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Evolution of Cancer Stem-like Cells in Endocrine-Resistant Metastatic Breast Cancers Is Mediated by Stromal Microvesicles

Pasquale Sansone¹, Marjan Berishaj¹, Vinagolu K. Rajasekhar¹, Claudio Ceccarelli², Qing Chang¹, Antonio Strillacci^{1,3}, Claudia Savini^{1,2,4}, Lauren Shapiro⁵, Robert L. Bowman⁶, Chiara Mastroleo¹, Sabrina De Carolis^{2,4}, Laura Daly¹, Alberto Benito Martin⁷, Fabiana Perna⁸, Nicola Fabbri⁹, John H. Healey⁹, Enzo Spisni³, Monica Cricca², David Lyden^{7,10,11}, Massimiliano Bonafé^{2,4}, and Jacqueline Bromberg^{1,12}

Abstract

The hypothesis that microvesicle mediated miRNA transfer converts noncancer stem cells into cancer stem cells (CSC) leading to therapy resistance remains poorly investigated. Here we provide direct evidence supporting this hypothesis, by demonstrating how microvesicles derived from cancer associated fibroblasts (CAF) transfer miR 221 to promote hormonal therapy resistance (HTR) in models of luminal breast cancer. We determined that CAF derived microvesicles horizontally transferred miR 221 to tumor cells and, in combination with hormone therapy, activated an ER^{lo}/Notch^{hi} feed forward loop responsible for the generation of CD133^{hi} CSCs. Importantly, microvesicles from patients with HTR metastatic disease expressed high levels of miR 221. We further determined that the IL6 pStat3 pathway promoted the biogenesis of onco miR 221^{hi} CAF microvesicles and established stromal CSC niches in experimental and patient derived breast cancer models. Coinjection of patient derived CAFs from bone

metastases led to *de novo* HTR tumors, which was reversed with IL6R blockade. Finally, we generated patient derived xenograft (PDX) models from patient derived HTR bone metastases and analyzed tumor cells, stroma, and microvesicles. Murine and human CAFs were enriched in HTR tumors expressing high levels of CD133^{hi} cells. Depletion of murine CAFs from PDX restored sensitivity to HT, with a concurrent reduction of CD133^{hi} CSCs. Conversely, in models of CD133^{neg}, HT sensitive cancer cells, both murine and human CAFs promoted *de novo* HT resistance via the generation of CD133^{hi} CSCs that expressed low levels of estrogen receptor alpha. Overall, our results illuminate how microvesicle mediated horizontal transfer of genetic material from host stromal cells to cancer cells triggers the evolution of therapy resistant metastases, with potentially broad implications for their control.

¹Department of Medicine, Memorial Sloan Kettering Cancer Center, New York, New York. ²Department of Experimental, Diagnostic and Specialty Medicine, AlmaMater Studiorum, Università di Bologna, Bologna, Italy. ³Department of Biological, Geological and Environmental Sciences, Università di Bologna, Bologna, Italy. ⁴Center for Applied Biomedical Research Laboratory, Policlinico Universitario S. Orsola Malpighi AlmaMater Studiorum, Università di Bologna, Bologna, Italy. ⁵Department of Radiation Oncology, Kaiser Permanente, Oakland, California. ⁶Cancer Biology and Genetics Program, Memorial Sloan Kettering Cancer Center, New York, New York. ⁷Department of Pediatrics, Cell and Developmental Biology, Children's Cancer and Blood Foundation Laboratories, Weill Cornell Medicine, New York, New York. ⁸Molecular Pharmacology and Chemistry Program, Memorial Sloan Kettering Cancer Center, New York, New York. ⁹Orthopedics Service, Memorial Sloan Kettering Cancer Center, New York, New York. ¹⁰Drukier Institute for Children's Health, Meyer Cancer Center, Weill Cornell Medicine, New York, New York. ¹¹Department of Pediatrics, Memorial Sloan Kettering Cancer Center, New York, New York. ¹²Department of Medicine, Weill Cornell Medicine, New York, New York.

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D. Lyden, M. Bonafé, and J. Bromberg contributed equally to this article.

Corresponding Author: Jacqueline Bromberg, Memorial Sloan Kettering Cancer Center, 1275 York Avenue, Box 397, New York, NY 10065. Phone: 646 888 3112; Fax: 646 888 3200; E mail: bromberj@mskcc.org

Introduction

Tumor heterogeneity and resistance to therapy may occur from microvesicle mediated transfer of genetic material between cells (1-3). Thus, the characterization of this phenomenon could have important clinical ramifications most notably in the development of new therapeutically relevant compounds.

Although adjuvant hormonal therapy (HT) improves disease free survival in luminal breast cancer patients, HT resistant (HTR) metastatic disease commonly develops in the bones of these patients. This observation suggests that the bone microenvironment may foster estrogen receptor (ER) independent growth of luminal breast cancer leading to HTR metastases.

The interaction of stromal cells (cancer associated fibroblasts; CAF) with tumor cells has been shown to mediate and modulate estrogen receptor dependent (e.g., fibronectin, collagen) and independent proliferation (e.g., laminin) of luminal breast cancer cells, suggesting that stroma-tumor communication may play a pivotal role in the ER independent self renewal of breast cancers (4). In the metastatic microenvironment, we hypothesize that chronic inflammation incurred by anti estrogen therapy and the effects of disseminated tumor cells on the local microenvironment will lead to the activation of resident stromal cells or circulating mesenchymal stem cells to become CAFs. Once

activated, the CAFs may sustain a feed forward circuit of self renewal, proliferation, and differentiation of CSCs, resulting in metastasis.

As tumors become more metastatic and resistant to targeted therapies, the number and types of CSCs increases, suggesting that CSCs evolve from non CSC cells in a given tumor niche (5, 6). The role of stroma microvesicles (MV) in the generation of therapy resistant cancer and the regulation of self renewal remains poorly investigated.

Here, we investigated the hypothesis that HT and CAF derived microvesicles converge to promote HT resistance and ER independent self renewal in luminal breast cancer. By employing patient derived xenografts from breast cancer bone metastases and experimental models of luminal breast cancer, we uncovered a unique process of CAF mediated resistance to HT. Our data demonstrate the formation of therapy resistant stromal tumor niches via an IL6/Stat3 driven expansion of CAFs, CAF MV mediated oncomiR 221 transfer to cancer cells leading to the expansion of Notch3^{hi}/ER^{lo}/CD133^{hi} CSCs. These data reinforce the concept of targeting the stromal niche to prevent both HT resistance and metastatic progression (7-9).

Materials and Methods

Microvesicle isolation and *in vivo* education experiment

Plasma (10 mL) was collected and processed within 4 hours from patients with metastatic disease (Supplementary Table S1) and in healthy controls who were consented to an MSKCC biospecimen protocol #12 137. The plasma and conditioned media (CM) from cancer and CAF cultures was collected from 10⁷ cells grown in 5 × 10 cm dishes and centrifuged for 20 minutes at 3,000 × *g* at 4°C. The supernatant was subsequently centrifuged for 30 minutes at 12,500 × *g* at 4°C. The supernatant was transferred and centrifuged at 100,000 × *g* for 90 minutes at 4°C. The supernatant was discarded while the pellet, containing microvesicles, was resuspended in 25 mL of PBS and loaded onto a 5 mL 30% sucrose cushion to deplete microvesicles from extracellular proteins (300 g/L sucrose, 24 g/L Tris base, pH 7.4). Samples were centrifuged at 100,000 × *g* for 90 minutes 4°C. Cushion (3.5 mL), containing microvesicles, was diluted with 10 mL of PBS and centrifuged at 100,000 × *g* for 90 minutes at 4°C. The supernatant was discarded and the pellet resuspended in 25 μL of PBS. Microvesicles were treated with 0.1 mg/mL of DNase I solution (Epicentre) to eliminate contaminating DNA bound to the microvesicles surface or present in solution. Nanosight (Lyden laboratory, Cornell Medical Center) and electron microscopy (MSKCC Electron Microscopy Core) were used to characterize the physical structures of these microvesicles (size and distribution). Confocal microscopy (MSKCC Microscopy Core) of cancer cells educated with pre-labeled (PKH67 Green Fluorescent Cell Linker Kit, Sigma) CAF MVs was performed to ensure transfer and uptake. The *in vivo* role of CAF MVs in the promotion of HT resistant luminal breast cancer was determined by injecting CAF MVs (Mu CAFs, isolated from HT resistant xenografts and cultured *in vitro*) and control microvesicles (from MCF7 cells) into the arterial circulation (retro orbital injection, 3 × 10⁹ particles/mouse/weekly) of tumor bearing mice (MCF7 cells). Once mammary fat pad (MFP) xenografts were established (after 4 months), mice were treated with HT (fulvestrant; 100 μg/injection/once a week for 3 months).

Primary cultures and patient derived xenografts of endocrine resistant luminal breast cancer bone metastases

Patient derived xenografts (PDX) were established from *n* = 2 of 5 HT resistant and 1 *de novo* stage IV breast cancer bone metastatic tissue isolated at MSKCC (Supplementary Table S2). Patients who developed metastatic breast cancer in the bone were enrolled in the study (IRB protocol #97 094). Following surgery, tissue was processed by the pathologist, 60% of the specimen was used for confirmation of diagnosis and molecular analyses (IHC, IMPACT analyses), while 40% was used for further analysis. Tissues were placed in sterile Epicult (Voden Medical), minced with sterile razor blades and incubated at 37°C for 8 to 12 hours in the presence of Collagenase/Hyaluronidase enzyme mix (1,000 Units, Voden Medical). To grow tumor cells devoid of its cognate stroma, we performed serial centrifugations to separate epithelial cells as mammosphere cultures (MS). Secondary and tertiary MS potential (II or III MS) was performed as follows: 7 day primary MS started to form after 4-6 days, then they were disaggregated in 1 × Trypsin EDTA (StemCell Technologies), washed in complete MEGM, filtered through a 40 μm nylon mesh, and seeded to form second generation MS. Number of MS was assessed by counting the total number of spheres (size > 100 μm) from cells seeded in low attachment plates (from 100 to 1,000). To establish primary CAF cultures from patient derived tissue, MS depleted supernatant was centrifuged at 450 × *g* for 10 minutes; this pellet was enriched with stromal cells was plated onto 10 cm plates supplemented with DMEM 10% serum media. CAF primary cultures were expanded *in vitro* for *n* = 10 passages. III MS primary cultures were used to establish PDXs: 50-100 MS (size ~100 μm) were injected in the MFP of NOD/SCID mice and tumor growth was determined over a period of 5 months. At the endpoint of the experiment, xenograft tissues were collected and primary PDX cultures were established. Multiple passaged PDX were generated following repeated orthotopic injection of PDX derived EpCAM^{pos} cancer cells (recognizes only human cells) in the MFPs of NOD/SCID mice (from 1st to 4th generation). Luminal breast cancer cells expressing a vector for GFP/Luciferase were generated and used for all the *in vivo* experiments. Tumor growth was determined using *in vivo* bioluminescence technology (BLI: Xenogen, IVIS System). Luminal cancer xenografts from the coinjection of human CAFs and MCF7 cells were also generated to determine the effect of the stroma on the generation of *de novo* resistant endocrine tumors

Xenograft assays and preclinical trials

All cancer cell lines were engineered to express a GFP positive luciferase expression vector. Prior to *in vivo* inoculation, cancer cells were FACS purified (for GFP) and injected bilaterally in the MFPs of 5- to 7 week old NOD/SCID mice (obtained from NCI, Frederick, MD). For each *in vivo* experiment, cancer cells were mixed with an equal volume of Matrigel (BD Biosciences) in a total volume of 50 μL. Bioluminescence was used to monitor both tumour growth (weekly) and metastatic burden (at necropsy). Luminal cancer xenografts from the coinjection of human CAFs (HTR bone metastases) and MCF7 cells (10³ cells) were also generated to determine the effect of the stroma on the generation of *de novo* resistant endocrine tumors. In addition, human bone marrow stromal cells HS27a, HS27shC, and HS27shIL6 (100 cells/injection) were coinjected with MCF7 cells (10³ cells/injection) into the MFP. For immunostaining assays, organs were

collected and fixed overnight in 4% paraformaldehyde, washed, embedded in paraffin, and sectioned (Histo Serve Core). Hematoxylin and eosin (H&E) staining was performed by standard methods. For the detection of metastases at secondary sites, we performed *in vivo* BLI as well as immunofluorescence/IHC staining for GFP and ER. All the surgical procedures and animal care followed the institutional guidelines and an approved protocol from our IACUC at MSKCC. For the preclinical studies, injectable fulvestrant (Faslodex, AstraZeneca) was given intramuscularly in the tibialis posterior/popliteal muscles (100 µg/injection/once a week) for 2 months. Tocilizumab (Actemra, Roche Pharmaceuticals) was diluted in PBS at a final concentration of 20 mg/mL. A dosage of 100 µg/g/mouse was administered intraperitoneally every week (this is ~5 fold higher than the physiologic range, patients receive 8 mg/kg i.v.). Control mice received isotype control (placebo) or PBS injection.

