher in breast cancer as compared with normal mammary tissue (8, 9) and SRC3 protein levels are elevated in 16% to 83% (depending on the study) of human breast tumors (10).

SRC3 expression has been associated with decreased risk of relapse in patients with breast cancer who did not receive adjuvant endocrine therapy (11). Conversely, in tamoxifen-treated patients SRC3 expression was associated with poor outcome and increased risk of relapse (11). Others have found no such association with tamoxifen treatment, although an increased risk was seen with high SRC3 when expressed alongside HER1, HER2, or HER3 (12). In a group of women of whom 82% were postmenopausal, SRC3 expression has been associated with early recurrence on tamoxifen treatment (13), although high SRC3 in premenopausal women was an independent predictive factor of improved response to tamoxifen (14). ER α -associated SRC3 is typically found at enhancer regions, where expression of genes proximal to shared ER α /SRC3 chromatin binding events correlates with poor outcome of ER α -positive patients with breast cancer treated with tamoxifen (15). This feature was only found for genes where SRC3 was selectively enriched as compared with the other two p160 family members, SRC1 and SRC2.

Phosphorylation of SRC3 is required for it to function as a potent transcriptional activator, and SRC3 possesses multiple phosphorylation sites that may be functionally involved in this process (16, 17). Estradiol treatment increases SRC3 phosphorylation, but the kinase responsible for this remains elusive. However, a number of kinases are known that induce phosphorylation at these residues of SRC3 (16, 17). In addition, dephosphorylation of serine-101 and -102 regulates SRC3 protein stability by preventing proteas-ome-dependent turnover (18). Of note, the C-terminal domain of SRC3 has weak histone acetyltransferase activity raising the possibility that it may have a role in chromatin remodeling (10).

Previous studies have shown a requirement of SRC3 phosphorylation for SRC3 activity, including phosphorylation of SRC3 at serine residue 543 (SRC3-pS543 (16)). Furthermore, the region around S543 is evolutionarily conserved in mammals, birds, and reptiles, although no sequence homology around S543 is found in SRC1 or SRC2 (19). In addition, thus far no studies explored the genomic consequences and clinical significance of SRC3 phosphorylation. Given all the above, we decided to investigate the chromatin associations of SRC3 in vitro and in vivo when phosphorylated at serine 543 (SRC3-pS543) together with its clinical significance in ER α -positive breast cancer. To separate the prognostic from the predictive potential of SRC3 phosphorylation, tamoxifen-treated patients were compared with patients who did not receive any adjuvant endocrine therapy.

Results

Specificity of SRC3-pS543 antibody

An antibody for SRC3-pS543 was generated, which recognized wild-type phosphorylated SRC3, but not S543-mutated SRC3 (S543A; **Fig. 1A**). The SRC3-pS543 antibody detected SRC3 in immunoprecipitates from MCF7 cell lysates (**Fig. 1A and B**; full size blot Supplementary Figs. S1 and S2), where signal was lost after the phosphatase treatment (**Fig. 1B**). Consistent with a previous report (17), we find estrogen treatment to increase S543 phopshorylation in a dose-dependent manner, further validating our SRC3-pS543 antibody (**Fig. 1C**). To ensure the specificity of the SRC3-pS543 antibody

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Figure 1: Validation of phosphostate-specific antibody.

(A) CHO cells were transfected with pCDNA3, Flag-tagged wild-type (WT) SRC3 or a Flag-tagged SRC3 mutant in which serine at residue 543 has been substituted by alanine. MCF7 cells were used as control. The lysates were immunoprecipitated with SRC-3 antibody. Immunoblotting was performed for SRC-3 and phospho-S543. Note that the Flag-tagged SRC3 migrates slower on Western blot as compared with endogenous SRC3. (B) MCF7 cell lysate was immunoprecipated with SRC-3 antibody. The beads were then divided into two halves, one untreated and the other treated with λ -phosphatase. Immunoblotting was performed for SRC-3 and phosphorylation state-specific antibody. (C) CHO cells were grown in DMEM with dextran-coated charcoal stripped FCS for 72 hours and then transfected with vector (pCDNA3) or SRC-3 for 24 hours. Cells were then treated with vehicle or increasing concentrations of 17 β -estradiol (E2) for 1 hour before making lysates for immunoprecipitation. Western blotting was carried out with the antibodies indicated. MCF7 cells were used as control. (D) Representative tumor sections following preincubation with phospho-peptide and nonphospho-peptide and phosphatase treatment prior to staining with SRC3-pS543 in breast cancers with negative, low/medium, and high expression of SRC3-pS543.

by IHC, the antibody was incubated with either peptide containing the phospho-moiety or the nonphospho-peptide. Subsequently, the preincubated antibody (bound to nonphospho-peptide and phospho-peptide) was tested on four tumors with different levels of signal for SRC3-pS543 (negative, low/ medium, and high; **Fig. 1D**). In addition, phosphatase treatment was applied. Although the phospho-peptide strongly decreased pS543-SRC3 signal, this was not observed with the nonphospho-peptide. In addition, phosphatase treatment strongly diminished pS543-SRC3 signal in all tested specimens (**Fig. 1D**).

