Evidence of lateral transmission of aggressive features between different types of breast cancer cells

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Abstract. Breast cancer (BrC) is a major public health problem worldwide. The intra-tumoral heterogeneity and tumor cell plasticity importantly contribute to disease progression and treatment failure. However, the dynamic interactions between different tumor clones, as well as their contribution to tumor aggressiveness are still poorly understood. In this study, we provide evidence of a lateral transmission of aggressive features between aggressive and non-aggressive tumor cells, consisting of gain of expression of cancer stem cell markers, increased expression of CXCL12 receptors CXCR4 and CXCR7 and increased invasiveness in response to CXCL12, which correlated with high levels of secretion of pro-inflammatory mediators G-CSF, GM-CSF, MCP-1, IL-8 and metalloproteinases 1 and 2 by the aggressive cells. Noteworthy, we found no evidence of a TGF-β participation in the inducible-invasive phenotype. Altogether, our results provide evidence of communication between tumor cells with different potentials for aggressiveness, which could influence intra-tumoral population dynamics promoting the emergence of clones with novel functions. Understanding these interactions will provide better targets for diagnosis, prognosis and therapeutic strategies.

Introduction

Cancer is one of the leading human health problems worldwide. Since the term includes an extensive variety of different diseases, there are still many types of cancer with a dismal prognosis. Breast cancer (BrC) is the most common cause of cancer in working age women. Since most cases are diagnosed at advanced stages, BrC mortality is particularly high in underdeveloped countries (1,2). Similar to most cancers, BrC is a highly heterogeneous disease, and although we currently have several classification and staging strategies to distinguish BrC sub-types with variable clinical outcomes, diseases originally thought as non-aggressive are often treatment resistant or relapse with highly aggressive characteristics (3). BrC cells also exhibit a large genetic and epigenetic intra-tumoral heterogeneity, conferring individual cells with specific immunosuppressive, proliferative, survival, metabolic, and invasive characteristics, that signal specific tumor clones with cancer dissemination and treatment resistance capacities (4). Furthermore, tumor clones are surrounded by a highly dynamic microenvironment formed by tumor-associated macrophages, neutrophils, T-cells, NK cells, fibroblasts, among other cells (the tumor microenvironment or TME). Although, in recent years we have seen a hefty increase in our understanding of critical interactions between tumor cells and TME cells, that has propelled the design of therapeutic molecular inhibitors (5), we significantly lack behind in our comprehension of relevant interactions between tumor clones with different aggressive features and the impact of these interactions on disease progression and prognosis.

Miller et al in a 1983 hallmark study observed that there is cooperation between metastatic and non-metastatic tumor clones. This group reported in a syngeneic mouse model that the presence of a metastatic subpopulation enabled non-mobile subpopulations to metastasize (6). More recently, a similar observation was also made by Calbo et al (7). Experiments using the Drosophila melanogaster (fruit fly) in which different cells were engineered to express either RASV12 or scrib common oncogenic mutations, revealed intra-clonal cooperation that promoted tumor growth and invasion (8). Similarly, Cleary et al observed in a mouse model of BrC that two different cellular clones had to be transferred to propagate the tumor in new mice, one clone with an Hras genetic mutation and the other with the capacity to secrete high levels of the
Wnt1 signaling molecule but harboring a wild-type \textit{Hras} (9). Soluble factors secreted by chemoresistant tumor cells and also by cancer stem cells (CSCs) promote resistance of chemosensitive cancer cells (10). Moreover, Mukherjee \textit{et al.} showed that non-migratory CSCs confer metastatic potential to sensitive cancer cells (10). Moreover, Mukherjee \textit{et al} also by cancer stem cells (CSCs) promote resistance of chemoresistant tumor cells and harboring a wild-type Hras (9).

Understanding the origin of intra-tumoral heterogeneity is one of the greatest challenges nowadays. There is evidence supporting tumor cell plasticity to microenvironmental stimuli and to genetic and epigenetic changes. Differentiated tumor cells seem capable to acquire stem cell-like properties, and conversely, CSCs can lose stemness and form more differentiated populations (12). This bi-directionality among highly adaptable cells shapes the tumor with highly organized cell populations that directly impact disease evolution and prognosis (13). The epithelial to mesenchymal transition (EMT) is a conserved embryonic developmental process that also occurs in cancer. During EMT, epithelial cells lose their typical adhesive characteristics while gaining properties more related to mesenchymal mobile cells (14). The best-understood biomolecule associated with triggering EMT is TGF-\(\beta\) (transforming growth factor-\(\beta\)), and mounting evidence supports a TGF-\(\beta\) role in cancer cell invasion, metastasis, chemoresistance and relapse (15). EMT has been shown to correlate with acquisition of a CSC-like phenotype (16,17), and circulating BrC cells often share characteristics of both stem-like cells and of EMT cells (18).

