Selection of Bone Metastasis Seeds by Mesenchymal Signals in the Primary Tumor Stroma

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SUMMARY

How organ-specific metastatic traits arise in primary tumors remains unknown. Here, we show a role of the breast tumor stroma in selecting cancer cells that are primed for metastasis in bone. Cancer-associated fibroblasts (CAFs) in triple-negative (TN) breast tumors skew heterogeneous cancer cell populations toward a predominance of clones that thrive on the CAF-derived factors CXCL12 and IGF1. Limiting concentrations of these factors select for cancer cells with high Src activity, a known clinical predictor of bone relapse and an enhancer of PI3K-Akt pathway activation by CXCL12 and IGF1. Carcinoma clones selected in this manner are primed for metastasis in the CXCL12-rich microenvironment of the bone marrow. The evidence suggests that stromal signals resembling those of a distant organ select for cancer cells that are primed for metastasis in that organ, thus illuminating the evolution of metastatic traits in a primary tumor and its distant metastases.

INTRODUCTION

A key question in understanding the origin of metastasis is how cancer cells in a primary tumor acquire the ability to colonize a particular distant organ. Primary tumors release large numbers of cancer cells into the circulation, yet only a small proportion of these cells survive the stress of invading distant organs and progress to metastases (Chambers et al., 2002; Fidler, 2003; Gupta and Massagué, 2006). Moreover, different tumor types metastasize with distinct patterns of organ preference. It was postulated that these metastatic traits are gained through random prometastatic mutations in primary tumors and remain rare until selection in secondary organ sites leading to expansion of the mutant cell clones (Fidler, 1973; Nowell, 1976). In this model, the molecular determinants of metastasis would not be overtly manifest in the bulk cell population of a primary tumor. However, no driver mutations specific for metastasis to particular organs have been identified to date. Growing evidence shows, on the contrary, that the likelihood of metastasis in general, and of metastasis to certain organs in particular, can be predicted from gene expression patterns of primary tumors (Chang et al., 2005; Chiang and Massagué, 2008; Minn et al., 2005; van ’t Veer et al., 2002; Weigelt et al., 2003). These findings imply that prometastatic activities expressed in large segments of the primary tumor cell population increase the probability that the cancer cells will colonize particular organs.
Src hyperactivity, however, does not confer a net growth advantage in mammary tumors (Zhang et al., 2009), nor is Src frequently mutated or genomically amplified in breast cancer (The Cancer Genome Atlas Network, 2012). These observations raise questions about the mechanisms that lead to the accumulation of Src-hyperactive cells in breast tumors. We addressed these questions by considering two alternative hypotheses. Src hyperactivity in SRS+ breast cancer cells could be biochemically linked to the primary oncogenic alterations driving the tumor or, alternatively, it could result from clonal selection of a Src-dependent growth advantage that is not manifest in the primary tumor. Prior findings, which we confirmed, lent support to the first of these hypotheses in ER+ breast tumors (Collins and Webb, 1999) and HER2+ breast tumors (tumors driven by the ERBB2 oncogene) (Ishizawar et al., 2007; Zhang et al., 2011). However, our search for answers in the third major class of breast cancer, TN tumors, revealed a process of stroma-driven selection for clones that are primed for bone metastasis. This process is driven by CXCL12 and IGF1 from mesenchymal stromal cells, and it selects for Src-hyperactive cancer cell clones that thrive in environments containing these signals. We delineate this process of metastasis seed preselection in experimental models and provide evidence for the existence of its defining traits in human breast tumors.

RESULTS

Src Activation in Different Subtypes of Breast Cancer

To determine what drives Src activation in primary tumors, we investigated the features associated with SRS+ status in different subtypes of breast cancer. Most ER+ breast tumors and at least half of HER2+ tumors score as SRS+ (Figure 1A; Figures S1A and S1B available online) (Zhang et al., 2009). Mechanistically, this is in line with the ability of Src to interact with ER and activate nontranscriptional tumorigenic signaling (Collins and Webb, 1999) and the capacity of Src and ERBB2-ERBB3 dimers for reciprocal activation (Ishizawar et al., 2007; Zhang et al., 2011). We confirmed these links with biochemical and pharmacologic approaches (Figures S1C–S1H). We concluded that Src activation, SRS+ status, and the propensity to metastasize in bone are intrinsically coupled to the oncogenic drivers in these breast cancer subtypes (Figure 1A; Table S1).

TN breast tumors have a propensity to metastasize in visceral organs. However, a significant proportion of these patients develop metastasis to bone as the first site (Figure 1A; Table S1) (Dent et al., 2009). SRS+ status in TN tumors is a strong prognostic indicator of bone relapse (Zhang et al., 2009). Unlike the ER+ and HER2+ subtypes, TN tumors are driven by diverse oncogenic mutations. PIK3CA mutations are frequent in breast cancer (The Cancer Genome Atlas Network, 2012). In the TCGA breast cancer cohort, comprising 409 primary tumor samples with matched mutation and expression data, 43% of SRS+ tumors harbor PIK3CA mutations, versus 32% of all tumors (p = 2.7 × 10⁻⁵). Mutations in other PI3K pathway components such as PTEN did not correlate with SRS status (p = 0.51). Within TN tumors, 16% had PIK3CA mutations, versus 36% in the SRS+ TN subset (p = 0.00027), suggesting a limited contribution of PIK3CA mutations to SRS+ status. Thus, SRS+ status represents an output of diverse upstream mechanisms in TN breast cancer.