ChIP-seq analysis shows a genomic preference of SRC3-pS543 to bind promoter regions

Next, we determined the genome-wide distributions of this SRC3-pS543 phosphorylation as compared with total SRC3 and ERa in asynchronously proliferating luminal breast cancer cells MCF7 as exemplified for wellknown ERa binding sites proximal to the XBP-1 locus (Fig. 2A). For ERa genomic locations, data from ref. 24 were used, although total SRC3 data were from ref. 15. In total, 47,214 ERa binding sites were detected, of which 15,329 were also bound by SRC3 (ref. 15; Fig. 2B). Between ERa, SRC3, and SRC3-pS543, 2,365 peaks were shared (for raw data visualizations, see Fig. 2C). Not all SRC3 binding events were phosphorylated and 2,795 peaks for SRC3-pS543 were apparently not shared with total SRC3. However, there was weak total SRC3 signal observed at these positions, which did not reach the detection threshold for peak calling (Supplementary Fig. S3A and S3B). Based on these results, we conclude that differential binding patterns between SRC3-pS543 and total SRC3 should be considered as "enriched" rather than "unique." P160 family members share substantial sequence homology. Even though the sequence surrounding S543 is not conserved in SRC1 and SRC2, the SRC3-pS543 antibody might still detect the other two p160 members. Therefore, we determined potential overlap of SRC3-pS543 enriched sites with SRC1 and SRC2 in MCF7 cells (15). The overlap of SRC3-pS543 enriched sites with SRC1 and SRC2 was limited, rendering it unlikely that the SRC3-pS543 enriched sites can be explained merely by cross-reactivity with one of the other p160s (Supplementary Fig. S4). Differential chromatin binding between SRC3-pS543 and total SRC3 could be validated by qPCR analyses on asynchronous MCF7 cells (Supplementary Fig. S5A), using four primer sets that were designed for each subset of binding sites, that is, SRC3-pS543enriched, total SRC3-enriched, and shared sites (Supplementary Table S1 for

SRC3 enriched -pS543 enriched All shared ERa enriched ERa and SRC3 ERc and pS543 enriched _____ SRC3 and pS543 enriched _____ ш C ⊳ SRC3 pS543 SRC3 enriched ERo 5'UTR 3'UTR ERα Coding exon
Intron
Distal intergenic SRC3 pS543 П Β Genes with SRC3-pS543 at promoter (%) SRC3-pS543 735 16] 00 165 ER regulated SRC3 P < 0.0001 12,450 514 All Refseq ERa 31,595 ERa/SRC3-pS543 enriched ERa/SRC3-pS543 shared A PARTY AND A PART

Figure 2: SRC3-pS543, SRC3, and ERa chromatin binding patterns in MCF7 cells. (A) Genome browser snapshot of ChIP-seq samples for ERa (red), SRC3 (blue), and SRC3-pS543 (green) in proliferating MCF7 cells. Y bar shows tag count. Genomic

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coordinates are indicated. (B) Venn diagram of ERa (red), SRC3 (blue), and SRC3pS543 (green). Numbers of shared and unique peaks are shown. (C) Heatmap visualization of ERa (red), SRC3 (blue), and SRC3-pS543 (green). (D) Motif analyses of ERa/SRC3-pS543 shared and SRC3-pS543 enriched binding sites. Z-score for each motif is shown in a MRP. Motif enrichment is shown for chromatin binding sites shared between ERa and SRC3-pS543 (red) or selectively enriched for SRC3-pS543 (green). The radial data points represent the absolute value of Z-score. (E) Genomic distributions of ERa, SRC3, and SRC3-pS543 sites. (F) Percentage of ERa-regulated genes with a SRC3-pS543 binding event at the promoter, compared with all refseq genes. $\chi 2$ test: P < 0.0001.

primer sequences and Supplementary Fig. S5B for genome browser snapshots of SRC3 and SRC3-pS543 ChIP-seq signal at these sites). To assess hormone dependency of SRC3-pS543 chromatin interactions, SRC3-pS543 ChIP-qP-CR was performed on cells that were hormone deprived for 3 days, and subsequently treated for 3 hours with vehicle, E2, or tamoxifen (Supplementary Fig. S6). SRC3-pS543/chromatin interactions at SRC3-pS543-enriched sites (that were not bound by $ER\alpha$) were hormone independent and not affected by ligand. For SRC3-pS543 sites shared with total SRC3, no chromatin binding was observed in the absence of hormone and in tamoxifen-treated cells, but this interaction was actively induced by E2 treatment. Phosphatase treatment abrogated SRC3-pS543 enrichment at these sites (Supplementary Fig. S6). Genomic distributions of ERa and SRC3 were comparable as reported by others (ref. 29; Supplementary Fig. S7). To further illustrate functional interplay between ERa and SRC3-pS543, ERa was depleted from MCF7 cells using Fulvestrant (Supplementary Fig. S6). SRC3-pS543/SRC3 shared, but not SRC3-pS543-enriched chromatin interactions were dependent on ERα levels, as shown by Fulvestrant-mediated ERa degradation (Supplementary Fig. S8). In order to identify potential selectivity of transcription factor usage, motif analyses were performed on the sites that were either or not shared between SRC3-pS543 and ERa, and visualized in a MRP (Fig. 2D). For the shared events, motif enrichment was found for classical luminal breast cancer-selective transcription factors, including ESR1 and forkhead motifs. For SRC3-pS543-enriched binding events, no motifs for ESR1 or FOXA1 were enriched, but a clear enrichment was observed for ELK, ELF, and ETV motifs (see also Supplementary Table S4 for the total list).