In this study, we report dynamic interactions between BrC cells with different aggressive potential leading to lateral transmission of aggressive features. We used four BrC cell lines, two characterized by an epithelial phenotype and the inability to induce metastasis in mice (MCF-7 and T47D; identified therein as non-aggressive or NA-BrC cells) and two with a mesenchymal phenotype and highly metastatic potential (HS578T and MDA-MB-231; identified as highly aggressive or HA-BrC cells). We found that aggressive cells promoted an EMT/CSC-like and invasive phenotype in non-aggressive cells. Altogether, the experimental observations fit within a molecular regulatory network in which G-CSF, GM-CSF, IL-8 and MCP-1 inflammatory cytokines induce a stem-like invasive phenotype in NA-BrC cells, which respond increasing the activity of the CXCL12/CXCR4/CXCR7 chemokine signaling axis.

Materials and methods

\textbf{Cell culture.} All cell lines were obtained from the American Type Culture Collection (ATCC). Culture media and supplements were obtained from Gibco BRL Life Technologies. BrC cells were estrogen receptor (ER)-positive cells MCF-7 and T47D, and triple-negative HS578T and MDA-MB-231. MCF-7 (HTB-22) and HS578T (HTB-126) were cultured in Dulbecco’s modified Eagle’s medium (DMEM) with High Glucose \(4.5 \text{ g/l}\) (ref. 11965-092), T47D (HTB-133) with RPMI-1640 medium (ref. 11875-093) and MDA-MB-231 (CRM-HTB-26) were cultured in Dulbecco’s modified Eagle’s medium with Nutrient Mixture F-12 (DMEM/F12, ref. 11039-021), the media were supplemented with 10% fetal bovine serum (FBS) (ref. 16000-044), 100 U/ml penicillin, 100 \(\mu\text{g/ml}\) streptomycin, and 0.25 \(\mu\text{g/ml}\) fungizone (ref. 15240-062) and were cultured at \(37^\circ\text{C}\) in 5\% CO\(_2\). All BrC cell lines were submitted to short tandem repeat analysis to verify the authenticity of each cell line. To obtain conditioned media \(2\times10^6\) cells of each cell line were plated in 182 cm\(^2\) flasks in their standard supplemented medium. Supernatants were discarded when cultures reached 80\% of confluence, cells were rinsed with PBS 1X (Phosphate Buffered Saline, Gibco, ref. 20012-027), and then 30 ml of their respective culture media without FBS was added. Conditioned media were harvested after incubation for an additional 48 h, centrifuged at 1500 rpm/5 min, aliquoted, and stored at -20\(^\circ\text{C}\) until use.

\textbf{Sera from breast cancer patients.} Sera from BrC patients and healthy blood donors (controls) were obtained from the tissues and sera bank of the Unidad de Investigación en Virología y Cáncer, Hospital Infantil de México Federico Gómez. This study was approved by the Scientific, Ethical and Biosecurity Review Boards of the Hospital Infantil de México Federico Gómez (Comité de Investigación, Comité de Ética en Investigación and Comité de Bioseguridad). All patients and healthy controls were prospectively enrolled and were informed about the nature of the study. Those willing to participate signed a written informed consent prior to specimen collection and were treated according to the ethical guidelines and best clinical practice of our institution. The identity of the participants’ names was anonymized for the duration of the study. Patients were on average 55.5 years old (range: 37-88), 22 were diagnosed with invasive ductal carcinoma, 2 with lobular and 4 with mixed carcinoma, and they had not received neoadjuvant therapy at the time of sample collection. All patients were classified as luminal A, luminal B, Her-2 positive and triple-negative and as clinical stages I, II, and III, except for 4 patients for whom we could not obtain this information. Consecutive numbers were given to the samples as they were collected.