CAF Content of TN Tumors Is Linked to Bone Metastasis

Seeking additional clues for how Src activity accumulates in SRS+ TN tumors, we compared the gene expression profiles of SRS− and SRS+ TN tumors in independent clinical data sets and focused on genes that are significantly enriched in SRS+ tumors but are not part of the SRS gene set itself (Figure 1B). A total of 43 genes scored as SRS-associated genes, among which 38 were validated in the TCGA data set (p < 0.001, FDR < 0.05; Figure 1B; Table S2). Notably, these genes included CXCL12, CXCL14, IGF1, and IGF2 (Figure 1C), which are also overexpressed in bone metastasis tissues compared to metastases in lung, liver, or brain (Figure 1D) (Zhang et al., 2009). The enriched expression of these four cytokines in ER− tumors was especially pronounced in TN cases compared to HER2+ cases (Figure 1E). A two-gene classifier comprised of CXCL12 and IGF1 alone was also prognostic of bone metastasis (Figure 1F). These observations pointed at an intriguing resemblance in cytokine content between SRS+ TN tumors and the bone marrow where these tumors preferentially metastasize.

We sought to identify the source of these cytokines in SRS+ TN tumors. None of the TN breast cancer cell lines, including MDA-MB-231 (MDA231 for short), CN34, BT549, MDA-MB-435, or MDA-MB-436 cells, expressed a high level of CXCL12, CXCL14, IGF1, or IGF2 mRNAs (transcriptomic data not shown). To determine whether the association of SRS with CXCL12/ CXCL14 and IGF1/IGF2 mRNA expression in TN tumors reflects an abundance of a particular cell type, we applied gene signature classifiers that designate specific breast tumor stromal cell types. These gene signatures were previously derived from purified cell populations from breast tumors of multiple patients (Allinen et al., 2004) and, therefore, represent consensus profiles of specific stromal types from individually heterogeneous tumors. Notably, SRS+ status in TN tumors was associated with a high score for fibroblast and myofibroblast gene signatures, but not leukocyte or myoepithelial gene signatures (Figure 2A). Indeed, tumor-associated mesenchymal cells (fibroblast and myofibroblast), also referred to as cancer-associated fibroblasts (CAFs), abundantly express CXCL12 (Orimo et al., 2005). In line with this, previous transcriptional data sets showed no CXCL12 mRNA expression and low expression of IGF1 mRNA in cancer cells compared to mesenchymal stroma (Allinen et al., 2004).

The CAF gene set and the SRS-associated gene set included CXCL12, CXCL14, and IGF1 (Figure 2B). Unsupervised hierarchical clustering by the combined mesenchymal stroma gene signature (CAF signature for brevity) identified a set of TN tumors comprising almost exclusively SRS+ cases (Figures 2C and S2A). Gene set enrichment analysis (GSEA) confirmed this association (Figures 2D and S2B). Exclusion of tumors with PIK3CA mutations did not diminish the correlation between CAF+ status and SRS+ status (p = 0.0031, versus p = 0.004 for all tumors included). Importantly, the CAF signature score of TN breast tumors was tightly correlated with the content of stromal cells staining positive for the CAF marker z smooth muscle actin (zSMA) (Figures 2E and 2F). The SRS score was also associated
Figure 1. Cytokine Similarities between SRS+ TN Breast Tumors and Bone Metastases

(A) Src activation in different breast cancer subtypes is shown. Upper panel illustrates hierarchical clustering by SRS, ER, and ERBB2 status of 615 primary breast tumors with known bone metastasis outcomes. Red sticks indicate bone metastasis incidence. Lower panels show that, in ER+ tumors, ER in the cytoplasm interacts with Src to trigger cell survival and proliferation (Collins and Webb, 1999). In HER2+ tumors, ERBB2-ERBB3 heterodimers signal cell proliferation and survival by activating the PI3K-Akt pathway. Src potentiates the ERBB2-ERBB3 heterodimers, and conversely, ERBB2 activates Src (Ishizawar et al., 2007; Zhang et al., 2011). The link to Src in TN tumors is unknown. Neg., negative; Pos., positive. See also Figure S1.

(B) Comparison of SRS+ and SRS- tumors in the TN subtype to identify genes associated with SRS across independent data sets. EMC-MSK (GSE2603, 5327, 2034, and 12276), GSE7390, GSE4922, and GSE1456 data sets were used as discovery data sets, and TCGA data set was used as an independent validation set. The corresponding sample sizes are indicated by pie charts.