Differential motif enrichment could form the basis of differential gene expression programs. Therefore, GO pathway enrichment was assessed

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for genes with an SRC3/chromatin binding event proximal (<20 kb) to their transcription start sites. GO terms highly varied for the differential SRC3 subclasses (Supplementary Fig. S9), with 1 out of 80 GO terms for SRC3 enriched sites ("biologic process") shared with the other two subsets. Interestingly, the majority of SRC3-pS543/SRC3 shared peak GO terms, 28 out of 37, were overlapping with those found for the SRC3-pS543-enriched sites, including "metabolic process," "gene expression," and "translation." For a complete list of all GO terms, see Supplementary Table S5.

Because the SRC3-pS543 chromatin binding events only recapitulated a proportion of the total SRC3 binding events, genomic distributions could deviate as well. As described before (15, 30), the majority of ER α and SRC3 binding events are found within enhancers and introns (Fig. 2E). In contrast, SRC3-pS543 binding was clearly enriched at promoter regions and 5'UTR, with promoter-bound SRC3-pS543 colocalizing with SRC3, ERa, and histone modifications indicative for active gene transcription (H3K4me2, H3K4me3, and H3K27Ac; Supplementary Fig. S10). As expected, SRC3 and ERa signal was found centered on the SRC3-pS543 signal. Because transcription factors bind accessible chromatin, histone marks indicative for active promoters (H3K4me3 and H3K27Ac) are found around the SRC3-pS543 sites instead of directly centered on them. Interestingly, this was not the case for H3K4me2, which was clearly enriched on the SRC3-pS543 peak. The enhancer-selective histone mark H3K4me1 was not enriched at SRC3-pS543 promoters. Even though less pronounced than the total of SRC3-pS543 peaks, the sites shared between ERα and SRC3-pS543 were also more promoter enriched (13.9%) as compared with the ER α /SRC3 shared regions (4.9%). Furthermore, SRC3pS543 signal was highly enriched at promoters of E2-regulated genes (31) as compared with all Refseq genes (P < 0.0001; Fig. 2F).

SRC3-pS543 and ERα chromatin binding events in primary tumor tissue Next, we determined phosphorylated SRC3/chromatin interactions in primary tumors (Fig. 3). ChIP-seq was performed on two independent ER+/PR+/ HER2– breast tumor specimens as described (21, 24), and directly compared with the MCF7 cell line data. Peak intensities between the two tumors and the MCF7 sample were comparable, and peaks are well conserved (Fig. 3A). Between the two tumors and MCF7 cells, 2,670 binding events of SRC3-pS543 were shared (Fig. 3B, visualized in a raw heatmap in Fig. 3C). Furthermore, peaks were also found selectively enriched for only MCF7 (2,523 sites), tumor #1 (1,816 sites), or tumor #2 (4,631 sites). The SRC3-pS543 binding

sites from all samples were annotated for their genomic distributions, and again enrichment for promoters and 5'UTR regions was found (Fig. 3D). This was even more strikingly observed for binding events shared by all three samples, with 52% of all sites being localized to promoters and 20% at 5'UTR. For tumor #1, the lysate was divided in two prior to immunoprecipitation, and the second half of the lysate was used for ERa ChIP-seq, enabling a direct comparison of SRC3-pS543 and ERa binding patterns within the same tumor (Fig. 3E). Between ERα and SRC3-pS543, 1,727 binding events were shared, although 15,050 binding events were only found for ERa and 5,537 binding sites were only found for SRC3-pS543 (Fig. 3F). The shared and enriched binding patterns were not dictated by different thresholds in peak calling algorithms, as shown by heatmap analyses (Fig. 3G). The genomic distributions of ER α in the tumor sample closely resemble what was found in the cell line (Fig. 2E) and previously reported in breast tumor tissue (21, 24), with approximately 5% of ERa binding events being found at promoters, with the vast majority of sites located at distal enhancers and introns (Fig. 3H). The SRC3-pS543 sites (either or not shared with ER α), were highly enriched at promoters and 5'UTR regions.

Motif analyses for the ER α binding events in the tumor specimen revealed enrichment in ER α , FOXA1, and AP-2 motifs, as expected (refs. 30, 32; **Fig. 31**). Enrichment for these specific motifs was found both for ER α -enriched sites as well as for sites shared between ER α and SRC3-pS543. Furthermore, practically all motifs found at SRC3-pS543–enriched sites were also found at the regions shared by ER α and SRC3-pS543 bounds, including motifs for ELK, ETV, and ETS factors as were observed in the MCF7 cells (see Supplementary Table S6 for the total list).