Cell lines and FACS

Human cancer cell lines (Namalwa, lymphoma; HeLa, cervical carcinoma), human breast cancer cell lines (MCF7, ZR751, T47D, and BT474), human bone marrow stromal cell lines (HS5, HS27a), and human normal fibroblasts (MRC5, HMF) were purchased from the ATCC and authenticated by short tandem repeat (STR) DNA profiling (Genomic Core MSKCC). Murine CAFs (Mu CAF) were isolated from HTR xenografts and PDXs by FACS purification (GFP⁻/EpCAM⁻). Cells were mycoplasma free and maintained in minimum essential medium and RPMI (ATCC and Media Core) supplemented with 5% FBS (Media Core), 2 mmol/L glutamine, 100 U mL⁻¹ penicillin, and 0.1 mg mL⁻¹ streptomycin (Media Core). Cancer cells from xenografts were isolated from primary and metastatic tissues by enzymatic digestion (Collagenase/Hyaluronidase, Sigma Aldrich), sorted (GFP⁺/DAPI⁻), and cultured *in vitro*. The following reagents: 4 hydroxytamoxifen and fulvestrant were purchased from Sigma (Sigma Aldrich). For FACS/flow analyses, tumors were digested in sterile Epicult media (StemCell Technology), minced with sterile razor blades, and incubated for 3 hours in the presence of collagenase/hyaluronidase (1,000 Units/sample). Cells were washed with sterile filtered PBS supplemented with 1% BSA (PBS BSA 1%) and filtered through a 40 µm nylon mesh (BD Biosciences). For the detection of CD44 and CD133, EpCAM antigens, cells were stained in a volume of 100 µL (PBS BSA 1%) with each antibody CD44 APC (100 ng/10⁶ 10⁸ cells Clone IM7, eBiosciences), CD133/1 PE (100 ng/10⁶ 10⁸ cells, clone AC133, Miltenyi Biotech) and EpCAM FITC (250 ng/10⁶ 10⁸ cells, Clone VU 1D9, StemCell Technologies). Cells were labeled on ice for 30 minutes and analyzed (BDFACS Aria III, Flow Core). Samples were analyzed for cell population distribution and sorted for GFP/viability (GFP⁺/DAPI⁻) and CD133/CD44 expression. For flow plot analyses, samples were run using FlowJo 7.5 software (Tree Star). shRNAs for Notch3 and IL6 were previously described (10, 11).

Microarray and miRNA analyses

Normalized gene expression values were downloaded from the GEO under accession number GSE17705 and probes were aggregated to median gene level expression. A CAF gene set from Allinen and colleagues (12) was used in a single sample gene set enrichment analysis (ssGSEA). ssGSEA scores were z scored and a "CAF score" was assigned to each patient. Patients were split at the

median into CAF high and CAF low groups. PROM1 high and PROM1 low groups were split based on PROM1 (CD133) median expression. Statistical significance for differences in PROM1 expression and ESR1 expression were assessed with a Student *t* test. The heatmap for CAF signature gene and PROM1 was plotted in R with the heatmap.2 function. For real time PCR (qPCR), we extracted RNA using TRIzol (Invitrogen). RNA concentration was determined with a NanoDrop 2000. For microarray analysis of published datasets, normalized gene expression data were downloaded from the Gene Expression Omnibus (GEO). Each gene was mean centered and scaled by SD. All analyses were conducted in R. Normalized gene expression data was downloaded from the NCBI for dataset GSE69280 (5). For qPCR, 1 µg of total RNA was reverse transcribed to cDNA using iScript Select cDNA Synthesis Kit (Bio Rad) following the manufacturer's protocol. Reverse transcription PCR (RT PCR) analysis was performed using the following primers: ERα; forward 5' TGAAAGTGGGATACGAAAA GAC 3', reverse 5' CAGGATCTCTAGCCAGGCACAT 3'; β2µ forward 5' ACCCCCACTGAAAAGATGA 3', reverse 5' ATCTCA AACCTCCATGA 3'. DNA was isolated using phenol/chloroform technique from PDX derived EpCAM positive/negative cells. The presence of murine and human cells was determined on 2 ng of DNA by PCR for GAPDH (murine: forward 5' AGCAGCCG CATCTTCTGTGCAGTG 3', reverse 5' GGCCTTGACTGTGCC GTTGAATTT 3'; Human: forward 5' CTCTGCTCCTCTGTTC GAC 3', reverse 5' ACGACCAAATCCGTTGACTC 3'). miRNA expression was analyzed as described previously (13). Briefly, miRNA were reverse transcribed using stem loop RT PCR technology (14) and amplified by real time PCR using SYBR Select Master Mix (Applied Biosystems) and ViiA 7 Real Time PCR System (Applied Biosystems) according to the manufacturer's instructions. The melting curve data were collected to check PCR specificity. miRNA expression was normalized against RNA U6 levels: (RT miR 221) 5' GTCGTATCCAGTGCAGGGTCCGAG GTATTCGCACTGGATACGACGAAACCC 3'; (RT miR 222) 5' GTCGTATCCAGTGCAGGGTCCGAGGTATTCCGACTGGATAC GACACCCAGT 3'; (RT miR 101) 5' GTCGTATCCAGTGCA GGGTCCGAGGTATTCCGACTGGATACGACTTCAGTT 3'; (forward miR 221) 5' AGCTACATTGTCTGCTGGGTTTC 3'; (forward miR 222) 5' AGCTACATCTGGCTACTGGGT 3'; (forward miR 101) 5' GCCGCTACAGTACTGTGA 3'; (forward U6) 5' CTTCGGCAGCACATATACT 3'; (reverse U6) 5' AAAATATG GAACGCTTCACG 3' (reverse all miRs) 5' TGCAGGGTCC GAGGTAT 3'. All primers were purchased from Eurofins MWG Operon. miRNA expression was analyzed as described elsewhere (13).

Protein and *in vitro* studies

For immunoblotting assays, cells were lysed in buffer (50 mmol/L Tris at pH 7.5, 150 mmol/L NaCl, 5 µg/mL aprotinin, pepstatin, 1% NP 40, 1 mmol/L EDTA, 0.25% deoxycholate, and protease inhibitor cocktail tablet, Sigma). Proteins were separated by SDS PAGE, transferred to polyvinylidene difluoride membranes and blotted with specific antibodies (Supplementary Table S3). For functional interference studies, anti-miR 221 and control RNA oligonucleotide were purchased from Applied Biosystems. MCF7 cells were seeded in a 6 well plate (8 × 10⁵ cells/well) at 60% confluence. After 24 hours, cells were transfected using Lipofectamine 2000 transfection reagent (Invitrogen) according to the manufacturer's instructions (RNA final concentration, 200 nmol/L). After 6 hours of

incubation at 37°C, transfection medium was replaced with 2 mL of complete medium containing 10% FCS supplemented with/without CAF MVs. For determination of cell viability, we seeded 2,500 cells per well in 96 well plates and treated them with fulvestrant (10 µmol/L). Viable cells were determined 7-14 days after treatment using Trypan blue and cell counting was done using bright field microscopy or DAPI staining by flow cytometry (Dako Cytomation). Crystal violet assay was performed to obtain information of the relative cell density at the endpoint of proliferation potential experiments. IL6 ELISA assays were performed using the conditioned medium collected from 5 day cultures of CAF derived cells seeded at 2×10^6 cells/plate. Proliferation assay was carried out using CalceinAM technology (Invitrogen): cells were seeded in 96 well plates treated with the prefluorescent compound for 20 minutes and fluorescence was read using a plate reader (SpectraMax plate platform). To determine the selective growth potential of cancer cells over stroma cells, we analyzed the proliferation potential of luciferase positive cancer cells by *in vitro* BLI: cells were seeded in 96 well plates in presence/absence of distinct CAFs/normal fibroblast (1:10 ratio of CAFs to tumor cells) and treated with fulvestrant (10 µmol/L/weekly for 3 weeks). Luciferase activity was measured weekly. Cytokine Arrays were performed on 10 µg of CM derived proteins according to manufacturer's protocol (Antibody Array 3, Ray Biotech. Inc.).

Immunostaining analysis

Serial sections of formalin fixed paraffin embedded samples were immunostained using monoclonal anti CD133 diluted 1:70 (clone W6B3C1, Miltenyi Biotec), anti ER α RTU (clone SP1, Ventana), anti Pankeratin RTU (clone AE1/AE3/PCK26, Ventana), and polyclonal anti Notch 3 diluted 1:400 (M 134, Santa Cruz Biotechnology). CD133 and Notch3 immunostaining was performed as follows: sections were dewaxed, rehydrated, and subjected to antigen retrieval treatment. Antigens were unmasked with a Tris EDTA pH 9.0 buffer at 98°C for 20 minutes in a waterbath. Endogenous peroxidase activity was inhibited using a 0.5% H₂O₂ solution in methanol for 20 minutes at room temperature. Sections were processed using a non biotin amplified method (Novolink, Novocastra) according to the manufacturer's protocols. When mouse tissue was used, a short treatment (30 minutes at room temperature) with MOM blocking solution (Vector Laboratories Inc.) was conducted prior to primary antibody overnight incubation at 4°C. The reaction was visualized using the UltraView DAB Detection System. The immunologic reaction was developed using a 3,3'-diaminobenzidine (DAB)/H₂O₂ PBS pH 7.2-7.4 solution for 10 minutes. Sections were then washed in distilled water and counterstained in Harris hematoxylin. Anti ER α (ER) and Pankeratin (CK) immunostaining was performed on an automated immunostainer (Benchmark Ultra, Ventana) using the UltraView DAB Detection kit according to the manufacturer's protocol. Antigen retrieval was performed onboard with UltraCC1 buffer (pH 8.2-8.5) at 95°C for 52 minutes (ER) or 20 minutes (CK). Primary antibodies were incubated 28 minutes at 37°C (ER) or 8 minutes at room temperature (CK). For CD133 and Notch3 evaluation, each section was examined at 400 \times . In each microscopic field, the neoplastic cells were classified according to both positive percentage and staining intensity: [percentage 0 if <1%, 1 if >1% < 25%, 2 if >25% <

50%, 3 if >50% <75%, 4 if >75%; intensity 1 (weak), 2 (moderate), and 3 (strong)]. A final classification was obtained by multiplying the two mean values (percentage and intensity, IRS score). As for ER evaluation, the neoplastic population was scanned using Image Cytometry and reported as percentage of positive cells (%) (IMAGE Pro Plus V5.0.1, Media Cybernetics Inc.). A detailed histologic examination of xenograft tissues was performed at the collaborating institution (University of Bologna, Bologna, Italy). Xenograft tissue was stained with hematoxylin and eosin and examined by three independent pathologists (C. Ceccarelli, Donatella Santini, and M. Bonafe, from the University Hospital of Bologna, Bologna, Italy). For each microscope field (200 \times), the area occupied from cancer cells, stromal cells, and necrotic components was evaluated and represented as percentage.