\textbf{Induction of the invasive/stemness phenotype.} BrC cells were plated at a density of \(3\times10^4\) cells/ml/well in 6-well flat bottom culture plates. Once cells attached to the plate, supernatants were discarded, cells were rinsed with PBS 1X, and then cultured with 3 ml of conditioned media or with their respective media supplemented with 5, 10 or 20 ng/ml of human recombinant TGF-\(\beta\) (ref. 100-21) or 100 ng/ml of the following cytokines: G-CSF (ref. 300-23), GM-CSF (ref. 300-03), IL-8 (ref. 200-08) and MCP-1 (ref. 300-04) all cytokines were from PeproTech) individually or all combined in a cocktail of cytokines. After incubation for 72 h cells were harvested for analyses. To neutralize the biological activity of TGF-\(\beta\) of the HA-CMs, 2 \(\mu\text{g/ml}\) of the rabbit anti-human TGF-\(\beta\), anti-\(\beta_2\), and anti-\(\beta_3\) neutralizing antibody (anti-TGF-\(\beta\), R&D Systems, Inc., ref. MAB1835) were added according to the guideline provided by the manufacturer.

\textbf{Immunofluorescence.} Cells (\(3\times10^4\)) were seeded on coverslips for 24 h, the invasive/stemness phenotype was experimentally induced as explained above, after which cells were fixed with paraformaldehyde 4\% for 10 min, and permeabilized with 0.2\% Triton X-100 in PBS for 20 min. Cells were blocked for 1 h and then stained overnight at 4\(^\circ\text{C}\) with the primary antibodies: mouse monoclonal anti-E-cadherin
Flow cytometry. For extracellular staining 3x10⁴ cells were blocked with an unspecific IgG antibody 1:100 diluted in PBS 1X supplemented with 3% FBS for 15 min and then incubated for 30 min with mouse monoclonal anti-human CXCR3-PE-Cy7 (1:50, Clone: 1C6/CXCR3, ref. 560831), CXCR4-PE-Cy7 (1:50, Clone: 12G5, ref. 560669), CXCR7-APC (1:50, Clone: 10D1-J16, ref. 391406), CCR7-Alexa 647 (1:50, Clone: 150503, ref. 560816), CD44-PE (1:50, Clone: G44-26, 555479) (all antibodies were from BD Biosciences; except for anti-CXCR7 that was from BioLegend). Finally, cells were incubated with DAPI (4',6-diamidino-2-phenylindole dihydrochloride; Thermo Fisher Scientific, ref. D1306) for 25 min. Cells were observed using a fluorescence microscope Olympus BX51 and images were acquired with a digital camera (Camedia C4040, Olympus). FITC staining intensity was quantified using the Image Pro Plus software, and the integrated optical density (IOD) of green cells was obtained.

Invasion assay. Cells (2x10⁴) were resuspended in 200 µl of their respective media with or without FBS and placed in the upper chamber of a Transwell [6.5-mm diameter, 8 µm pore size (Corning Inc.) filled with Matrigel (Corning Inc., ref. 356237)]. Then, the Transwells were placed in a 24-well culture dish containing 800 µl of their respective media supplemented with any of the following chemoattractants: 10% FBS, the different CMs or 100 ng/ml of CXCL12 (C-X-C motif chemokine ligand 12, also known as SDF-1α, PeproTech, ref. 300-28A). To neutralize the biological activity of CXCL12 of the HA-CMs, 0.5 µg/ml of the goat anti-human CXCL12/SDF-1 neutralizing antibody (R&D Systems, Inc., ref. AF-310-NA) were added according to the guideline provided by the manufacturer. After 24 h of incubation at 37°C, invasive cells were stained with crystal violet and observed using a fluorescence microscope Motic AE31 and images were acquired with a digital camera (Motiscam 5.0 MP). When cells invaded in groups, reliable cell counts were not possible, in those cases crystal violet staining intensity was quantified using the Image Pro Plus software, and the IOD of invading cells was reported.

Analysis of TGF-β and cytokine profile. The level of TGF-β in the BrC CMs was analyzed with the human TGF-β Quantikine ELISA kit (R&D Systems, ref. DY240) according to the protocol provided by the manufacturer. To determine the cytokine profile of CMs and sera from patients with BrC, the concentration (in pgs/ml) of G-CSF, GM-CSF, IL-1β, IL-2, IL-4, IL-6, IL-8, IL-10, IL-17, IL-12 p70, Interferon-α2 (INF-α2), MCP-1, RANTES, EGF, VEGF, CXCL12 (also known as SDF-1), and metalloproteinases MMP-1, MMP-7, MMP-9, and MMP-10 were determined with the MILLIPLEX HCYTOMAG-60K kit (EMD Millipore Corp.) following the manufacturer’s recommended procedure. The analysis of data was performed in the xPONENT® Software.