Association of total SRC3 and SRC3-pS543 with clinicopathologic features

Total SRC3 and SRC3-pS543 expression was evaluated in 1,650 breast carcinomas, with protein expression predominately nuclear (Supplementary Fig. S11). The total SRC3 mean histoscore was 157 (interquartile range, 0–300) and for SRC3-pS543 mean histoscore was 31 (interquartile range, 0–300). For subsequent analysis, total SRC3 expression was categorized as high (above mean, H-score > 157) or low (below mean, H-score < 157) and for SRC3pS543 high (above mean, H-score > 31) or low (below mean, H-score \leq 30; Supplementary Table S3). A scatterplot analysis of total SRC3 versus SRC3pS543 in these tumors shows that the relative levels between total SRC3 ver-

D Q Þ SRC3-pS543 ERa unique Shared SRC3 pS543 SRC3 fumor #2 umor #1 ER 5 kb SRC3 pS543 chr21:30,357,497-30,451,412 has side and the ERa unique т - dan ш SRC3 ERa ER*ul* SRC3-pS543 shared 54 Tumor #1 ω SRC3-pS543 SRC3 pS543 unique 1,816 Distal intergenie 2,509 Promoter Downstream 5'UTR 3'UTR Coding exon Intron 2,670 2,523 MCF chr16:23,426,822-23,470,813 4,631 Imor #2 A CONTRACT OF CONTRACT C MCF7 MOF7 and į SRC3 П 5,537 KINK 1,727 MCF7 15,050 ERa enriched SRC3-pS543 enriched ERa/SRC3-pS543 shared Tumor 1 Tumor 2

Figure 3: SRC3-pS543 binding events in MCF7 cells and breast tumors. (A) Genome browser snapshot, ChIP-seq for SRC3-pS543 on MCF7 cells (green),

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tumor sample #1 (blue), and tumor sample #2 (purple). Y bar shows tag count. Genomic coordinates are indicated. (B) Venn diagram of SRC3-S543P Chlp-seq for *MCF7 cells (green), tumor sample #1 (blue), and tumor sample #2 (purple). Number* of shared and unique sites is illustrated. (C) Heatmap visualization of SRC3-S543P ChIp-seq for MCF7 cells (green), tumor sample #1 (blue), and tumor sample #2 (purple). Number of shared and enriched sites is illustrated. (D) Genomic distributions of shared and unique SRC3-pS543 binding events for MCF7 cells, tumor sample #1, and tumor sample #2. (E) Genome browser snapshot, ChIP-seq on a primary breast tumor for ERa (red) and SRC3-pS543 (blue). (F) Venn diagram of ERa (red) and SRC3-pS543 (blue) binding events. (G) Heatmap analysis, depicting all shared and enriched binding events for $ER\alpha$ (red) and SRC3-pS543 (blue). Shown are all binding events with a window of 5 kb around the binding site. (H) Genomic distributions of the shared and enriched binding events of $ER\alpha$ and SRC3-pS543. (I) Motif analyses of the shared and enriched binding events of ERa and SRC3-pS543. Z-score for each motif is shown in a MRP. Motif enrichment is shown for chromatin binding sites enriched for ER α (red), SRC3-pS543 (green), or shared between ER α and SRC3-pS543 (blue). The radial data points represent the absolute value of Z-score.

sus SRC3-pS543 were strongly variable between tumors. Most importantly, a large proportion of tumors were identified that were positive for total SRC3, but negative for SRC3-pS543. Furthermore, no tumors could be identified that were positive for the phospho-SRC3 but negative for total SRC3, again highlighting specificity of the antibody (Supplementary Fig. S12).

Although no association on IHC between total SRC3 and ER α was found, there was a strong association for SRC3-pS543 with ER α (P = < 0.0001) and high expression of ER α -associated proteins (progesterone receptor; P = < 0.0001 and Bcl2; P = < 0.0001). Total SRC3 expression was significantly associated with HER2 (P = < 0.0001) and HER3 (P = 0.023), and correlated the absence of basal-like (P = 0.021) and triple negative phenotypes (P = 0.005; Table 1). SRC3-pS543 was significantly associated with well-differentiated (P ≤ 0.0001), low proliferative (P ≤ 0.0001) tumors, as well as androgen receptor (P ≤ 0.0001) expression. In addition, it was associated with the presence of p53 (P = 0.025) and the absence of MDM4 (P < 0.001) and MDM2 (P < 0.001), as well as the absence of SRC3-pS543 was significantly associated with the overexpression of HER4 (P < 0.0001), the presence of lympho-vascular invasion (P ≤ 0.0001), and loss of expression of the key DNA repair proteins, including BRCA1 (P < 0.0001), ATM (P < 0.0001), and TOP2A (P < 0.0001; Table 1). Apart from being phosphorylated by estrogen treatment, SRC3-S543 is also phosphorylated by JNK and p38 MAPK (16). In line with these previous observations, we find a significant correlation between the phosphorylated JNK and p38 MAPK with SRC3-pS543 (P < 0.0001; Table 1). Additional tumor biomarkers and their correlation with SRC3 and SRC3-pS543 are shown in Supplementary Table S7.