(C) Genes that achieve significance (p < 0.001 and fold change >2) in the EMC-MSK data set were subjected to further analysis. For the EMC-MSK data set, p values are log transformed and represented by the color scale. For other data sets, genes with p < 0.05 are indicated in red. All mapped genes are confirmed in TCGA data set (p < 0.001 and FDR < 0.05; see Table S2).

(D) Venn diagram shows the overlap between cytokine genes enriched in bone metastases and SRS-associated genes in primary tumors. The number of genes in each category and the four overlapping genes are indicated.

(E) Microarray gene expression heatmap of the four cytokines within all ER- tumors, including HER2+ and TN tumors. The lower heatmap shows unsupervised clustering of ER-/SRS+ tumors using the four cytokines and ERBB2. Red sticks indicate bone metastasis incidence.

(F) Kaplan-Meier plot illustrates the probability of cumulative bone metastasis (met)-free survival in patients with TN cancer according to CXCL12/IGF1 expression. Numbers of cases in each category are indicated in parentheses. p value was calculated according to log rank test.

See also Table S1.
with the abundance of αSMA+ stromal cells (Figure S2C). The CAF signature was overrepresented in bone metastases compared to metastases at other sites (Figure 2G). These results showed a concordance among CAF stromal content, CAF gene signature score, and CAF-derived cytokines in TN SRS+ tumors.

Notably, both the CAF gene signature (Figures 2H and 2I) and CAF cell content in TN breast tumors (Figures S2D and S2E) were associated with bone relapse, but not with lung relapse. Thus, by molecular and histological criteria, CAF-rich TN tumors were predisposed to metastasize in bone. CAF signature (p = 0.027; hazard ratio 2.36, 95% CI 1.24–4.47) and SRS (p = 0.023; hazard ratio 7.11, 95% CI 1.87–27.10) as continuous variables also correlated with bone relapse as the endpoint. An association of the CAF signature with SRS+ status was also present though less pronounced in ER+ and HER2+ tumors (Figure S2F). Collectively, these results linked the CAF content with the propensity of TN tumors to relapse in bone.

Premises for CXCL12 and IGF1-Driven Selection

The repeated observation of an association among SRS status, CAF gene signature, CXCL12/IGF1 expression, and risk of bone metastasis led us to consider the possibility of a selection process whereby stroma-derived CXCL12 and IGF1 in breast tumors skew cancer cell populations toward a preponderance of Src-hyperactive clones and, thus, toward bone metastasis potential. This “metastasis seed preselection” hypothesis required three premises: first, the presence of a selectable trait—Src hyperactivity—in the cancer cells; second, a heterogeneous distribution of this trait in the cancer cell population; and third, the presence of limiting concentrations of CXCL12/IGF1 that would act as a selector of cell fitness to respond to these survival signals.

To determine the clonal distribution of Src activity in a heterogeneous breast cancer cell population, we analyzed the level of activated Src (Src phosphorylated at Tyr416, pY416-Src) in 100 clonal cell lines isolated from the MDA231 parental population. We quantitated the pY416-Src level in each clone by ELISA. The distribution of values varied over a 12-fold range (Figure 3A). These values remained stable over 25 population doublings in culture (data not shown). Thus, pY416-Src was a heterogeneously distributed, stable trait in the cancer cell population.

To define CXCL12 and IGF1 as positive selectors, we determined the Akt response in cancer cells incubated with a range of CXCL12 and IGF1 concentrations (Figure S3A). Based on the sensitivity of Akt activation, we chose intermediate (30 ng/ml CXCL12, 10 ng/ml IGF1) or saturating (300 ng/ml CXCL12, 100 ng/ml IGF1) concentration of these factors and determined their combined effect on MDA231 by RNA-seq. The transcriptomic responses (fold change >2; p < 0.05) to these additions (data not shown).

CXCL12 and IGF1 Select for Src-Hyperactive Cancer Cells

With these premises established, we modeled CXCL12/IGF1-driven selection by subjecting cancer cells to long-term culture in mitogen-poor media (0.2% fetal bovine serum) with or without supplementation with CXCL12 and IGF1 (Figure 3D). Without supplementation, very few colonies developed. These background clones (MDA231-0 cells) were pooled and expanded in regular media. The level of pY416-Src in MDA231-0 cells was the same as in the parental MDA231 cells (Figure 3E). With CXCL12/IGF1 addition, more colonies grew (Figure S3B). After expansion in regular growth media, the resulting cell population (CXCL12/IGF1-selected MDA231 cells, MDA231-CI) showed a gain in pY416-Src as determined by immunoblotting (Figure 3E) and phospho-flow cytometry (Figure 3F). The level of pY416-Src in MDA231-CI cells matched the upper quartile of the parental MDA231 population (Figure 3A) and had a high SRS score (Figure 3G). Both parameters were stable over at least 25 population doublings in regular culture conditions without CXCL12 or IGF1 additions (data not shown).