Characterization of CAFs

Serial sections (5 µm) of paraformaldehyde fixed paraffin-embedded samples underwent antigen retrieval using Leica Bond ER2 Buffer (Leica Biosystems) for 20 minutes at 100°C before staining with 1 µg/mL Desmin rabbit polyclonal antibody (Abcam catalog no. ab8592) and 1 µg/mL pStat3 (clone D3A7, Cell Signaling Technology) for 1 hour using Leica Protocol F (Molecular Imaging Core facility, MSKCC, New York, NY). Quantification of Desmin/pStat3 staining was performed using ImageJ/FIJI (NIH, Bethesda, MD). At least 19 fields at 400 \times were randomly selected and evaluated. The results were expressed as percentage of immunostained cells/over total area of tissue. To discriminate between cancer and stromal cells, fortified H&E staining was also performed (His toServ Inc). A color deconvolution algorithm was then used, with RGB vectors for the stromal component and counterstain/background stain created from regions of interest drawn from example images (Molecular Imaging Core facility, MSKCC, New York, NY). Appropriate thresholds were then set for each cell type of interest and area measurements were taken for all images. To rule out possible non CAFs/noncancer cells component, specific staining for CAFs (desmin murine CAFs) and cancer cells (human pankeratin) was also performed in serial section slides. Stroma tumor niches were evaluated as area of tissue slide with the copresence of pankeratin positive cells and stroma cells.

ALDEFLUOR assay

ALDEFLUOR analysis was performed using the ALDEFLUOR Kit (StemCell Technologies) according to the manufacturer's protocol. Cancer cells from PDX primary cultures were washed with 5 mL 10% PBS supplemented with Accumax (Innovative Cell Technologies), and single cell suspensions were first stained with anti CD133 PE conjugated antibody for 20 minutes, washed twice with PBS BSA (5%), and then incubated with ALDEFLUOR reagent.

CM preparation and phenotypic assays

CM was isolated from CAFs and cancer cell lines (10^8 cells), concentrated using Amicon Ultra 15 centrifuge tubes (Millipore), and protein levels were measured by the Lowry technique; 10 µg of total extracellular protein was loaded for zymographic/protein (MMP 2, MMP 9) and *in vitro* studies (invasion capacity). Cell growth of cocultured cancer cells with CAFs was determined with

and without anti Jagged1/Notch3 blocking antibody (AF1277, R&D Systems 500 ng every 72 hours). Briefly, luciferase positive breast cancer cells (MCF7) grown with CAFs (1:50) were seeded in 96 well plate and treated with fulvestrant (10 $\mu\text{mol/L}$ /weekly) in the presence of mouse anti Jagged1 blocking antibody. BLI signals were measured every 48 hours and growth curves were generated accordingly.

Luciferase assays

Cells were plated in 6 well plates at a density of 2×10^5 cells per well. Cells were transfected with 0.3 μg of promoter luciferase (*CD133*; ref. 15) and the activated form of *Notch3* (pNICD3 2 μg ; ref. 10). To normalize transfection efficiency, cells were also cotransfected with 0.1 μg of the pRL CMV (*Renilla* luciferase, Promega). Forty eight hours after transfection, luciferase activity was measured using the Dual Luciferase Assay Kit (Promega). Three independent experiments were performed, and the calculated means and SDs are presented.

TaqMan gene expression profile and RT PCR

qPCR was performed on 100 ng of cDNA using TaqMan precustom probes (Applied Biosystems, ER α Hs00174860 62 bp, GATA3 Hs00231122 80 bp, FOXA1 Hs0418755 59 bp, GREB1 Hs00536409 67 bp, EGR3 Hs00231780 91 bp, CCL5 Hs00174575 63 bp, PGR Hs01556702 77 bp) and SYBR Green technique (α sma forward 5' CAGGGCTGTTTCCCATCCAT 3', reverse 5' GCCATGTTCTATCGGGTACTTC 3'; SDF 1 α forward 5' CCATGAACGCCAAGGTCGTG 3', reverse 5' CCAGGTACT CCTGAATCCAC 3'; Vimentin forward 5' TGGCAGCTCTT GACCTTAAAA 3', reverse 5' GGTCATCGTGATGCTGAGAA 3'; Slug 5' AGATGCATATTC GGACCCACA 3', reverse 5' CCT CATGTTTGTGCAGGAGA 3'; CD44 forward 5' CAGCAACCC TACTGATGATGACG 3', reverse 5' GCCAAGAGGGATGCCAA GATGA 3'). ViiATM 7 Real Time PCR System was used (Applied Biosystems) in accordance with the manufacturer's instructions. For analysis, ΔC_t method was applied and fold change was calculated ($2^{-\Delta\Delta C_t}$). All values were normalized to GAPDH expression (TaqMan, Hs02758991). RT PCR for Notch3 (forward 5' TCAGGCTCTCACCTTGG 3', reverse 5' AGTCACTGGCAC GGTGTAG 3'), Jagged1 (forward 5' TCGCTGTATCTGTCCAC CTG 3', reverse 5' AGTCACTGGCACGGTTGTAG 3') and $\beta 2\mu$ as internal control 5' ACCCCCCTGAAAAAGATGA 3', reverse 5' ATCTTCAAACCTCCATGA 3' was performed in MCF7 cells control and shNotch3 and mCAF/fibroblast cell lines.

Statistical analysis

Statistical analysis was performed by SPSS (SPSS Inc.). Continuous variables were analyzed by unequal variance *t* test, paired *t* test (samples, $n = 2$), general linear model (GLM) ANOVA, or GLM for repeated measures (samples, $n > 2$). Mann Whitney and Wilcoxon tests were used to analyze ordinal variables. *P* values were adjusted for multiple comparisons according to Bonferroni correction. Association among quantitative variables was quantified by Pearson correlation coefficient. Categorical variables were analyzed by Monte Carlo χ^2 test. All the tests were two sided. $P < 0.05$ was considered significant. Elda software was used to measure the statistics of limiting dilution experiments ([<http://bioinf.wehi.edu.au/>](http://bioinf.wehi.edu.au/)software/elda/).

Results

Microvesicles from CAF mediated HT resistance

The presence of CAFs have been assessed as prognosticators in breast cancer and an "active stromal signature" in normal fibroblasts exhibits a tumor promoting phenotype (16). Many stromal secreted factors including IL6, SFD 1 α , and HFG participate in the communication between CAFs and tumor cells within the tumor microenvironment.

Stromal microvesicles have also been implicated in tumor progression in glioblastomas and ovarian cancers (17, 18). However, the molecular and pathologic relevance of CAF derived microvesicles in luminal breast cancer remains unclear.

To study tumor progression in luminal breast cancer, we established long term xenografts of highly tumorigenic MCF7 and ZR751 cells (5). Following tumor establishment (1 cm), all mice received HT (fulvestrant a selective estrogen receptor degrader commonly given to patients with ER $^+$ metastatic disease, 10 $\mu\text{mol/L}$ for 3 months. Although the majority of xenografts displayed sensitivity to HT (HTS, stable disease or remission), approximately 10% of the xenografts (data not shown) grew in the presence of therapy (Fig. 1A, HTR resistance to HT). Interestingly, the histologic analysis of these tissues revealed the enrichment of CAFs in the HTR xenografts (Fig. 1B; Supplementary Fig. S1A). Furthermore, we could isolate and *in vitro* passage CAFs from HTR derived xenograft tissues. These CAF cell lines displayed the upregulation of CAF markers by Western blot analysis and the capability of growth for multi passages (more than 20 passages; Fig. 1B and data not shown). Although we were able to isolate CAFs from HTS lesions in a small fraction of xenografts (5%, $n = 3$ out of 60), we could not propagate them in culture for more than 2 passages (2 weeks). Therefore, no CAF cell lines (0%) were established from HTS xenografts.

To further characterize these tumor associated stromal cells, we cultured HTR tumors and isolated stromal cells by FACS (negative selection with EpCAM, which recognizes epithelial cells) and determined that EpCAM $^{\text{neg}}$ cells were morphologically spindle shaped, were murine in origin (expressed murine genomic DNA, data not shown) and expressed markers of activated CAFs including Fap, vimentin, fibronectin, and activated Stat3 (Fig. 1B, phospho tyrosine 705 Stat3 pStat3).

Next, we asked whether these CAFs could promote *de novo* HTR disease. We cocultured murine CAFs and human HS27a "CAF" like cells (bone marrow derived immortalized mesenchymal cells) with HT (Luciferase $^{\text{pos}}$) naïve cancer cells in the presence/absence of HT (fulvestrant, 10 $\mu\text{mol/L}$ /weekly) and cancer cell growth was analyzed by *in vitro* bioluminescence after 2 weeks (Fig. 1C). We found that CAFs promoted tumor cell growth following HT, whereas no difference was found in the absence of therapy (Fig. 1D; Supplementary Fig. S1B). In contrast to HS27a cells and murine CAFs, normal fibroblasts (mammary and lung) did not confer resistance to HT in cocultures (Supplementary Fig. S1C).

In addition to growth factors, stromal cells have been shown to secrete microvesicles, which can horizontally transfer numerous pro-survival factors and confer resistance to radiotherapy (19). Here we determined that the number of microvesicles produced by murine CAFs and HS27a cells was much greater than MCF7 tumor cells (Fig. 1E). To determine whether these microvesicles could confer a protumorigenic advantage, we set up an *in vivo* model (Fig. 1F). MFP xenografts from HT naïve cells (MCF7) were