Association of total SRC3 and SRC3-pS543 expression with outcome

With regard to outcome, no association of total SRC3 with outcome was observed in the whole cohort (Supplementary Fig. S11A), the ER α -positive group in total (Supplementary Fig. S11B), or on the basis of tamoxifen treatment (Supplementary Fig. S11D–S11F). Furthermore, no association of total SRC3 with outcome was observed in the ER α -negative cohort (Supplementary Fig. S11C). However, expression of SRC3-pS543 in the whole cohort was associated with significantly longer disease-free survival (DFS) (P < 0.00001) and breast cancer specific survival (BCSS) (P = 0.0001; Fig. 4A). SRC3-pS543 expression in the ER α -positive cohort correlated with a longer DFS (P = 0.003) and BCSS (P = 0.001; Fig. 4B), although SRC3-pS543 did not associate with survival in ER α -negative breast cancers (Fig. 4C). Furthermore, in low-risk ER α -positive cancers, no correlation of SRC3-pS543 with outcome was found (Fig. 4D).

Importantly, SRC3-pS543 was associated with a longer DFS (P =0.035) and BCSS (P = 0.005) in ER-positive high-risk breast cancer not treated with tamoxifen (Fig. 4E). In high-risk patients who did receive tamoxifen, SRC3-pS543 did not associate with a longer DFS (P = 0.57) or BCSS (P =0.21; Fig. 4F). For ER-positive high-risk patients with low SRC3-pS543, exposure to tamoxifen reduced the risk of death from breast cancer [HR (95% CI) = 0.68 (0.49-0.94), P = 0.019] by 32% (Fig. 4G), whereas for those with high pS543, there was no effect (Fig. 4H), the interaction was statistically significant (P = 0.001). Similarly, exposure to tamoxifen reduced risk of recurrence by 42% [HR (95% CI) = 0.58 (0.44–0.77), P = 0.0001] in ER+ high-risk patients with low pS543, whereas no effect on those with high pS543 patients, the interaction was statistically significant (P = 0.000.1). Tamoxifen-treated patients with HER2-positive tumors that expressed high levels of SRC3 had a significantly shorter disease-free and BCSS. This was not observed for the patients who did not receive endocrine treatment (Supplementary Fig. S14). SRC3-pS543 was confirmed as an independent prognostic factor in breast cancers after adjustment for endocrine therapy and other validated prognostic



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Figure 4: DFS and BCSS in patients with breast cancer based on SRC3-pS543 protein expression.

(A) Entire Nottingham cohort. (B) $ER\alpha$ -positive cohort. (C) $ER\alpha$ -negative cohort NPI > 3.4. (D) $ER\alpha$ -positive patients NPI < 3.4 who did not receive tamoxifen. (E) $ER\alpha$ -positive patients $NPI \ge 3.4$ who did not receive tamoxifen. (F) $ER\alpha$ -positive patients $NPI \ge 3.4$ who received tamoxifen. (G) BCSS based on SRC3-pS543 protein expression and tamoxifen treatment in the $ER\alpha$ -positive high-risk group with low pSRC3-pS543. (H) BCSS based on SRC3-pS543 protein expression and tamoxifen treatment in the ER\alpha-positive high-risk group with high pSRC3-pS543.

factors; and absence of SRC3-pS543 signal correlated with recurrence (HR = 1.4, CI 95%, 1.1-2.5) and death from breast cancer (HR = 1.6, CI 95%, 1.1-2.5; Table 2).

Discussion

Coactivators for ER α have been the focus of many clinical and cell biologic studies. SRC3 is of particular interest, because the SRC3 gene is frequently amplified in breast cancer (7), and its expression was found to correlate with HER2 expression levels and poor outcome (11). SRC3 is a member of the p160 coregulator protein family (along with SRC1 and SRC2), which all have shared and preferred genome-wide chromatin binding patterns, which are strongly induced by ER α activation (15).

Here, we focus on one distinct phosphorylation event on SRC3, Serine 543 phosphorylation. This phosphorylation has spatial selectivity on a genomic scale, where chromatin binding was selectively enriched at active promoter regions. This genomic preference at promoters is on contrast to total SRC3, which is mainly found at enhancers. It is likely that SRC3 and SRC3-pS543 antibodies have different sensitivities, and the level of overlap detected between these two signals is possibly an underestimate of the actual proportion of SRC3 that is phosphorylated. Relative expression levels and antibody sensitivity are likely to alter the threshold of detection in ChIP-seq experiments, forming a major source for false-negative signal. Because the strongest total SRC3 sites are enhancer-selective, whereas the strongest SRCpS543 were mostly enriched at promoters, this genomic selectivity of phosphorylated SRC3 to promoter regions cannot be merely explained by relative expression levels or antibody sensitivity.

Because ER α typically binds enhancers (30), the determination of which ER α binding events are functionally orchestrating what genes has re-

mained a challenge in the field. Because most ER α binding events are observed within a 20-kb window from the transcription start sites of responsive genes (33), this spacing around the promoter is typically applied to provide an indication of functionality. All p160 coactivators, including SRC3, show comparable genomic features as compared with ER α and are typically found at enhancers as well (15). Because the SRC3-pS543 phosphorylation is promoter-enriched, commonly shared with ER α and induced by estradiol, this phosphorylation event could be considered as a novel ER α functionality probe, highlighting ER α complexes that are actively involved in transcriptional events at gene promoters.