The gain in pY416-Src was verified by immunoprecipitation with Src-specific antibody followed by anti-pY416-Src immunoblotting (Figure S3C). Src was not activated by CXCL12 and IGF1, but it rather enhanced the activation of the PI3K-Akt pathway by these cytokines, and this enhancement was inhibited by the broad-spectrum Src kinase inhibitor dasatinib (Figure S3D) (Zhang et al., 2009). Indeed, when inoculated into the arterial circulation of mice, the MDA231-CI cells were significantly more metastatic to bone compared to parental MDA231 and MDA231-0 cells (Figure 4A). Bone lesions formed by MDA231-CI typically affected femora and vertebrae (Figure 4B). MDA231-CI showed no gain in lung metastatic activity (Figures 4A and 4B). The lesions formed by MDA231-CI cells were mostly restricted to the bone marrow cavity, with little engagement of osteoclasts or bone matrix destruction, as compared to lesions formed by the aggressive bone metastatic derivative MDA231-BoM1833 (Kang et al., 2003, 2005; Lu et al., 2009) (Figure 4C). In line with this result, MDA231-CI showed little or no gain in the expression of typical pro-osteolytic genes (Figure 4D). Thus, MDA231-CI cells spontaneously presented a survival and outgrowth advantage in the bone marrow prior to the onset of the osteolytic stage.

Several lines of evidence supported the specificity of these effects. Src knockdown in MDA231-CI (Figures 4E, S4C, and S4D) or administration of dasatinib 2 weeks after MDA231-CI inoculation (Figure 4F) inhibited bone metastases. Bone
Figure 2. CAF Content in TN Tumors Is Linked to Bone Metastasis

(A) Box-whisker plots show the expression of various cancer-associated stromal signatures (Allinen et al., 2004) in TN tumors of the EMC-MSK data set, according to SRS status. The expression level of each signature in each tumor is represented by the average probe intensity of the genes comprising the signature. Differences between SRS+ and SRS− tumors were tested using Wilcoxon signed rank test. **p < 0.0001. Myofibrobl., myofibroblast; n.s., no statistical difference.

(B) Venn diagram shows the overlap between the SRS-associated genes and the CAF signature genes, and identities of the six overlapping genes. p value was calculated by Fisher’s exact test.

(C) Unsupervised hierarchical clustering of TN tumors from the EMC-MSK data set by CAF signature is shown. SRS+ status is indicated by red sticks above the heatmap. Small arrows point to CXCL12 and IGF1 gene probes. Red and green bars left of the heatmap indicate genes that are over- or underexpressed in CAF signature, respectively.

(legend continued on next page)
metastasis inhibition by dasatinib in MDA231 cells is specifically mediated by inhibition of Src in the cancer cells (Zhang et al., 2009). No EGF family members were present in the SRS+--associated gene set or in the CAF signature, and MDA231 cell populations selected with EGF in low-serum media showed little gain in pY416-Src content (Figure S4E) or bone metastatic activity (Figures 4G and S4F).

Cancer cell clones with high Src activity (Src\textsuperscript{high}, refer to Figure 3A) showed an intrinsically high bone metastatic ability independently of osteolytic gene expression (Figures 4H, S4G, and S4H), thus matching the phenotype of the CXCL12/IGF1--selected MDA231-CI population. To test whether Src-hyperactive cells show elevated bone metastatic ability from the orthotopic site, we labeled Src\textsuperscript{high} cells with turbo-GFP, which generates a strong fluorescence signal that facilitates visualization of lesions. Nine weeks after mammary tumor implantation, no significant difference was observed in the size of primary tumors formed by Src\textsuperscript{high} or Src\textsuperscript{low} cells (Figure S4I) or in the accumulation of cancer cells in the lungs (Figure S4J). Notably, Src\textsuperscript{high} mammary tumors showed a much higher rate of spontaneous bone metastases compared to Src\textsuperscript{low} tumors (Figures 4I and 4J), and the proportion of GFP+ cancer cells in bone marrow aspirates subjected to flow cytometry (Figure S4K) was much higher in mice bearing Src\textsuperscript{high} tumors (Figure 4K).

Figure 3. CXCL12 and IGF1 Select for Src-Hyperactive Cancer Cells
(A) Clonal heterogeneity of pY416-Src levels in the MDA231 cell population is presented. A total of 100 random single-cell clones were subjected to pY416-Src ELISA. Values were normalized to the value of the parental population and ranked. The pY416-Src level of the in-vitro-selected subpopulation MDA231-CI is indicated by the red dot.
(B) Principal component analysis of gene expression profiles of MDA231 cells treated with CXCL12 and IGF1 at intermediate (30 plus 10 ng/ml) and saturating (300 plus 100 ng/ml) concentrations is presented. The first three principal components are shown.
(C) Schematic portrays the in vitro selection procedure. Cell selection in media containing 0.2% serum lasted 3 weeks for MDA231 cells or 9 weeks for CN34 cells. CI, CXCL12/IGF1. (D) Schematic illustrates the in vitro selection procedure. Cell selection in media containing 0.2% serum lasted 3 weeks for MDA231 cells or 9 weeks for CN34 cells. CI, CXCL12/IGF1.
(E) pY416-Src levels of MDA231 subpopulations resulted from in vitro selection as determined by western immunoblotting. MDA231-P, parental cells; MDA231-0, cells selected without CXCL12/IGF1 addition; MDA231-CI, cells selected with 30 ng/ml CXCL12, 10 ng/ml IGF1.
(F) Histograms show pY416-Src distribution at the single-cell level as determined by phospho-flow cytometry of the indicated cell lines.
(G) SRS scores derived from gene expression profiles of MDA231-0 and MDA231-CI cells are presented. Scores are −0.76 and −1.0 for two biological replicates of MDA231-0 cells, and 1.0 and 0.58 for MDA231-CI cells.
See also Figure S3 and Table S3.