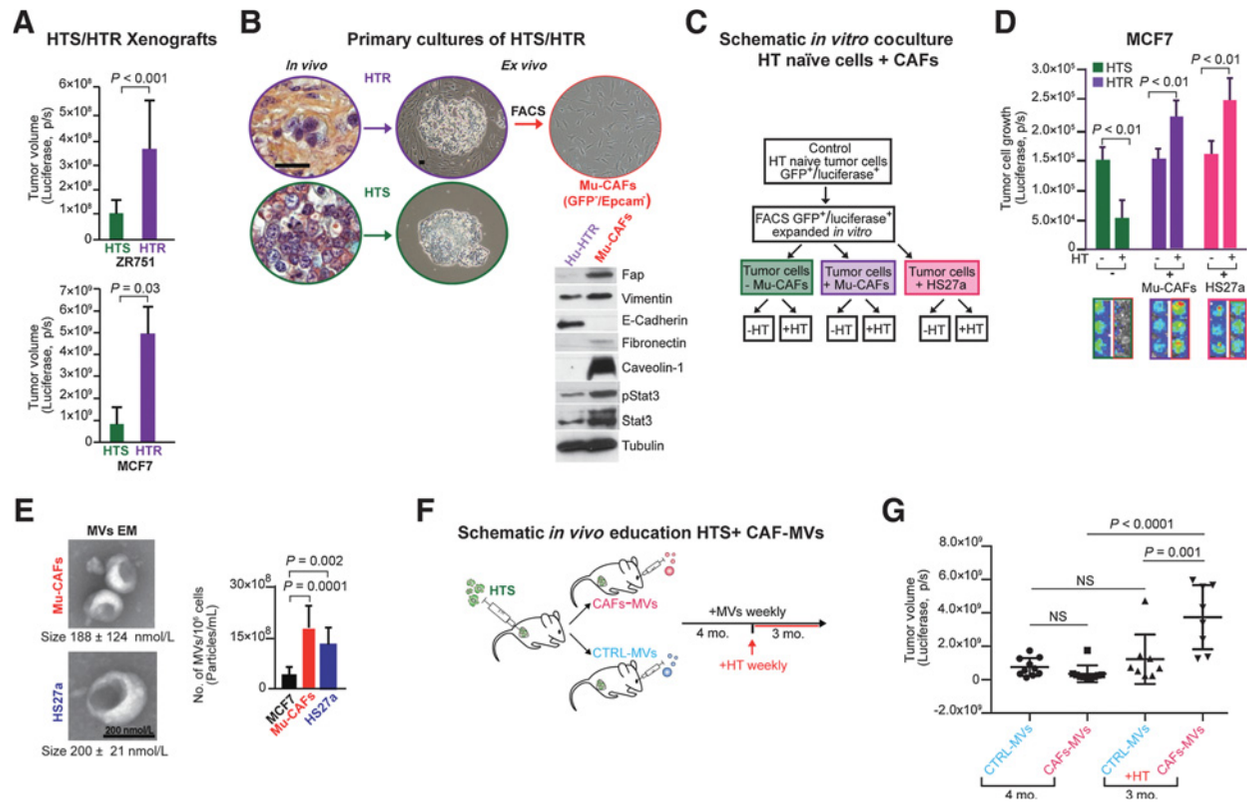


Figure 1. Microvesicles from CAFs mediated HTR. **A**, Generation of HTR (purple) and HTS (green) xenografts in luminal breast cancer. Highly tumorigenic luminal breast cancer cells (MCF7 and ZR751 luciferase positive) were injected in the MFP; xenograft bearing mice were treated with fulvestrant starting at 4 months for 3 months (HT, fulvestrant intramuscular injection 100 μ g/mouse/weekly). Some tumors grew in the presence of HT (HTR), while the majority of them was sensitive to HT (HTS). Data reported as error bars mean bioluminescence (BLI value \pm SEM) at the endpoint of the experiment (7 months, $n = 10$ mice/group). **B**, HTR cultures were enriched with murine CAFs (Mu CAFs, red). Representative fortified H&E staining of HTR xenografts (from **A**, MCF7, scale bar, 50 μ m; yellow, stroma-desmoplastic reaction), bright field images of primary cultures from HTS and HTR xenografts and Western blot analysis of FACS purified cancer cells (Hu HTR) and murine cells (Mu CAFs). **C**, Schematic of the experiment. HT naïve cancer cells (GFP⁺/luciferase⁺) were cocultured with FACS purified Mu CAFs (purple), human BMSC/CAF HS27a cells (red), or in the absence of stromal cells (green). Cells were treated with or without HT (fulvestrant, 10 μ mol/L/weekly). **D**, Proliferation potential (after 2 weeks, BLI) was determined \pm HT (fulvestrant, 10 μ mol/L). Data are reported as error bars, mean \pm SD of $n = 3$ independent experiments. *, $P < 0.05$ (Student t test). **E**, Electron microscopy images and quantification (number by nanosight) of CAF derived microvesicles (murine and human, HS27a). Scale bar, 200 nm; data reported as error bars, mean \pm SD of number of particles/mL for 10^6 cells. *, $P < 0.05$ (Student t test) of $n = 3$ independent experiments. **F**, Schematic of the experiment. Mice were injected in the MFP with HTS cells (MCF7) and subsequently injected weekly with either Mu CAFs microvesicles or control microvesicles (MCF7 derived; see Materials and Methods). After 4 months, HT was administered for 3 months (fulvestrant intramuscular injection 100 μ g/mouse/weekly). **G**, Dot plot of tumor growth of HTS cells (luciferase/p/s) grown in the MFP of mice educated with CAF MVs or CTRL MVs (MCF7) in the presence/absence of HT (fulvestrant, see **F**; microvesicles, retro orbital injection, 3×10^9 particles/mouse/weekly); error bars, mean \pm SEM of the last point of the growth curve before (4 months) and after HT (7 months). *, $P < 0.05$ values refer two way ANOVA (**G**). NS, not significant.

established; CAF MVs (3×10^9) and control microvesicles (tumor derived microvesicles, MCF7) were injected retro orbitally weekly for 7 months. Once tumors were established (after 4 months) all mice received HT (fulvestrant weekly). Although there was no difference in tumor growth before HT, those animals treated with CAF MVs had tumors resistant to HT (HTR) while MCF7 MVs provided no benefit as tumors regressed with HT (Fig. 1G). Overall, these data demonstrated that circulating stromal microvesicles can induce resistance to HT *in vivo*.

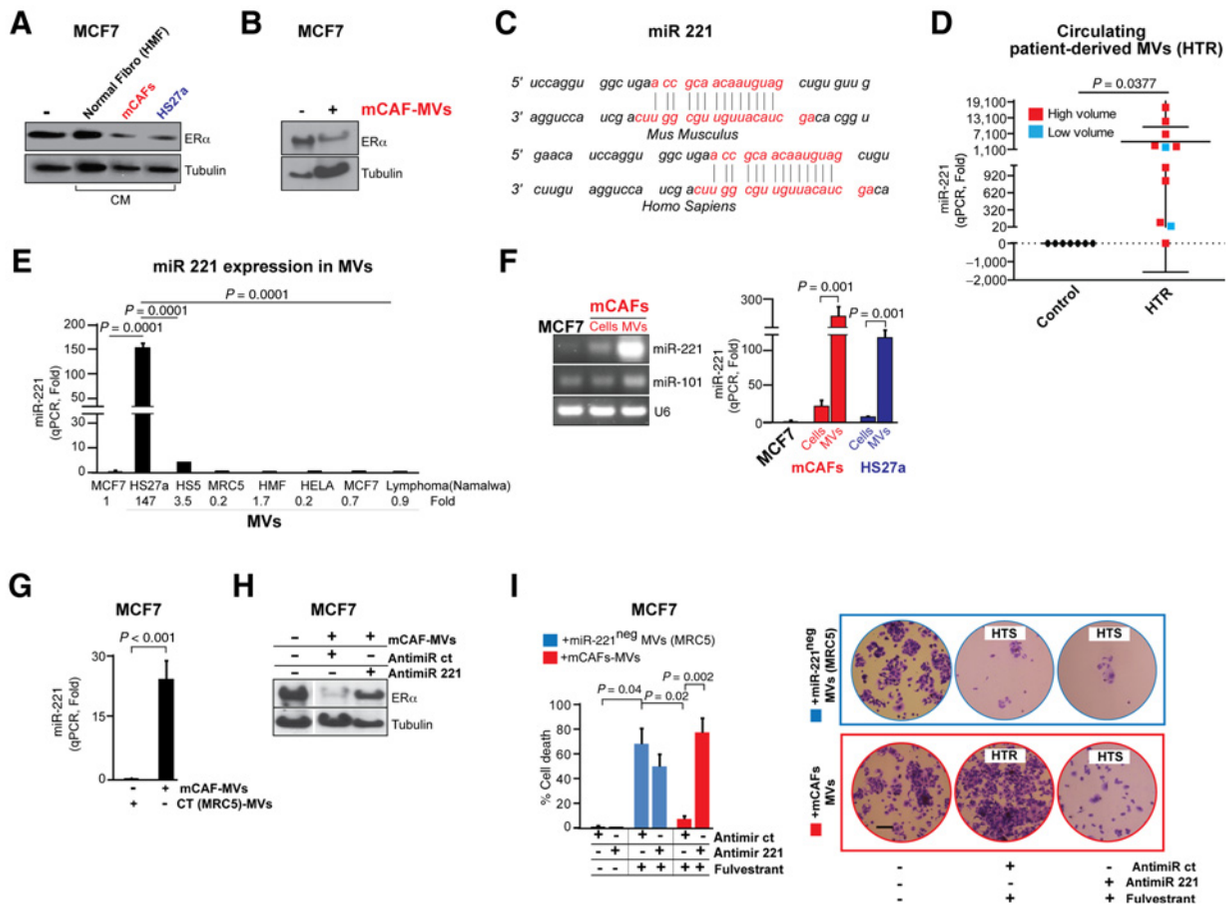
CAF derived microvesicle transfer of oncomiR 221/222 to cancer cells promotes *de novo* HT resistance

Distinct genes and pathways have been associated with resistance to HT including the activation of mutations in the *ESR1* gene (20), increased Her2 expression (21), decreased ER

levels, and ER transcriptional signatures (22, 23), increased expression of oncomiRs including the ER repressor miR 221/222 (24) and, more recently, increased Notch signaling in CSCs (5, 25).

As a reduction in ER expression is associated with resistance to HT (22), and CD133^{hi} CSCs have lower ER levels (mRNA and protein) as compared with CD133^{lo}/CD44^{lo} cells, we reasoned that the suppression of ER signaling could be a mechanism of stroma mediated expansion of therapy resistant CSCs (CD133^{hi}/Notch3^{hi}).

To test our hypothesis, we first demonstrated that the CM from CAFs (murine and human), but not normal fibroblasts, led to a decrease in ER protein expression and ER dependent transcripts (e.g., GATA3, FOXA1, GREB1, EGR3, CCL5, PGR) in MCF7 cells (Fig. 2A; Supplementary Fig. S2A). As the



Figure

2.

Microvesicle mediated OncomiR 221 transfer from CAFs to cancer cells promotes an ER^{lo} phenotype, leading to HTR resistance **A**, Western blot analysis of ER α protein in MCF7 cells following treatment with the CM of human normal mammary fibroblasts (HMF), mCAFs (red font), and HS27a cells (blue font, 48 hours). **B**, Western blot of ER α protein in MCF7 cells treated with and without mCAF derived microvesicles (10⁸ particles, 48 hours). **C**, Image showing the OncomiR 221 sequence conservation in *Mus Musculus* and *Homo Sapiens*. **D**, MV miR221 expression as determined by qPCR as fold increase from patients with HTR disease and healthy controls (reference MCF7 MVs was used and normalized to total RNA expression, $n = 3$ replicates; Supplementary Table S1). Patients with high volume disease (>10% of organ involvement) are denoted in red and low volume disease (<1% of organ involvement) is denoted in blue. **E**, Bar graphs representing oncomiR 221 expression (qPCR) in microvesicles from indicated sources (10⁸ cells). Data are reported as fold increase (221/U6 expression) \pm SD of $n = 3$ replicates (MV MCF7 is used as reference). **F**, RT-PCR analysis of miR 221 and 101 in MCF7, mCAFs, and mCAF derived microvesicles (RNA was isolated from 10¹⁵ particles) and bar graph showing oncomiR 221 expression (qPCR, Fold, normalized to U6 values) in mCAFs and HS27a cells and their respective microvesicles. The expression of oncomiR 221 in MCF7 was used as a reference control. **G**, Bar graph of oncomiR 221 (qPCR, Fold) in MCF7 cells following chronic mCAF MV education (one month, 10⁸ particles weekly). Microvesicles were isolated from MRC5 cells (normal lung fibroblasts) and used as control microvesicles (CT MVs). **H**, Western blot analysis of ER α protein in MCF7 cells treated with mCAF MVs (10⁸ particles, 48 hours) previously transfected (24 hours before microvesicle education) with the anti miR 221 or control (CT; see Materials and Methods); **I**, Bar graph representing cell death (by Trypan blue, %) of MCF7 cells transfected with anti miR 221 and controls, treated with fulvestrant (10 μ mol/L, 7 days) in the presence/absence of mCAF and MRC5 microvesicles (10⁸ particles every 48 hours). miR 221 negative microvesicles were used as control microvesicles and were obtained from normal lung fibroblasts (MRC5). Representative images of Crystal violet staining of MCF7 cells at the endpoint of the experiment described in **H**. Data are reported as mean \pm SD of three independent experiments ($n = 3$). NS, not significant. P values, Student t test (**D-G, I**).

suppression of ER protein occurred with CM from both murine and human CAFs, we hypothesized that rather than soluble factors (which can be typically species specific; refs. 26, 27), CAF derived miRNAs might be mediating the downregulation of ER expression. Microvesicles have been suggested to be mediators of nucleic acid transfer including miRNAs (27). Among different miRNAs, the forced overexpression of oncomiR 221/222 in luminal breast cancer cells has been found to reduce ER expression and promote HT resistance (24). In addition, increased plasma levels of miR 221 were found in ER negative breast cancer patients (28). The administration of 10⁸ microvesicles from mCAFs to ER⁺

cancer cells (MCF7) reduced ER levels after 48 hours (Fig. 2B). We showed that oncomiR 221/222 sequences are conserved between human and mouse species, suggesting possible functional cross species interactions (Fig. 2C). Importantly, miR 221 expression was found in circulating microvesicles from patients with HTR metastatic disease (independent of tumor burden) as compared with healthy controls (Fig. 2D, $n = 11$; Supplementary Table S1).