Smaller previous studies on SRC3 expression and its correlations with ERα and members of the EGFR/HER family have reported varying results (11–13). We find no correlation of SRC3 with ER α but do find a correlation with HER2 and HER3. However, SRC3-pS543 does significantly correlate with ER α , potentially reflecting the role of E2/ER α in phosphorylation of SRC3. Consistent with these results was the observed correlation of SRC3pS543 with known estrogen-regulated genes, including PR. Although previous studies have explored the influence of SRC3 expression on patient outcome as well how it may influence endocrine therapy (11-13, 34-39), the current study is the largest and most well-characterized series to date, with a long median follow-up. We find no statistically significant association between SRC3 levels and DFS and BCSS in patients with ERa-positive breast cancer treated with tamoxifen, consistent with other reports (12, 13). Of note, in those women with $ER\alpha$ -positive tumors not treated with tamoxifen, there is a trend to an improved outcome and survival for SRC3-positive tumors, analogous to the first report relating SRC3 and response to endocrine therapy (11). The poor outcome seen in HER2-positive/SRC3-high tumors for patients treated with tamoxifen is consistent with previously published data (11, 12). One previous study on ER α -negative tumors, which utilized automated quantitative analysis in a subgroup of 133 patients, found SRC3 associated with a poorer overall survival (38). However, in the current study of 377 ERα-negative tumors no correlations with recurrence or outcome were found for SRC3. Of note, we show a significant correlation of p-JNK and p-p38 with SRC3-pS543. These kinases have previously been described to phosphorylate SRC3-S543 (16), and activated (phosphorylated) JNK has previously be linked to endocrine therapy resistance (37) (40), as well as poor outcome in breast cancer (41). The current study linking pJNK with high SRC3-pS543 and a good prognosis group is not comparable with these previous studies for

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technical and methodological reasons. In the previous study, which investigated the expression of pJNK in human breast cancer (39), differing reagents, scoring system, as well as a smaller number of cases (n = 68) was used. Furthermore, 40% of the cases were ER α negative, and in those that were ER α positive no clinicopathologic correlation was presented (39). Although the other studies either investigated the activity of JNK and not its expression in the context of breast cancers exposed to tamoxifen (37) or where levels of pJNK were assessed via immunoblotting in the context of a MCF7 breast cancer xenograft model (38).

Our study is the first to explore the clinical significance of a phosphorylated form of SRC3. SRC3-pS543 correlates with an improved DFS and BCSS for the whole cohort and is a predictor of outcome in ER α -positive breast cancer. Furthermore, in high-risk ER-positive disease it is demonstrated that SRC3-pS543 is predictive of benefit with adjuvant tamoxifen. With no benefit being seen with adjuvant tamoxifen in the presence of high SRC3pS543, although those tumors with low SRC3-pS543 had improved outcomes with tamoxifen treatment. The lack of benefit observed in high SRC3-pS543 tumors treated with tamoxifen indicates that the transcriptional events driven by SRC3-pS543 at gene promoters of ER α responsive result in a phenotype with an excellent clinical outcome, this would be consistent with the clinicopathologic features associated with SRC3-pS543. Although insufficiency or inability to enrich at these promoters as a result of low or absent SRC3-pS543 results in a more aggressive phenotype, which can be abrogated by tamoxifen.

The patient group studied here is a well-characterized historical cohort treated with or without tamoxifen. However, estrogen withdrawal therapy using aromatase inhibitors (AI) is now a standard clinical practice in the adjuvant setting for postmenopausal breast cancer, and correlations of response to AIs with (phosphorylated) SRC3 levels need to be assessed within this setting, particularly because the oncogenicity of SRC3 appears modulated by the hormonal milieu in mouse models (42). Currently, the translational arm of the Intergroup Exemestane Study (so-called Path-IES) is exploring the potential role of SRC3 and pS543 in this context (43).

In summary, we have found a novel biomarker for ER α -positive breast cancer and provide mechanistic insights in its functional involvement in the ER α transcription complex. SRC3-pS543 phospho-specific antibodies could aid in identifying patients with a functional ER α signaling pathway, which predicts for a favorable outcome in the absence of adjuvant therapy and where tamoxifen might better be withheld.

Material and Methods

Cell lines, plasmids and antibodies.

MCF-7 and CHO cells were from the American Type Culture Collection (ATCC), which utilises short tandem repeat profiling with no authentication being performed by authors. All experiments were performed within 10 passages from the original stock. Cell lines were grown in DMEM with 10% FCS and standard antibiotics. Wild type SRC3 and mutant S543A plasmids kindly provided by BW O'Malley (Baylor College of Medicine). The antibodies used were SRC3 (Immunohistochemistry, BD Transduction Laboratories; 611105), SRC3 (ChIP-seq experiments, SC-9119; Santa Cruz), ER α (SC-543; Santa Cruz). A rabbit polyclonal antiserum for SRC3-pS543 was generated by immunising animals with the peptide 537-LLSTLSSPGPKLDN-550, where the residue in bold is phospho-serine (Moravian Biotechnologies, Czech Republic).