(D) GSEA revealed a strong association between SRS status and the enrichment of CAF signature genes.
(E) Representative examples and scores of αSMA IHC staining of TN breast tumor samples are shown. Scale bar, 200 μm.
(F) Box-whisker plots show the CAF signature scores (see Experimental Procedures) in tumor tissue microarrays with matched microarray data. The definition and sample size of each category are indicated. Statistical significance is assessed using one-way ANOVA.
(G) Box-whisker plots show the expression of CAF signature in 65 breast cancer metastasis samples from different sites. The number of metastases at each site is indicated in parentheses. The expression in bone metastases was significantly higher (p < 0.0001, Wilcoxon test) compared to all other sites combined.
(H and I) Kaplan-Meier plots illustrate cumulative bone (H) and lung (I) metastasis-free survival in patients with TN according to CAF signature. The median CAF signature score is used as a cutoff to separate CAF+ and CAF− cases. Cases without clinical follow-up information are omitted. The cases included in this analysis are all TN samples with available microarray data. Numbers of relapse cases over total cases in each category are indicated in parentheses. p value was calculated according to log rank test.
See also Figure S2.
CXCL12 and IGF1 Select for Clones that Persist in the Bone Marrow

To further characterize the phenotype selected by CXCL12 and IGF1, we applied the in vitro selection protocol to the CN34 model, which is more indolent than MDA231 and allows analysis of the latent metastasis state (Zhang et al., 2009). Similar enrichment of Src-hyperactive clones was seen after long-term culture of CN34 cells in CXCL12/IGF1-supplemented media (Figures 5A and 5S). We determined the load of human cells in the bone marrow by quantitation of human β2-microglobulin (B2M) mRNA levels (Figure 5B), which can detect as few as ten human cancer cells in 5 x 10^6 mouse cells (Figure 5S). Eight weeks after inoculation of the cells into the arterial circulation of mice, the CXCL12/IGF1-selected CN34 cells (CN34-CI) had a superior ability to persist in the bone marrow (Figure 5C). Under experimental conditions that allow the incipient outgrowth of these cells in the marrow, administration of dasatinib suppressed the outgrowth (Figure 5D).

Transcriptional Shift of Mammary Tumors under a Mesenchymal Stroma

We investigated whether supplementing experimental mammary tumors with mesenchymal cells would lead to the selection of Src-hyperactive cancer cells and a gain in bone...
RNA-seq analysis of the cancer cell transcriptome from freshly harvested breast cancer cells did not secrete CXCL12 or IGF1. Transcript 2011). In their basal state, MSCs secreted CXCL12, and when stimulated with cancer cell-conditioned media, MSCs addition-secreted IGF1 (Figures S6 A and S6B). In contrast, the cancer cell ribosomes, allowing the specific retrieval of cancer cell transcripts from tumor lysates by polysome immunoprecipitation with anti-EGFP antibody (Figure 6A; translating ribosome affinity purification [TRAP]). TRAP (Heiman et al., 2008) enables RNA-seq analysis of the cancer cell transcriptome from freshly excised tumors without intervening cell isolation steps that could confound the results.

MDA231 cells expressing EGFP-L10a were implanted alone as a control or MDA231 admixed with MSCs at a 1:1 ratio into the mammary fat pads of mice. RNA-seq analysis of the TRAP samples showed that tumor supplementation with MSCs caused a transcriptomic shift in the cancer cells that resembled the shift associated with CAF+ status in clinical tumor samples (Figures 6B and 6C). Consistently, MSC-supplemented tumors showed a 3- to 10-fold increase in the proportion of cells staining positive for the CAF marker αSMA (Figure 6D) and a higher level of CXCL12 and IGF1 immunostaining (Figures 6E and 6F). The TRAP RNA-seq data revealed no ribosome-associated CXCL12 or IGF1 transcripts in cancer cells (Figure 6D).

The gene expression profile of cancer cells from MSC-supplemented tumors resembled the profile of MDA231 cells treated with limiting CXCL12 and IGF1 concentrations (Figure 6G). This similarity was further validated by Ingenuity pathway analysis of the differentially expressed genes. Upstream regulator analysis revealed CXCL12- and IGF1-induced signaling and their downstream PI3K-Akt signaling in the cancer cells (Figure 6H). These genes are enriched in biological processes associated with metastasis and bone biology (Figure 6I). Thus, the model tumors cotransplanted with MSCs gained a bioactive CXCL12/IGF1 microenvironment and response features that are characteristic of CAF+ TN tumors.