We determined that CAF derived microvesicles were enriched for miR 221 compared with normal fibroblasts and distinct cancer cells lines (breast, cervical, and lymphoma),

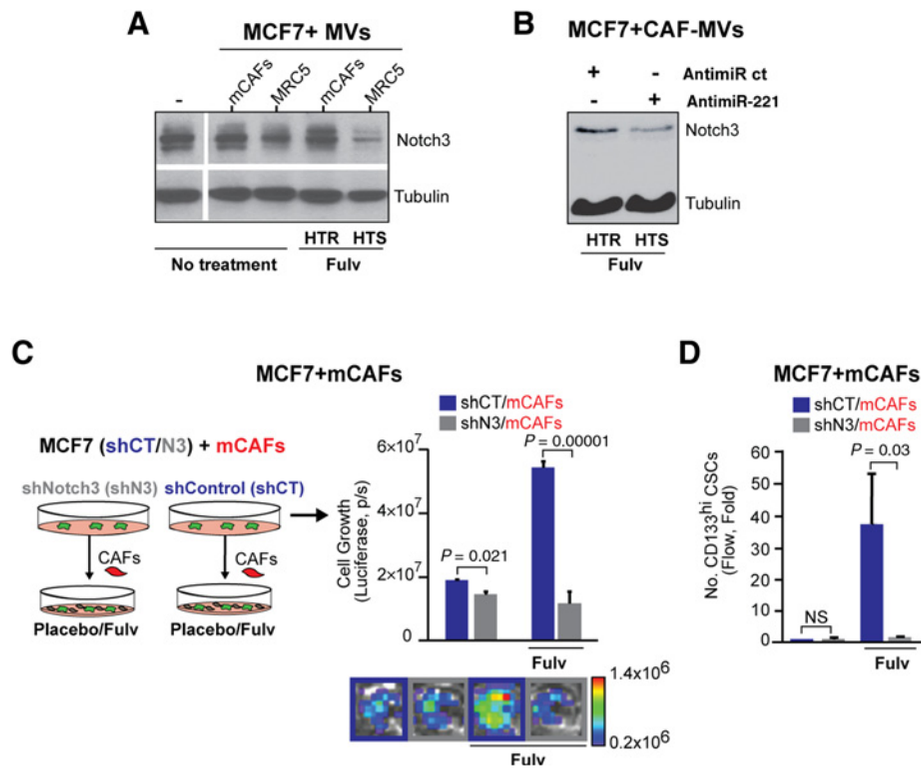


Figure 3.

CAF mediated expansion of CD133^{hi} CSCs via an OncomiR 221^{hi}/ER^{lo}/Notch3^{hi} loop. **A**, Western blot analysis of Notch3 protein from MCF7 cells in the presence/absence of mCAF and MRC5 microvesicles (10⁸ particles every 48 hours) ± fulvestrant (Fulv; 10 μmol/L for 7 days). Tubulin was used as a loading control. **B**, Representative Western blot analysis of Notch3 protein from MCF7 cells transfected with the anti miR 221/CT (1 μg/well) and treated with fulvestrant and mCAF MVs (48 hours). Tubulin was used as a loading control. **C**, Schematic of coculture experiments. Control (shCT) and Notch3 knocked down MCF7 GFP/Luciferase⁺ cells (shN3) were cultured with PDX derived mCAFs (1:10; mCAFs:cancer cells) in the presence/absence of fulvestrant (Fulv, 10 μmol/L, weekly administration for a month); growth of MCF7 shCT/shN3 cells in cocultures was then determined. Data are reported as mean (Luciferase, p/s photons/seconds) ± SEM of the last time point of the growth curve or 28 days (three biological replicates with three technical replicates each). **D**, Fold Increase in CD133^{hi} CSCs (flow analysis) from MCF7 shCT/shN3 cell cocultures treated with fulvestrant (1 month, endpoint, **C**). Data are reported as total number of CD133 positive cells (mean ± SD of fold change) of *n* = 3 different specimens (as a reference, total numbers of CD133^{hi} cells from 10⁹ naïve MCF7 cells were used). NS, not significant. *P* values, Student *t* test (**C**, **D**).

suggesting that stroma derived microvesicle could account for the increased level of miR 221 expression in microvesicles (Fig. 2E). Accordingly, miR 221, but not a control miRNA (miR 101), was highly expressed (100 to 200 fold enrichment) in microvesicles from CAFs as compared with micro vesicles from normal fibroblasts, CAFs and cancer derived cell lines (Fig. 2F; Supplementary Fig. S2B). Administration of murine CAF MVs compared with normal fibroblast MVs (MRC5) to MCF7 cells (oncomiR 221 negative) led to the transfer of oncomiR 221 (Fig. 2G, 20 fold increase in expression in educated cancer cells).

To address the consequences of microvesicle derived oncomiR 221 transfer in mediating HT resistance, anti miR 221 was transfected in cancer cells before microvesicle administration. We demonstrated that anti miR 221 transfection abrogated mCAF MV dependent downregulation of ER protein (Fig. 2H) and HT resistance following chronic mCAF MV education (Fig. 2I). Restored sensitivity to HT in anti miR 221 transfected cells was associated with an increase in ER expression/activity (Supplementary Fig. S2C and S2D). These data suggest that CAF microvesicle mediated HT resistance occurs via the transfer of onco miR 221 promoting an ER^{lo} phenotype.

CAF mediated expansion of CD133^{hi} CSCs via an oncomiR 221^{hi}/ER^{lo}/Notch3^{hi} loop

Increased expression of Notch and downstream signaling events as well as a higher number of CSCs are found in HT resistant breast cancer (29, 30). Among Notch proteins, Notch3 and Notch4 are crucial mediators of resistance to HT in distinct models of luminal breast cancers (5, 25, 31, 32).

Given the pivotal role of CAF derived microvesicle in promoting the switch from sensitive to resistant disease (HTS to HTR), we asked whether these stromal microvesicles could also modulate Notch3 expression. We cultured MCF7 cells with fulvestrant (10 μmol/L/weekly) for 2 weeks (see schematic Fig. 1C) with microvesicles (10⁸ particles/weekly) from either MRC5 or mCAFs. Although no effect was observed in the absence of HT (no treatment), mCAF MVs restored Notch3 expression and activity (Hes1, Hey 1 mRNAs) following fulvestrant, which associated with increased growth (Fig. 3A; Supplementary Fig. S3A, HTR cells see Fig. 2I). Conversely, MRC5 MVs administered cells did not overcome ER dependent downregulation of Notch3 expression and activity (Hes1, Hey 1 mRNAs) as well as the suppression of growth (Fig. 3A, for HTS cells, see Fig. 2I). The transfection of anti-miR 221 led to decreased Notch3 protein levels and restored

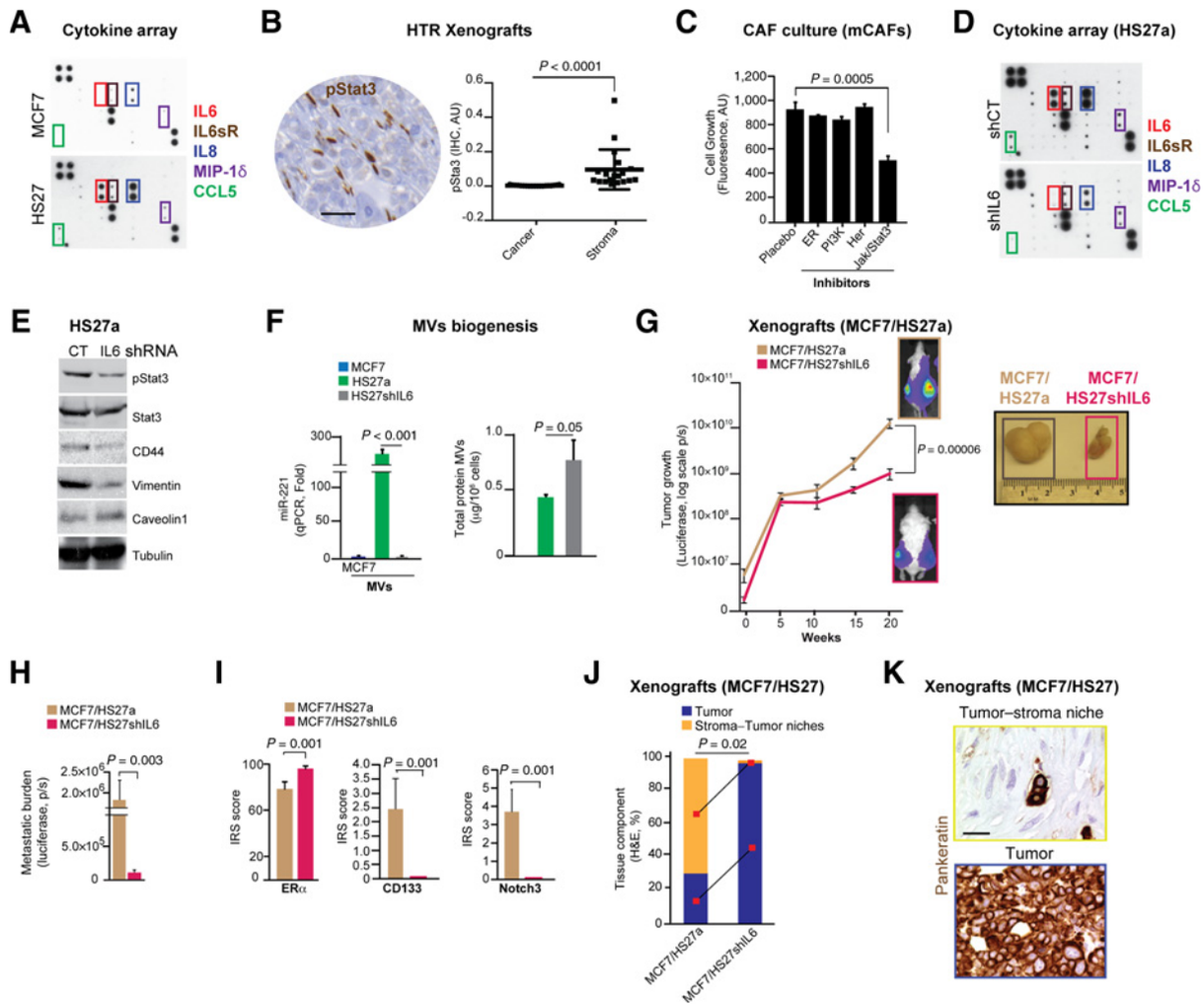


Figure 4.