Tumorsphere forming assay. After induction of the invasive/stemness phenotype, single-cell suspensions were plated in 96-well ultra-low attachment plates (Costar; Corning Inc.) at 100 cells in 100 µl of MammoCult medium plus growth factors (Stem Cell Technologies, ref. 05621). Cultures were grown for 7 days. Tumorspheres were observed and photographed using Motic AE31 microscope with a digital camera (Motiscam 5.0 MP). Spheres >50 µm were counted and graphed.

Results

Aggressive breast cancer cells promote loss of E-cadherin and invasive features in non-aggressive cells. We defined MCF-7 and T47D ER-positive cells as NA-BrC cells and HS578T and MDA-MB-231 triple-negative cells as HA-BrC cells based on expression of the EMT markers E-cadherin and vimentin and their invasive properties (Fig. 1A and B). In agreement, the former cell lines are non-metastatic in mouse, while the latter are highly metastatic in similar experimental conditions (19). Since during metastasis cellular communication is importantly mediated by secreted factors (20), we assessed whether NA-BrC cells cultured with conditioned media (CM) of HA-BrC cells, and vice-versa, modify each others aggressive characteristics. Fig. 1C shows that MCF-7 cells cultured with the HS578T CM reduced E-cadherin levels from a basal IOD of 22,000 to an IOD of 8,000, while with the CM of MDA-MB-231 cells E-cadherin levels were undetectable. This reduction of E-cadherin levels correlated well with an acquired capacity of cells to invade, observing an average of 10 and 30 MCF-7 invading cells/field when cultured in HS578T and MDA-MB-231 CMs, respectively. The same was true for T47D cells, finding a reduction of the E-cadherin IOD from 50,000 to <500, and 15-20 invading cells/field after culture...
with CM from aggressive cells (Fig. 1D). On the contrary, no significant changes were observed in HA-BrC cells cultured with CM derived from NA-BrC cells (data not shown). Thus, these results support the capacity of aggressive tumor cells to laterally transmit aggressive features into non-aggressive cells, characterized by a partial EMT phenotype with reduction of E-cadherin but not significant induction of vimentin, and increased invasiveness.

**CXCL12-CXCR4/CXCR7 axis in the inducible-invasive phenotype.** CXCL12 is an essential chemokine often involved in the migration of BrC cells (21); similarly CXCL12 receptors CXCR4 and CXCR7, but also CXCR3 and CCR7 mediate metastasis of BrC cells (22). To further explore the mechanism by which HA-BrC cells induce the migration of NA-BrC cells, we analyzed the concentration of CXCL12 in the CM of the BrC cells and the expression levels of the chemokine receptors on the NA-BrC cells treated with the CM of HA-BrC cells. Although high levels of CXCL12 were found in the CM of the BrC cell lines, no significant differences were found between aggressive and non-aggressive cells (Fig. 2A). Of the chemokine receptors, we found a discrete and more heterogeneous result with CXCR3, contrary to CXCR4, CXCR7 and CCR7 that significantly increased expression solely in response to HA-BrC CMs; CXCR4 and CXCR7 changed in both NA-BrC cells, while CCR7 only on induced T47D cells (Fig. 2B). Since CXCR4 and CXCR7 are CXCL12 receptors, CXCL12 was tested as chemotratractant in invasion assays.

We found that both induced invasive NA-BrC cells migrated in response to this chemokine (Fig. 2C). To address whether the CM of aggressive cells could also attract cells, NA-BrC cells were cultured with the HA-BrC CMs as before, and then subjected to an invasive assay in which the same CM used to stimulate them was placed as chemotratractant in the lower chamber of the transwell cameras. We observed that the induced invasive MCF-7 and T47D cells were also migrating in response to the HA-BrC CMs, with MCF-7 cells seeming more responsive than T47D cells. In addition, the CM of MDA-MB-231 cells gave a more potent response than the CM of HS578T cells, with an average IOD/field of 900,000 for MCF-7 and 320,000 for T47D invading cells (Fig. 2D). Cells did not invade in the absence of CXCL12 (data not shown). To confirm that CXCL12 had a role in the invasive phenotype, invasion assays were performed on the induced invasive MCF-7 and T47D cells using the CM of the BrC cell lines, either alone or plus a CXCL12 neutralizing antibody. We observed that the stimulated NA-BrC cells decreased their invasiveness in the presence of the inhibitory antibody (Fig. 2E). These data support the idea that HA-BrC cells induce migratory properties on NA-BrC cells together with upregulation of CXCR4 and CXCR7 expression, and that these induced invasive NA-BrC cells then move in response to CXCL12.