A Mesenchymal Stromal Selects for Bone Metastatic Cells in Mammary Tumors

Cancer cells were purified from MDA231 mammary tumors, expanded in culture for several passages, and characterized (Figure 7A). Cancer cells recovered from MSC-supplemented tumors showed a high level of pY416-Src compared to populations derived from control tumors (Figures 7B and 7C). Treatment of the mice with inhibitors of the CXCL12 receptor CXCR4 (AMD3100) (Hatse et al., 2002) and of the IGF1 receptor (IGF1R) (BMS754807) (Carboni et al., 2009) (Figure S7A) during tumor growth prevented the accumulation of pY416-Srchigh cells (Figures 7B and 7C), without significantly affecting overall tumor growth. Importantly, cancer cell populations derived from the MSC-supplemented tumors were significantly more metastatic to bone (Figure 7D), without a gain in lung-colonizing ability (Figure 7E).

Cancer cell populations derived from mice treated with IGF1R and CXCR4 inhibitors were poorly metastatic to bone (Figure 7D). To confirm the specificity of these effects, we knocked down CXCR4 and IGF1R simultaneously in MDA231 cells (Figures 7F, 7G, and S7B). Knockdown of the two receptors did not affect the level of pY416-Src in MDA231 cells (Figure 7G), but it prevented MSCs from selecting for carcinoma clones with high pY416-Src content (Figures 7H and 7I). The ability of admixed MSCs to select for clones with high pY416-Src and bone metastatic ability was confirmed in the CN37 cell line (Figures S7C and S7D). Furthermore, primary CAFs isolated from a human breast tumor secreted CXCL12 and IGF1 (12.6 ± 0.2 ng/ml and 0.44 ± 0.02 ng/ml,
respectively, after 96 hr in culture) and selected for pY416-Src-enriched MDA231 cells when admixed in mammary tumors in mice (Figure 7J). The evidence therefore supports a model in which CAFs in mammary tumors cause an accumulation of stromal CXCL12 and IGF1, and these factors act directly on cancer cells to drive the selection of Src-hyperactive clones that are automatically primed for adaptation to the bone metastatic microenvironment.

Figure 6. Transcriptional Shift in Mammary Tumors under a Mesenchymal Stroma
(A) Isolation of cancer cell-specific, polysome-associated transcripts by TRAP for RNA-seq analysis is shown.
(B and C) Principal component analyses of polysome-associated transcriptomic data of MDA231 from xenograft tumors and transcriptomic data of TN tumors from a clinical cohort are presented. MDA231 mammary tumors were formed with MSC supplementation (+MSC) or without (-MSC) supplementation; TN tumors scoring as CAF+ or CAF− based on the CAF signature were from the EMC-MSK cohort. Principal component 1 maximally separates CAF+ from CAF− tumors and was used to generate the plot in (C). p value was calculated with t test.
(D–F) Histological analysis of mammary tumors generated with or without admixed MSCs is presented. (D) IHC staining of αSMA is shown. (E) Immunofluorescent (IF) staining of IGF1 is presented. (F) IF staining of CXCL12 and DAPI nuclear staining are shown. Scale bars, 100 µm.
(G) Principal component analysis using combined data of TRAP RNA-seq of MDA231 in vivo mammary tumors and that of in vitro CXCL12+IGF1 treatment (Figure 3C) is presented. The results show that intermediate CI samples fall closest to MSC+ tumors. Saturating CI samples, on the other hand, are dissimilar. Intermediate CI, intermediate [CXCL12, IGF1]; saturating CI, saturating [CXCL12, IGF1], refer to Figure 3C.
(H and I) Ingenuity pathway analysis of the differentially expressed genes from polysome-associated transcriptomic data of MDA231 mammary tumors in (A)–(C) is shown. IGF1, CXCL12, and AKT signaling are predicted to be activated based on upstream regulator analysis (H). Metastasis-related biological processes are enriched (I).
See also Figure S6.
DISCUSSION

Preselection of Bone Metastatic Seeds by the Primary Tumor Stroma

TN breast cancer is a major and particularly aggressive subtype of the disease. Unlike hormone receptor-positive breast tumors, which have a marked tendency to metastasize mainly in bone, TN tumors as a whole prominently metastasize in nonskeletal organs including lungs, liver, and brain (Dent et al., 2009). We have focused here on a subset of TN tumors that deviates from this trend in that it also metastasizes prominently in bone. Cancer cells from this subset are characterized by a high level of Src activity and a high sensitivity to PI3K-Akt pathway activation by CXCL12 and IGF1 (Zhang et al., 2009). By identifying the primary tumor stroma as a driver of the emergence of this phenotype, the present work illuminates the larger question of how organ-specific metastatic traits may arise in cancer.

Our evidence suggests that, when the stroma of a TN breast tumor is rich in mesenchymal cells, it selects for cancer clones that are fit to thrive on the CAF-derived cytokines CXCL12 and IGF1 (Figure 7K). This selection process leads to a gain in the predisposition of cancer cells for metastasis to bone, a site that is relatively rich in stromal CXCL12 and IGF1 compared to other metastasis target sites. In other major subtypes of breast cancer, Src activation is integral to the oncogenic drivers like ERα (Collins and Webb, 1999) and ERBB2 (Ishizawar et al., 2007; Zhang et al., 2011) and might similarly contribute to bone metastatic tropism of these tumors.