IL6/Stat3 signaling from CAFs promotes the expansion of CD133^{hi} CSCs. **A**, Cytokine array expression of the CM from MCF7 and HS27a cells (10 µg total protein). Highlighted are overexpressed cytokines and chemokines (IL6, IL6sR, IL8, MIP 1δ, and CCL5). **B**, Dot plot showing phospho tyrosine 705 Stat3 (pStat3) IHC quantification in HTR derived primary tumor tissues (see Fig. 1B) in both the stroma and tumor compartments. Quantification of pStat3 was performed using ImageJ/FIJI (NIH) of $n = 19$ fields from $n = 5$ different tumors. The results were expressed as ratio of pStat3 IHC value/total tissue area. A representative IHC image is shown. Scale bar, 25 µm. **C**, Bar graph of the proliferation capacity (Calcein AM, fluorescence) of xenograft derived mCAFs isolated from HTR xenografts (Fig. 1) and cultured in the presence of vehicle (placebo) or signaling pathway inhibitors including ER (fulvestrant, 10 µmol/L) or PI3K (BYL, 100 nmol/L), HER (lapatinib, 100 nmol/L) or JAK/pStat3 (AZD1480, 500 nmol/L). Data are reported as mean (fluorescence) ± SEM of the last time point of the growth curve (14 days; three biological replicates with three technical replicates each). **D**, Cytokine array expression from the CM (10 µg) of HS27shIL6 versus HS27a cells. Highlighted are overexpressed cytokines and chemokines (IL6, IL6sR, IL8, MIP 1δ, and CCL5). **E**, Western blot analysis of pStat3, Stat3, CD44, vimentin, caveolin1, and tubulin protein levels in HS27a cells CT and shIL6. **F**, Bar graph of OncomiR 221 expression in microvesicles derived from 10⁸ HS27a and HS27shIL6 cells (qPCR, fold as reference, MCF7 microvesicles were used, normalized to U6 expression). Bar graph of protein levels (µg) in 10⁸ microvesicles isolated from HS27a and HS27shIL6 cells is also shown. **G** and **H**, Tumor growth and metastatic burden (luciferase, BLI) in MCF7/HS27a and MCF7/HS27shIL6 xenografts. 10³ cancer cells were inoculated in both 4th inguinal MFP with 10² HS27a cells (CT or shIL6). Tumor growth was examined over 20 weeks. Data are reported as mean BLI value ± SEM (log scale) for each time point ($n = 4/5$ mice/group). Metastatic burden is mean BLI value ± SEM of signal from metastatic tissues including lymph nodes, lungs, and bones. A representative image of primary tumors from MCF7/HS27a versus MCF7/HS27shIL6 is shown. **I**, Bar graph representing immunohistochemical quantification (IRS score) of ERα, CD133, and Notch3 in tumor derived tissues from MCF7/HS27a or MCF7/HS27shIL6 xenografts (**G**). Data are reported as mean ± SD of $n = 10$ tissue sections for each group. **J**, Bar graph showing percentage (H&E, %) of stroma tumor niches versus tumor compartment in primary tumor tissues derived from MCF7/HS27a and MCF7/HS27shIL6 xenografts (**G**). **K**, Representative Pankeratin staining of tissues slides from **G** depicting stroma tumor niches (cancer cells surrounded by stromal cells). Data are reported as mean ± SD of $n = 10$ tissue slides for each group. *P* values, *t* test (**B, F, H, I**), Wilcoxon two sample test (**J**), multiple comparisons corrected *post hoc t* test after GLM ANOVA (**C**), repeated measures GLM ANOVA (**G**).

sensitivity to HT (fulvestrant) of CAF MV treated cancer cells (Figs. 2I and 3B). These data demonstrate that miR 221 in CAF MVs can block HT mediated downregulation of Notch3 expression.

We recently described the enrichment of CD133^{hi}/ER^{lo}/Notch3^{hi} CSCs in HT resistant tumors, which also expressed high levels of Notch regulated genes such as *Hey1* and *Hes1* (GSE69280; ref. 5).

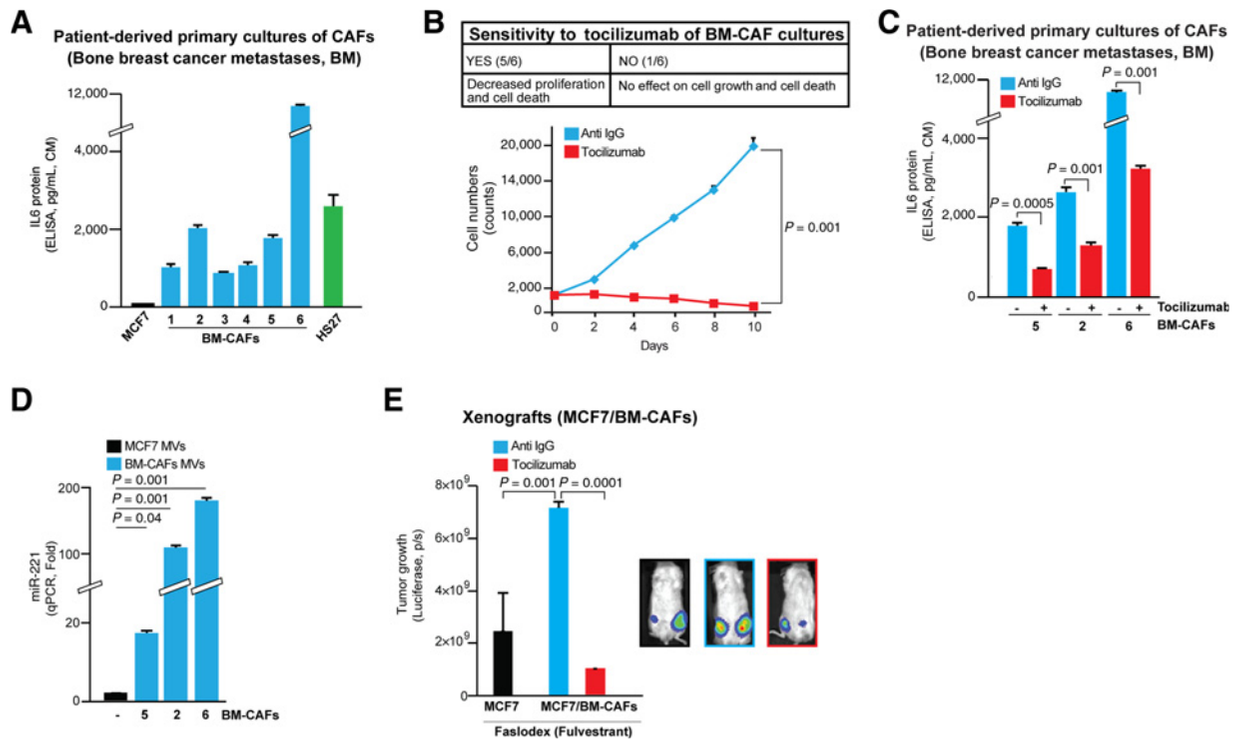


Figure 5.

Autocrine CAF derived IL6 triggers endocrine resistant disease. **A**, Bar graph showing IL6 levels measured by ELISA from the CM (10 μ g total protein) of primary CAF cultures (BM CAFs) isolated from patient bone derived breast cancer metastases (Supplementary Table S2). IL6 levels from MCF7 and HS27a are also shown. **B**, Growth (cell number) of BM CAFs (specimen 5, Supplementary Table S2) treated with an anti IL6 IL6R antibody tocilizumab (500 ng/mL every 48 hours) versus IgG control. **C**, Bar graph of IL6 protein (ELISA) from the CM (2 μ g total protein) of BM CAFs treated with tocilizumab or IgG for a week (500 ng/mL every 48 hours). **D**, Bar graph of oncomiR 221 expression by qPCR (Fold, normalized on U6 level) in microvesicles isolated from BM CAF cultures. Microvesicles isolated from MCF7 cells were used as a reference control. **E**, Tumor burden (luciferase) of MCF7 xenografts alone (black) and coinjected with BM CAFs (specimen 6, Supplementary Table S2) treated with fulvestrant and tocilizumab. Briefly, MCF7 xenografts were established in the MFP of NOD/SCID mice alone (10^3 cells) or in combination with CAFs (10^2 cells). When tumors reached \sim 1 cm (3 months), mice were randomized ($n = 4$ /group) to receive either fulvestrant (1 mg/weekly) or fulvestrant and tocilizumab (100 μ g/g/mouse) for 4 months. Data are reported as mean BLI value \pm SEM at the endpoint of the experiment (7 months). Data are reported as mean \pm SD of three independent experiments ($n = 3$). *P* values, Student *t* test (**C**, **D**, **E**), multiple comparisons corrected *post hoc t* test after repeated measures GLM ANOVA (**B**).

Given the role of CAF or stromal microvesicles in promoting a miR 221^{hi}/ER^{lo}/Notch3^{hi} phenotype, we tested the hypothesis that CAFs could promote the *in vivo* expansion of CD133^{hi} cells via Notch3 upregulation. The selective reduction of Notch3 expression in cancer cells (shNotch3) and activity (using an anti Jagged1 blocking antibody) abrogated CAF mediated HT resistance and the expansion of CD133^{hi} cancer cells (Fig. 3C and D; Supplementary Fig. S3B). In agreement with the knockdown experiment, overexpression of the activated form of Notch3 (pNICD3) in MCF7 cells led to an increase in CD133 promoter luciferase activity (pCD133) with HT (fulvestrant) in association with a reduction in ER protein levels (Supplementary Fig. S3C and S3D). These data suggest that higher Notch3 activation could promote a feed forward ER^{lo}/CD133^{hi} loop necessary for the generation of CD133^{hi} CSCs (Supplementary Fig. S3C and S3D). Overall, our data describe a novel mechanism of HT resistance: CAF MV mediated transfer of the onco miR 221 leading to reduced ER expression and Notch3 upregulation.

IL6/Stat3 activity is required for CAF CSC niche formation

As the biogenesis of oncomiR 221/222^{hi} microvesicles occurs preferentially in CAFs (not normal fibroblasts), we

hypothesized that the abrogation of a CAF phenotype would interfere with the generation of HT resistant CSCs. To investigate this hypothesis, we examined possible candidates responsible for CAF growth. Compared with breast cancer cells, the CM of CAFs (HS27a cells) expressed higher levels of chemokines (e.g., IL8, MIP 1 δ , CCL5) and cytokines, including IL6 an activator of Stat3 (Fig. 4A). These findings were further supported by evidence of high pStat3 levels in murine CAFs from HTR derived xenografts (Fig. 4B). Differently from other signaling pathways (HER, PI3K, ER), pStat3 activity was required for CAF proliferation as well as the generation of oncomiR 221^{hi} microvesicles (Fig. 4C; Supplementary Fig. S4A). In concordance with these data, reduced IL6 expression in HS27a cells (using an IL6 shRNA) lowered secreted IL8/IL6R/CCL5 levels as well as the expression of CAF markers including pStat3, vimentin, and CD44 (Fig. 4D and E). In addition, compared with HS27shCT cells, HS27shIL6 cells had reduced proliferative and invasive potential (Supplementary Fig. S4B and S4C) as well as lowered MMP2/9 expression and activity, indicating a loss of characteristic CAF features (Supplementary Fig. S4D). Moreover, microvesicles from HS27shIL6 cells had essentially no expression of oncomiR 221 compared with

HS27shCT cells with no change in microvesicle production (Fig. 4F, protein content as a surrogate marker of microvesicle yield). These data suggest that IL6/pStat3 signaling is crucial for the proliferation of CAFs and the production of oncomiR 221⁺ microvesicles.