(Chromatin) Immunoprecipitations

Chromatin immunoprecipitations were performed according to standard protocols (Schmidt et al., 2009). Primary breast cancer specimens and proliferation MCF-7 breast cancer cells were used. Tissue was cryosectioned in 30 slices of 30 µm, and subsequently defrosted in solution A (50 mM Hepes, 100 mM NaCl, 1mM EDTA, 0.5 mM EGTA) supplemented with 1% formaldehyde. For MCF-7 cells, formaldehyde as added to the culture medium at a final concentration of 1%. After 10 minutes of fixation, 100 mM Glycine as added, after which cells were washed in PBS in presence of Protease Inhibitor (Roche Diagnostics 11836145001) and Phosphatase Inhibitors (Roche Diagnostics: 04906837001). Tissue was homogenized, and nuclei were isolated, washed and sonicated using a Diagenode Bioruptor as described before (Zwart et al., 2013, Jansen et al., 2013). After sonication, the lysate was cleared by centrifugation. For each ChIP 100µl of Dynabeads (Invitrogen) was used, precoupled overnight with 10µg of SRC3 or SRC-pS543 antibody. Immunoprecipitation was performed overnight while rotating at 4 degrees. Subsequently, beads were washed 10 times using RIPA buffer and reverse crosslinked at 65 degrees for 6 hours. Supernatant was removed, cleaned using a phenol/chloroform column and precipitated as described before (Schmidt et al., 2009). For standard Immunoprecipitations, beads were resuspended in sample buff-

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er, boiled at 95°C for 5mins and the samples run onto 8% Polyacrylamide gel for protein analysis for S543P, Input SRC3 and Immunoprecipitated (IP) SRC3. Immunoprecipitations were performed in presence of Protease Inhibitor (Roche Diagnostics 11836145001) and Phosphatase Inhibitors (Roche Diagnostics: 04906837001). When indicated, samples were incubated with Lambda Protein Phosphatase (Millipore; cat # 14-405) for 30 min prior to ChIP. Primer sequences for QPCR analysis are in Supplementary Table S1. For MCF-7 experiments, ~6x107 asynchronous proliferating cells were used, and two independent replicates were performed for each ChIP-seq, where only consensus peaks that were found in both experiments were considered. For tumour specimen ChIP-seq, fresh snap-frozen primary ER α -positive, HER2-negative untreated breast cancers were obtained from the Imperial College Healthcare NHS Trust Tissue Bank which has the requisite ethical

Solexa sequencing and enrichment analysis

approval..

ChIP DNA was amplified as described (Schmidt et al., 2009). Sequences were generated by the Illumina Hiseq 2000 genome analyser (using 50 bp reads), and aligned to the Human Reference Genome (assembly hg19, February 2009). Enriched regions of the genome were identified by comparing the ChIP samples to mixed input using the MACS peak caller (Zhang et al., 2008) version 1.3.7.1. Total read count, percentage of aligned reads and number of peaks for each ChIP-seq is shown in Supplementary Table S2. For histone modification data, publically available MCF-7 ChIP-seq data for H3K4me1 (GEO: GSM588568), H3K4me2 (GEO: GSM822392), H3K4me3 (GEO: GSM945269) and H3K27Ac (GEO: GSM945854) was used. For ERα ChIP-seq data, peaks from (Ross-Innes et al., 2012) were used. ChIP-seq data for SRC1, SRC2 and total SRC3 were from (Zwart et al., 2011b). For heatmap visualizations, Seqminer software was used.

Motif analysis, heatmaps and genomic distributions of binding events

ChIP-seq data snapshots were generated using the Integrative Genome Viewer IGV 2.1 (www.broadinstitute.org/igv/). Motif analyses were performed through the Cistrome (cistrome.org), applying the SeqPos motif tool (He et al., 2010). The genomic distributions of binding sites were analysed using the cis-regulatory element annotation system (CEAS) (Ji et al., 2006). The genes closest to the binding site on both strands were analysed. If the binding region is within a gene, CEAS software indicates whether it is in a 5'UTR, 3'UTR, coding exon, or an intron. Promoter is defined as 3 kb upstream from RefSeq 5' start. If a binding site is >3 kb away from the RefSeq transcription start site, it is considered distal intergenic.

Nottingham Tenovus Primary breast cancer Series

Primary operable breast cancer cases (n = 1,650) from the Nottingham Tenovus Primary Breast Carcinoma Series was utilised as described before (Elston and Ellis, 1991, Ellis et al., 1992), supplementary table S3. Patients were under the age of 71 years (median, 55 years), diagnosed between 1986 and 1999, and treated uniformly in a single institution. Patients within the good prognosis group (Nottingham Prognostic Index (NPI) <3.4) did not receive adjuvant therapy. Pre-menopausal patients within the moderate (NPI >3.4 to <5.4) and poor (NPI >5.4) prognosis groups were candidates for chemotherapy. Conversely, postmenopausal patients with ER α positive breast cancer with moderate or poor NPI were offered hormonal therapy, while ER α -negative patients received chemotherapy. Clinical data were maintained on a prospective basis with a median follow-up of 130 months (Range 2-311 months).