The clinical evidence also links the CAF content with a predisposition for bone metastasis in TN tumors. In multiple human breast cancer cohorts, TN tumors that score high for cancer cells was dependent on CXCR4 and IGF1R expression in the tumor microenvironment, as determined by analysis of breast cancer cohorts, TN tumors that score high for cancer cells also does not entail an acquisition of specific metastatic traits in other organs.

The ability of the primary tumor stroma to drive the selection of organ-specific metastatic cells is distinct from previously described roles of stromal signals in tumor progression (Chung et al., 2010; Hanahan and Coussens, 2012; Hanahan and Weinberg, 2011; Joyce and Pollard, 2009), including roles in the resistance to stress and chemotherapy (Acharyya et al., 2012; Calon et al., 2012; DeNardo et al., 2011; Gocheva et al., 2010; Qian et al., 2011). These previously described stromal signals do not explain how organ-specific metastatic traits and their associated gene signatures emerge in primary tumors. The formation of premetastatic niches by tumor-derived systemic signals that condition distant organs for seeding by disseminated cancer cells also does not entail an acquisition of specific metastatic traits in the primary cancer cell population itself (Kaplan et al., 2005). The mechanism of metastasis seed preselection identified here may coexist with other effects of the tumor stroma but is distinct in that it selects for organ-specific metastatic traits.

The interplay between a primary tumor and a specific metastatic site revealed by the present work expands current views on the dynamic nature of metastasis. Coupled with processes of tumor self-seeding whereby previously disseminated cells may repopulate and expand in their tumor of origin (Kim et al., 2009), seed preselection at the primary site may further sway
Figure 7. A Mesenchymal Stroma Selects for Bone Metastatic Cells in Mammary Tumors

(A) Experimental strategy is presented. MDA231 cells alone or admixed with MSCs were implanted in mammary fat pads. One set of mice was treated with CXCR4 inhibitor (CRi, AMD3100) and IGF1R inhibitor (IRi, BMS754807). After reaching 1 cm³, tumors were excised, and cancer cells were recovered and expanded in culture for testing.

(B and C) pY416-Src levels in cancer cell lines from individual mammary tumors formed by MDA231 under the indicated conditions are presented: immunoblot (B) and quantification (C).

(D and E) Bone metastasis (D) and lung colonization (E) activity of the pooled cell lines derived in (B) is illustrated. Normalized bioluminescence signals were quantified at 21 days (D) or 30 days (E) after inoculation. Error bars, SEM (n = 8–10 in each group).

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the organotropic metastatic traits of a primary tumor mass with clinically manifest consequences. Our results also imply that breast cancer cases with a fibroblast-rich stroma or with incipient bone metastasis may benefit from therapies that target the PI3K-Akt cell survival pathway or its amplifier Src for the eradication of disseminated disease.

**EXPERIMENTAL PROCEDURES**

**Computational Analyses of Clinical Data Sets**

SRS was derived as previously described (Zhang et al., 2009). Stratification of the 615 patients with SRS, ER, and ERBB2 in the Erasmus Medical Center/Memorial Sloan-Kettering Cancer Center (EMC-MSK) cohort (GSE2603, GSE5327, GSE2034, and GSE12276 data sets) was done with unsupervised clustering implemented by the “heatmap.2” command of the “gplots” package of R statistical programming software. ER+, HER2+, (ER+/HER2+), and TN (ER-/HER2-) subtypes were determined either using the clinical annotation associated with the data set (if applicable) or by the transcript levels of the ESR1 and ERBB2, respectively. The continuous scores of SRS were computed with the sum of Z scores of 159 SRS genes and the DART algorithm (Liao et al., 2013).

To identify genes associated with SRS status, SRS was applied to EMC-MSK, Transibig (GSE7394), Uppsala (GSE4922), and Stockholm (GSE1458) data sets. SRSs and SRSs tumors were compared within the TN subset of each data set. Genes that are expressed at a significantly higher level in SRS+ tumors (p < 0.001 and fold change >2 for EMC-MSK, and p < 0.05 for the other three, t tests with Welch correction) were identified. To compute the combined p value for each gene across all four data sets, we employed Fisher’s combination (see Extended Experimental Procedures). To further evaluate the significance of these genes, we utilized the TCGA cohort as a validation data set (https://tcga-data.nci.nih.gov/tcga/tcgaDownload.jsp). We examined the p values and FDR of the 43 genes by comparing TN/SRS+ tumors with TN/SRS− tumors. Both p values and FDR were computed using the “signR” package in R statistical software based on SAM algorithm (Tusher et al., 2001). All 38 genes that can be mapped to TCGA platform reach significance (Table S2).