To address the phenotypic consequences of decreasing IL6 signaling in CAFs, we coinjected MCF7 cells with HS27shCT and HS27shIL6 CAFs into the MFP of mice. Compared with controls (MCF7/HS27a), the coinjection of MCF7/HS27shIL6 cells resulted in impaired tumor growth, lower metastatic burden, and decreased expression of CD133^{hi}/Notch3^{hi}/ER α ^{lo} CSCs (Fig. 4G-I; Supplementary Fig. S4E). In agreement with the loss of CD133^{hi}/Notch3^{hi} CSCs, MCF7/HS27shIL6 derived tumors had fewer stroma tumor niches (Fig. 4J and K) and decreased pStat3 expression (Supplementary Fig. S4F). Overall, our data suggest that IL6 mediated generation of stromal niches is required for the expansion of CD133^{hi} CSCs.

Anti IL6 therapy abrogates CAF mediated *de novo* resistance to HT

To extend our results to clinical specimens, we established primary cultures of stromal cells from patient derived bone metastases (Supplementary Table S2, BM CAFs). IL6 is a well known pleiotropic cytokine, secreted at high levels from the bone marrow microenvironment and CAFs (33, 34). We isolated and cultured CAFs from bone metastases (Supplementary Fig. S5A and data not shown) and detected high levels of IL6 protein from the CM of these primary cultures (Fig. 5A). These levels are similar to those found from the CM of HS27a cells. Consistent with the HS27 model, the abrogation of IL6 signaling, using the anti IL6R IL6 antibody (tocilizumab), abrogated the growth in 70% of CAF primary cultures (Fig. 5B), reduced IL6 secretion and the expression of CAF markers (Fig. 5C; Supplementary Fig. S5B), this finding suggests that autocrine IL6 maintains the CAF phenotype of these cells.

Although miR 221/222 expression is very low in luminal breast cancer tissues and cells (Fig. 2), high levels of miR 221 was found in circulating microvesicles from HTR patients (Fig. 2D). Next, we reasoned that, in agreement with other investigators, CAFs would be the major source of miR 221 (35). We then isolated microvesicles from CAF primary cultures derived from bone metastases and found increased levels of oncomiR 221 as compared with tumor microvesicles (Fig. 5D); these results were similar to those found in HS27a cells and mCAF (Fig. 4). We subsequently demonstrated that the coinjection of BM CAFs with MCF7 cells promoted HTR tumor growth (treated with fulvestrant) in an IL6 dependent manner, as combination fulvestrant/tocilizumab abrogated tumor growth (Fig. 5E).

CAF mediated expansion of CD133^{hi} CSCs

We and others have recently demonstrated the enrichment of CSCs expressing CD133 or ALDH^{hi} activity in experimental luminal breast cancers following HT (5, 25). In addition, expression of Prominin1 (CD133) was identified in tumors from patients who progressed on adjuvant HT (23). In agreement with other investigators, CD133^{hi} and CD44^{hi} cells are functionally CSCs as they were capable of engrafting with low cell numbers (<1,000, Supplementary Fig. S6A). We previously demonstrated that differently from CD44^{hi} cells, CD133^{hi}/CD44^{lo} cells expressed embryonic stem cell signatures and increased the expression of ABCG2, a CSC gene associated with therapy resis-

tance (GSE69280; refs. 5, 36). Here we determined that CD133^{hi}/CD44^{lo} cells expressed normal stem cell signatures by GSEA (Supplementary Fig. S6B, GSE69280; ref. 5). In addition, compared with CD133^{lo}/CD44^{lo} and CD44^{hi}/CD133^{lo} cells, the injection of CD133^{hi}/CD44^{lo} cells gave rise to slow growing tumors (Supplementary Fig. S6C) with an increased capacity to disseminate to the bone marrow (Supplementary Fig. S6D). These data suggest that the CD133^{hi} phenotype is a distinct CSC population (37).

Next, we investigated the hypothesis that extrinsic (stromal heterogeneity) factors could regulate the evolution of therapy resistant niches leading to metastatic progression (38, 39). We demonstrated high PROM1 (encoding CD133) expression by microarray (GSE17705) was associated with increased levels of a mammary CAF gene signature (12) in the setting of human HT resistant primary tumors (Fig. 6A, $P < 1.82 \times 10^{-4}$; Supplementary Fig. S7A).

To examine the importance of CD133^{hi} cells in clinically relevant models of metastatic breast cancer, we generated PDXs from HT resistant (HTR) luminal breast cancer bone metastases (Fig. 6B; Supplementary Table S2). Cultured tumor cells were serially transplanted into the MFP of immunocompromised mice in the presence of HT (fulvestrant: a selective estrogen receptor degrader). After three sequential passages, the enhanced tumorigenic capacity (% tumor take) of PDXs was associated with increased Notch3 expression (Supplementary Fig. S7B), enrichment of CD133^{hi} (~40 fold) cancer cells and murine stromal cells (~30 fold), which had infiltrated the tumor (Fig. 6B and C; Supplementary Fig. S7C and S7D). We further determined that the PDX derived CD133^{hi} cells had low ALDH activity, suggesting different and unique stem cell populations arising through a stroma mediated metastatic transition within ER⁺ breast cancer (Supplementary Fig. S7E).

Given the coenrichment of CD133^{hi} cells with CAFs, we hypothesized that CAFs could directly promote the expansion of HT resistant CD133^{hi} cells. To address this hypothesis, we depleted HT resistant PDX primary cultures from CAFs (by FACS for EpCAM^{pos}) and maintained these cancer cells in culture for several months. When long term depleted for murine CAFs, EpCAM^{pos} cells from PDX primary cultures were growth inhibited by HT as compared with tumor cells freshly isolated from cocultures (sorting for EpCAM^{pos}, Fig. 6D and E). Moreover, acquired sensitivity to HT resulted in decreased numbers of self renewing CD133^{hi} cells (Fig. 6E and F).

To determine whether CAFs could confer *de novo* HT resistance and promote the biogenesis of CD133^{hi} cells, we cultured HT sensitive MCF7 cells with murine CAFs, isolated from xenografts, with/without HT (fulvestrant) for 4 weeks (Fig. 6G). When cultured with CAFs, MCF7 cancer cells became resistant to HT (fulvestrant), as compared with MCF7 cells alone (Fig. 6G). Concomitant with HTR cancer cell growth, by FACS we observed a marked (30 fold) increase in CD133^{hi} CSCs as compared with MCF7 cells alone (Fig. 6H). Further characterization of the CD133^{hi} cells revealed that ER α mRNA levels were lower in these cells than in CD133^{lo}/CD44^{lo} cells (Fig. 6I). In agreement with these data, higher CD133/PROM1 expression was associated with decreased ESR1 mRNA expression and activity in HT resistant primary tumors (Fig. 6J, $P < 0.0001$). Overall, these findings demonstrate that CAFs, in the presence of HT, can promote the *de novo* formation of CD133^{hi} CSCs and HTR disease.

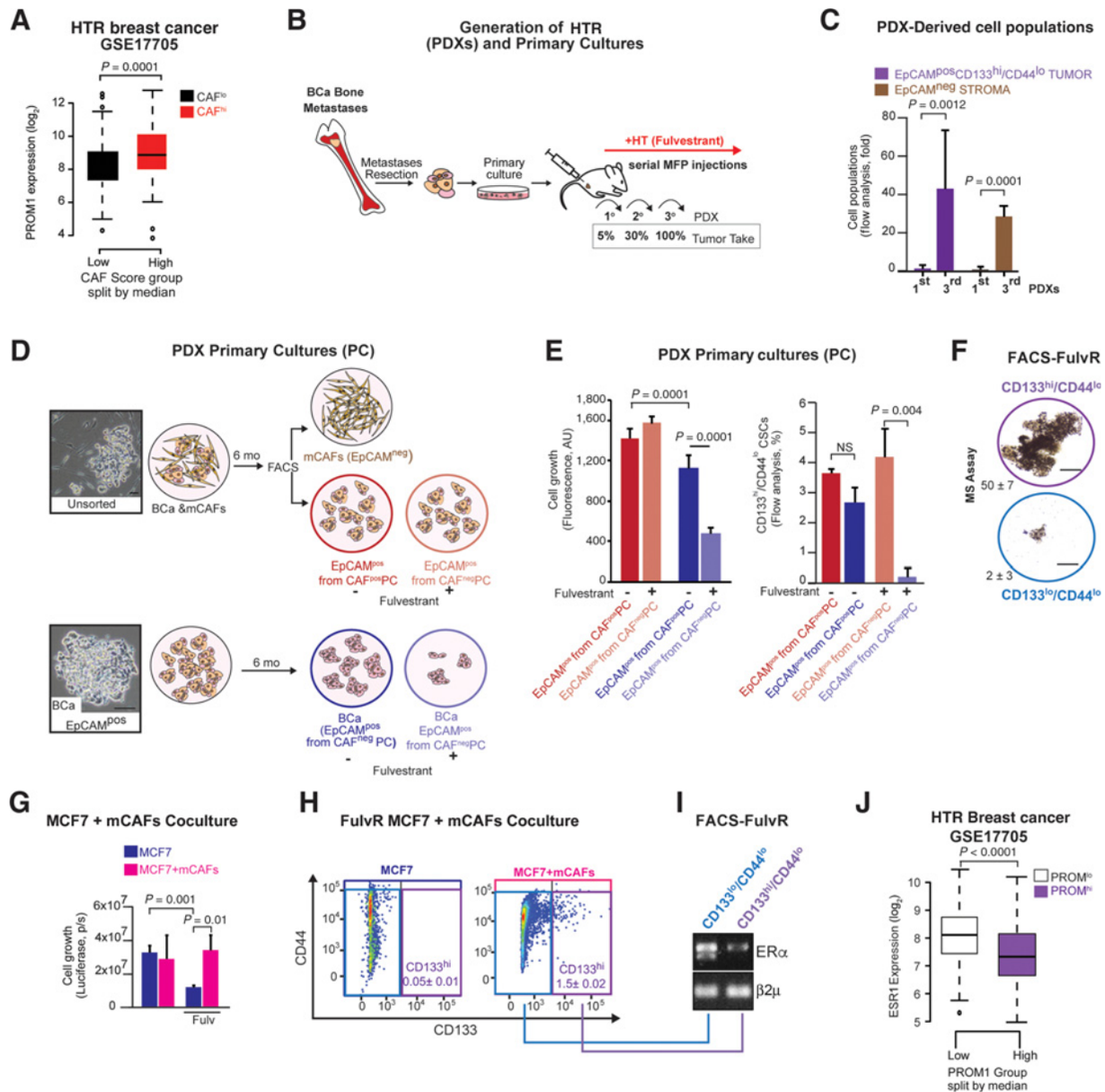
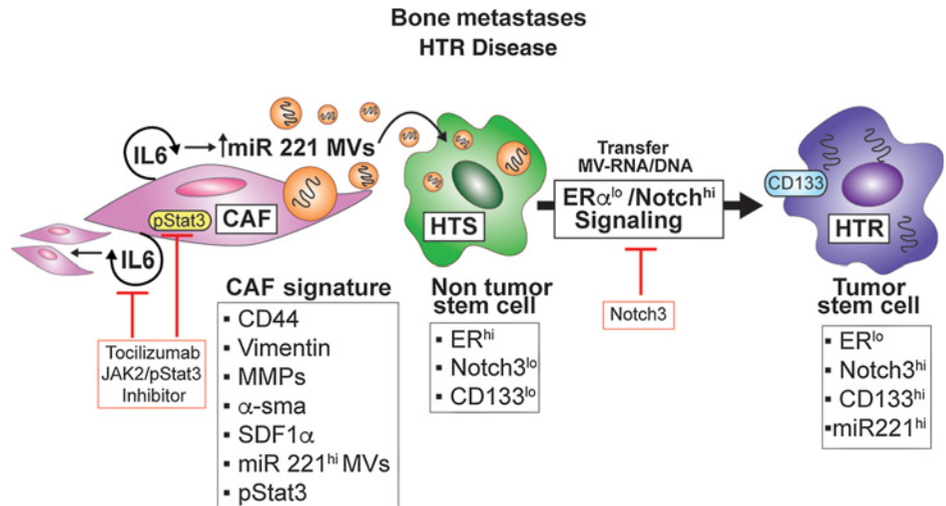


Figure 6.