Immunohistochemistry

Breast cancer tissue microarrays (TMAs) from 1,650 cases were immuno-stained with SRC3 (Dilution 1:20) and SRC3-pS543 (Dilution: 1:2000) antibodies overnight at room temperature, after antigen retrieval (citrate buffer (pH 6.0) and microwave (20 minutes at 750Watts). Negative controls were performed by omission of the primary antibody. Nuclear staining was scored based on H-score. Determination of the optimal cut-offs was performed using X-tile bioinformatics software (Yale University, USA) and according to histogram distribution of H-score. For immunohistochemistry blocking experiments, peptides were used which contained the phospho-moiety (CLL-STLS-phosphoS-PGPKLDN) or the non-phospho peptide CLLSTLSSPGP-KLDN). The peptides were pre-incubated overnight at 100 fold excess over antibody prior to staining as described previously.

Statistical analysis

Data analysis was performed using SPSS (SPSS, version 17 Chicago, IL). Where appropriate, Pearson's Chi-square, Fisher's exact, Student's t and ANOVAs one-way tests were used. Cumulative survival probabilities were estimated using the Kaplan–Meier method and differences between survival rates were tested using the log-rank test. Multivariate analysis for survival

onal hazards asratios (HR) and

was performed using the Cox hazard model. The proportional hazards assumption was tested using standard log-log plots. Hazard ratios (HR) and 95% confidence intervals (95% CI) were estimated for each variable, with p<0.05 as significant. For multiple comparisons, a stringent p value (<0.001) was considered significant.

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Conception and design: W. Zwart, S. Chan, R.C. Coombes, J.S. Carroll, S. Ali, C. Palmieri; Development of methodology: W. Zwart, B. Rudraraju, T.M.A. Abdel-Fatah, O. Gojis, S. Chan, J. Shaw, R.C. Coombes, J.S. Carroll, S. Ali, C. Palmieri; Acquisition of data (provided animals, acquired and managed patients, provided facilities, etc.): W. Zwart, K.D. Flach, B. Rudraraju, T.M.A. Abdel-Fatah, D. Moore, M. Opdam, I. Hofland, S. Chan, I.O. Ellis, R.C. Coombes, C. Palmieri; Analysis and interpretation of data (e.g., statistical analysis, biostatistics, computational analysis): W. Zwart, K.D. Flach, B. Rudraraju, T.M.A. Abdel-Fatah, S. Canisius, E. Nevedomskaya, M. Opdam, M. Droog, S. Chan, J. Shaw, I.O. Ellis, R.C. Coombes, J.S. Carroll; Writing, review, and/or revision of the manuscript: W. Zwart, K.D. Flach, T.M.A. Abdel-Fatah, S. Chan, I.O. Ellis, R.C. Coombes, S. Ali, C. Palmieri; Administrative, technical, or material support (i.e., reporting or organizing data, constructing databases): B. Rudraraju, T.M.A. Abdel-Fatah, M. Opdam, S. Chan, C. Palmieri; Study supervision: W. Zwart, S. Chan, R.C. Coombes The authors declare no conflict of interest.

ERa Cofactor phosphorylation

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Table 1: Patient characteristics and tumor biomarkers and correlation SRC3 andSRC3-pS543See online supplemental information

Supplementary information

Supplementary Figure 1-14 and table 1-6

See online supplemental information

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Table	2 : Multi	variate and	lysis u	sing Cox i	regression	analysis	confirms	that SR	<i>C3-</i>
pS543	protein	expression	is an	independe	nt prognos	stic factor	r for bo	th DFS	and
BCSS									

	BCSS at	10 years	DFS at 10 years		
Variable	HR (CI 95%)	Р	HR (CI 95%)	Р	
S543 (low expression)	1.6 (1.1–2.5)	0.024*	1.4 (1.0–1.9)	0.038*	
Tumor size	1.1 (1.0–1.3)	0.056	1.1 (1.0–1.2)	0.017 ^{-7a}	
Grade		0.018*		0.933	
G1	1.0		1.0		
G2	1.4 (1.3–3.1)		0.98 (0.7–1.4)		
G3	2.1 (2.6–5.7)		1.03 (1.2–2.0)		
Lymph node stage		$9.6\times10^{-11\star}$		$1.6 imes 10^{-10*}$	
Negative	1		1		
Positive (1-3 nodes)	1.7 (1.3–2.3)		1.4 (1.1–1.8)		
Positive (>3 nodes)	3.6 (2.5-5.3)		3.2 (2.3–4.4)		
Endocrine therapy (no)	1.1 (0.8–1.4)	0.661	0.9 (0.7–1.2)	0.565	
Chemotherapy (no)	1.2 (0.9–1.7)	0.259	1.3 (1.0–1.7)	0.03	
Bcl2 expression (positive)	0.4 (0.3–0.6)	6.8 × 10 ^{-8*}	0.5 (0.4–0.7)	9.2 × 10 ^{-7*}	
Ki67 expression (high)	1.5 (1.1–2.2)	0.022*	1.3 (1.0–1.7)	0.093*	
ER expression (negative)	1.5 (1.1–2.1)	0.016*	1.4 (1.0–1.8)	0.036*	
HER2 (overexpression)	1.7 (1.2–2.4)	0.002*	0.7 (0.5–1.0)	0.03*	

*Statistically significant

ERa Cofactor phosphorylation