Gene signatures associated with CAFs were obtained from a previous study by Allin et al. (2004). We used unsupervised hierarchical clustering, the mean expression of CAF genes, and GSEA to determine the CAF status of a tumor. Other stromal cell types were analyzed using the same approaches (see Extended Experimental Procedures for details). For analyses of correlation between somatic PIK3CA mutation status and SRS, somatic mutation data from the TCGA data set were matched with mRNA expression data. The overlap between PIK3CA mutation and SRS+ status was assessed with Fisher’s exact test.

To examine the correlation among CAF gene expression signature, SRS, and αSMA immunohistochemical (IHC), we divided the IHC staining into six groups depending on the percentage of αSMA+ fibroblastic cells (see Figure 2E). CAF and SRSs were then evaluated as continuous scores (described above) in different categories (Figures 2F and S2C). The statistical significance among the three groups was assessed by one-way ANOVA implemented by R.

**Cancer-Cell-Specific Polysome-Associated Transcript Profiling**

MDA231 cells were engineered to express an EGFP-L10a construct (Heiman et al., 2008). For the analysis of cytokine responses in vitro, cancer cells were treated for 6 hr with 30 ng/ml CXCL12 and 10 ng/ml IGFI1 or with 300 ng/ml CXCL12 and 100 ng/ml IGFI1. For the analysis of gene responses to MSCs in vivo, breast cancer cells were injected into the mammary fat pads of mice with or without a 1:1 admixture of MSCs, and tumors were harvested 3 weeks later. Cultured cells or tumors were lysed and immediately subjected to TRAP as previously described by Heiman et al. (2008) with modifications described in the Extended Experimental Procedures.

For RNA-seq data analysis, the raw sequence data (FASTQ files) were mapped using TopHat (v.1.4.0) (Trapnell et al., 2009) to the human genome (hg19 build) available from UCSC. Of all reads, 89% were uniquely mapped on average. Gene level expression counts were determined using HTSeq and normalized with DESeq (Anders and Huber, 2010). The normalized counts were then log transformed for analysis. Genes with zero counts were still assigned as zeros after log transformation to avoid exceptions in programming. The approaches of subsequent analyses are described in the Extended Experimental Procedures.

**Animal Studies**

All animal work was done in accordance with a protocol approved by the MSKCC Institutional Animal Care and Use Committee. Intracardiac injections of cancer cells for bone metastasis assays, intravenous injections for lung colonization assays, and orthotopic injections for mammary tumor formation were performed as previously described by Kang et al. (2003) and Minn et al. (2005). Animals were imaged weekly by bioluminescence imaging (IVIS; Xenogen). For quantification of bioluminescence signal, all signals were normalized to the signals at day 0 right after injection. For bone metastases, signals at hindlimbs and vertebrae were jointly quantified. Osteolytic lesions were visualized by X-ray radiography and were identified on radiographs as demarcated radiolucent lesions in the bone. Cancer cells from mammary tumors were retrieved by enzyme digestion (0.125% collagenase III and 0.1% hyaluronidase in DMEM/F12 medium) and GFP sorted for further experiments or inoculation into mice. Datasatiniti (Bristol-Myers Squibb) was administered by daily oral gavage with a dose of 10 mg/kg in PBS. AMD3100 (Tocris Bioscience; catalog S299), a CXCR4 inhibitor, was administered by subcutaneous injection twice a day at 2.5 mg/kg in PBS. BMS754807 (ChemieTek; catalog CT-BMS75), an IGF1R inhibitor, was administered by daily oral gavage at 6.25 mg/kg in polyethylene glycol:H2O (4:1).

Primers for SYBR Green Mix reaction (Thermo Scientific) are listed in Table S4.

**ACCESSION NUMBERS**

Microarray and RNA-seq data can be obtained online with GEO accession numbers GSE39102, GSE43296, GSE43306, and GSE47389.

**SUPPLEMENTAL INFORMATION**

Supplemental Information includes Extended Experimental Procedures, seven figures, and four tables and can be found with this article online at http://dx.doi.org/10.1016/j.cell.2013.07.036.

(F) Flow cytometry analysis of CXCR4 surface expression in MDA231 with two independent double knockdowns (DKD1, DKD2) of CXCR4 and IGF1R compared to control (ctrl) is shown.

(G) Western immunoblot analysis of IGF1R and pY416-Src protein levels in two DKD cell lines is presented.

(H and I) pY416-Src levels in cancer cell lines from mammary tumors formed by control MDA231 (ctrl) or two DKD cell lines with or without MSC supplementation are shown: immunoblot (H) and quantification (I).

(J) pY416-Src levels of cancer cell lines from individual mammary tumors formed by MDA231 cells with or without admixed CAFs from a human breast tumor are presented.

(K) A model of bone metastasis seed preselection by a mesenchymal-rich stroma in the mammary tumor is shown. Blue and gray cells represent mesenchymal and nonmesenchymal stromal cells, respectively. As the tumor grows in a mesenchymal-rich microenvironment, the cancer cell population is skewed toward a preponderance of clones that thrive on mesenchymal cytokines such as CXCL12 and IGFI1, which selects for a predisposition to metastasize to CXCL12- and IGFI1-rich bone marrow.

See also Figure S7.
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