CAF mediated expansion of CD133^{hi} CSCs in patient derived xenografts from HTR breast cancer bone metastases. **A**, Box plot of PROM1 expression (log₂) separated into "low" (black) and "high" (red) groups by the median CAF score as described in Materials and Methods using the GSE17705 dataset (23). Box plots are drawn such that horizontal lines indicate the sample median, the box spanning the interquartile distance (IQD), whiskers extending to 1.5× IQD, and remaining outlying points shown as open circles. **B**, Tumor take percentage (%) of PDXs following three sequential passages *in vivo*. Briefly, cancer cells were isolated from bone metastases (Supplementary Table S2, sample shown B Met3), grown in culture dishes for 2 weeks and then injected into the MFP of NOD/SCID mice (1st *n* = 1/5; 2nd *n* = 3/10; 3rd *n* = 40/40) treated with HT (fulvestrant, 1 mg/weekly). After 5 months, tumor tissues were digested and cancer cells were cultured *in vitro* and FACS sorted for EpCAM positivity (to eliminate murine cells) before reinoculating (10⁵ EpCAM^{pos} cells) these into new cohorts of mice. **C**, Number of CD133^{hi}/CD44^{lo} cancer cells (EpCAM^{pos}) and stroma cells (EpCAM^{neg}) derived from 1st and 3rd generation PDXs (flow analysis, fold). **D**, Schematic of the *in vitro* PDX culture established in presence/absence of CAFs (CAF^{pos} or CAF^{neg}). Unsorted (cancer cells and CAFs) or previously FACS purified EpCAM^{pos} cancer cells from PDX primary cultures were grown for 6 months (6 mo), then EpCAM^{pos} cancer cells were FACS purified from EpCAM^{neg} cells and treated with fulvestrant (Fulv, 10 μmol/L, 14 days). Scale bar, 10 μm. **E**, Bar graphs of cell growth (Calcein AM, left panel). Data are reported as mean (fluorescence) ± SEM after 14 days (three biological replicates with three technical replicates each). The number of CD133^{hi} cells (flow analysis, %, right) is also shown after 14 days of culturing. **F**, Images and quantification of secondary MS growth of FACS purified CD133^{hi} and CD133^{lo} cancer cells from experiment **E**. Scale bar, 100 μm. **G**, Schematic and growth of MCF7 (GFP⁺/luciferase⁺) cells, using BLI luciferase grown with and without mCAFs (PDX derived). A 1:10 mCAF to MCF7 cell ratio in the presence/absence of fulvestrant (Fulv, 10 μmol/L, added weekly for 30 days). **H**, Representative CD133/CD44 expression by flow analysis of fulvestrant resistant MCF7 cancer cells (FulvR) in the presence/absence of mCAFs at the endpoint of the experiment described in **G**. **I**, RT-PCR analysis of ERα in FACS isolated CD133^{hi} and CD133^{lo} cancer cells from experiment panels **G** and **H**. β2μ used as a loading control is shown. **J**, Box plot depicting ESR1 expression (log₂) separated into "low" (white) and "high" (purple) groups by the median PROM1 expression using the GSE17705 dataset (23). Box plots are drawn with a horizontal line indicating the sample median, box spanning IQD, whiskers extending to 1.5× IQD, and outlying points shown as open circles. Data are reported as mean ± SD of three independent experiments (*n* = 3). *P* values, Monte Carlo χ² test (**A**), Student *t* test (**B**, **C**, **E**, **J**), *t* test after GLM ANOVA (**G**), GLM ANOVA for repeated measures (**E**, left).

Figure 7.

Stroma microvesicle mediated CSC evolution in endocrine resistant metastatic breast cancer. Schematic of the proposed model of CAF mediated endocrine resistant disease in luminal breast cancer. Autocrine IL6/Stat3 signaling drives the proliferation of CAFs and the biogenesis of oncomiR 221/222^{hi} microvesicles; these microvesicles are taken up by ER^{pos} breast cancer cells and in conjunction with HT leads to the potent suppression of ER signaling promoting Notch3 upregulation, which in turn sustains the self renewal of CD133^{hi} CSCs in an ER independent manner.



In summary, our data led us to propose the following model: IL6 upregulation drives a CAF phenotype leading to the generation of stroma tumor niches *in vivo*. The presence of active CAFs promotes a skewing of the cancer cell population toward a CD133^{hi}/Notch3^{hi}/ER^{lo} CSC phenotype. This occurs via the production/action of CAF derived microvesicle, which reduces ER activity leading to HTR disease (Fig. 7; ref. 40).

Discussion

Significant progress has been made in identifying tumor cell specific factors (e.g., tumor secretome and gene signatures) that promote cancer cell survival and proliferation in the bone microenvironment (27, 41). In addition, cross talk between bone marrow derived myeloid cells, osteoclasts, and osteoblasts with tumor cells has identified many growth factors, chemokines, and miRNAs as critical regulators of bone metastasis (42). A recent manuscript by Luo and colleagues has also demonstrated that increased osteoblast derived IL6 promotes tumor cell seeding and bone metastases in a model of breast cancer cells injected in the arterial circulation (43). IL6 is a pleiotropic cytokine, which is secreted from several cell types, including CAFs and plays a crucial role in the expansion of cancer stem cells (44, 45) as well as in the proliferation potential of CAFs (46, 47).

The presence of CAFs have been assessed as a poor prognostic feature in breast cancer (48) and an "active stromal signature" in normal fibroblasts exhibits a tumor promoting phenotype (16). Moreover, PDX tumor tissue was shown to be enriched with host CAFs (49); but the molecular and pathologic relevance of any given CAF^{hi} phenotype in PDXs remains unclear.

The majority of breast cancers (~70%) are of the ER positive or luminal subtype. Although, the suppression of ER activity with HTs has led to improved survival, when these cancers recur they preferentially metastasize to the bone, and eventually acquire resistance to HT and are thus incurable (50, 51).

The upregulation of distinct pathways, including Her2, PI3K, and/or overactive estrogen signaling, are found in metastatic cancer cells escaping tumor dormancy from adjuvant HT. Although targeting these pathways in the metastatic setting leads to clinical responses (52), resistance to anti Her2/estrogens/PI3K regimens invariably occurs (27, 53). We and others have reported the clinical relevance of high numbers of CD133^{hi} cells in "ther

apy" resistant cancers including lung cancer and HT resistant luminal breast cancer metastases (5, 54).

Recently a crucial role for stromal microvesicle in tumor progression has been suggested (18, 19). However, whether CAF MVs could promote therapy resistant breast cancer and whether these microvesicles could recapitulate the phenotypic role of CAFs in the tumor microenvironment remain under debate.

Although HT itself suppresses ER signaling, HT alone cannot be a unique trigger of metastatic disease in luminal breast cancer. We propose that communication between CAFs and tumor cells promotes an ER dependent to an ER independent switch in metastatic disease. This can occur via the genetic transfer of miRNAs (221/222), leading to the posttranscriptional downregulation of ER and the expansion of HT resistant tumors. Recently, the overexpression of miR 221/222 was demonstrated to promote mammosphere generation in T47D cells. In addition, a manuscript from Shah and colleagues has proved the presence of miR 221 high microvesicles from CAFs (35). However, the phenotypic relevance of these CAF MVs in the context of HTR disease was not investigated.

In this article, we have demonstrated that the expansion of CD133^{hi} CSCs is functionally associated with the expansion of CAFs in experimental and patient derived HTR disease. We developed the hypothesis that CAF derived microvesicles could generate *de novo* HTR disease via a miR 221 mediated conversion of non CSCs (ER^{hi}) into therapy resistant CSCs (ER^{lo}). We generated PDX models of luminal breast cancer and isolated CAFs from HTR bone metastases (Supplementary Table S2) and through their analysis uncovered a step wise process of CAF MV mediated HT resistance: the (i) IL6 pStat3 dependent activation of CAFs, (ii) the biogenesis of oncomiR 221/222 high microvesicles, (iii) the transfer of these oncomiRs from CAF MVs to ER^{hi} cancer cells, (iv) the suppression of ER signaling, Notch3 activation, and the generation of CD133^{hi}/ER^{lo}/Notch^{hi} CSCs.

Given the pivotal role of IL6 in CAF cell growth and the generation of stromal tumor niches, we identified combination IL6R/IL6 blockade and HT as a therapeutic intervention to abrogate the establishment of stromal tumor niches and endocrine resistance in metastatic luminal breast cancer (Fig. 7). Taken together, our data suggest a novel pathologic role of stromal IL6 in

luminal breast cancer: the secretion of oncomiR 221/222 high microvesicles leading to the evolution of therapy resistant stromal tumor niches. We characterized the cellular components of this stroma tumor niche: "CD133^{hi} CSCs and CAFs" and determined the molecular machinery responsible for the niche generation: autocrine IL6 in CAFs and CAF derived microvesicle dependent downregulation of ER in cancer cells.

Disclosure of Potential Conflicts of Interest

N. Fabbri is a consultant/advisory board member for Illuminoss Medical Inc. No potential conflicts of interest were disclosed for the other authors.

Authors' Contributions

Conception and design: P. Sansone, D. Lyden, J. Bromberg
Development of methodology: P. Sansone, M. Berishaj, C. Savini
Acquisition of data (provided animals, acquired and managed patients, provided facilities, etc.): P. Sansone, M. Berishaj, V.K. Rajasekhar, Q. Chang, A. Strillacci, L. Shapiro, A. Benito-Martin, N. Fabbri, J.H. Healey, J. Bromberg
Analysis and interpretation of data (e.g., statistical analysis, biostatistics, computational analysis): P. Sansone, M. Berishaj, C. Ceccarelli, Q. Chang, A. Strillacci, R. Bowman, C. Mastroleo, F. Perna, E. Spisni, M. Cricca, D. Lyden, M. Bonafé
Writing, review, and/or revision of the manuscript: P. Sansone, V.K. Rajasekhar, L. Shapiro, L. Daly, D. Lyden, M. Bonafé, J. Bromberg
Administrative, technical, or material support (i.e., reporting or organizing data, constructing databases): P. Sansone, M. Berishaj, S.D. Carolis
Study supervision: P. Sansone, D. Lyden, J. Bromberg

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Evolution of Cancer Stem-like Cells in Endocrine-Resistant Metastatic Breast Cancers Is Mediated by Stromal Microvesicles

Pasquale Sansone, Marjan Berishaj, Vinagolu K. Rajasekhar, et al.

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