**A TGF-β independent but G-CSF, GM-CSF, IL-8 and MCP-1 dependent response.** We addressed whether TGF-β was responsible for the inducible-invasive phenotype. When we quantified TGF-β levels in the CM of the BrC cell lines, we found a higher concentration of this cytokine in the CM of the HA-BrC cell lines (average of 343 and 377 pgs/ml in HS578T and MDA-MB-231 cells, respectively) than in the NA-BrC CMs (80 and 14 pgs/ml in T47D and MCF-7 cells, respectively) (Fig. 3A). We then used a TGF-β neutralizing antibody to assess whether this cytokine was responsible for the inducible-invasive phenotype of NA-BrC cells. To do this, NA-BrC cells were stimulated for 72 h with CM from HA-BrC cells in the presence of the TGF-β neutralizing antibody. Surprisingly, when expression of E-cadherin was assessed we still observed a significant reduction of the E-cadherin IOD promoted by the HA-BrC CMs (Fig. 3B). Similarly, the stimulated NA-BrC cells remain invasive in spite of the presence of the neutralizing antibody (Fig. 3C). Additionally, we added exogenous TGF-β at various concentrations to MCF-7 and T47D cells for 72 h, with no observed variations in E-cadherin or vimentin expression. Accordingly, we did not observe a TGF-β-induced invasiveness (Fig. 3D). TGF-β is known to induce its own expression in Caski cervical cancer cells (23). Exogenous TGF-β increased its own gene expression in Caski cells, and the neutralizing antibody abolished this induction, indicating that both recombinant TGF-β and the anti-TGF-β antibody were functional (data not shown).

We then analyzed for the presence of the following growth factors, cytokines, chemokines and matrix metalloproteinases (MMPs) in the CM of the BrC cell lines, which have been previously associated with tumor cell invasion and/ or EMT (24,25): Granulocyte-colony stimulating factor (G-CSF), Granulocyte macrophage-colony stimulating factor (GM-CSF), IL-1β, IL-2, IL-4, IL-6, IL-8, IL-10, IL-17, IL-12p70, Interferon-α2 (INF-α2), Monocyte chemotactic protein-1 (MCP-1), Regulated on activation, normal T-cell expressed and secreted (RANTES, also known as CCL5), epidermal growth factor (EGF), Vascular endothelial growth factor (VEGF), MMP-1, MMP-2, MMP-7, MMP-9 and MMP-10. Important differences between the CMs of NA- and HA-BrC cells were found in the levels of G-CSF, GM-CSF, IL-8 and MCP-1 (Fig. 4A). These cytokines are known to induce differentiation, proliferation and activation of myeloid cells recruited to the tumor microenvironment, and often, they also induce autocrine activation of tumor cells (26-28).

To assess whether these factors participate in the inducible-invasive phenotype, MCF-7 and T47D cells were cultured in media supplemented with G-CSF, GM-CSF, IL-8 and MCP-1 for 72 h. We found that MCF-7 cells were highly responsive to these cytokines with G-CSF and GM-CSF inducing a partial loss, and IL-8 and MCP-1 an almost complete loss of E-cadherin levels (Fig. 4B right plot). We also observed an increase of CXCR4 expression in response to all cytokines, while CCR7 only on induced T47d cells, and specifically IL-8 and MCP-1 (Fig. 4C left plots), which closely correlated with induction of invasiveness (Fig. 4D left panels). Overall, these data show a great correlation between IL-8/MCP-1-induction of CXCR7 and E-cadherin reduction/increased invasion.

Somehow the results obtained with T47D cells were more complex. We only observed a very discreet but significant reduction of E-cadherin levels on T47D cells, and specifically upon stimulation with IL-8 and MCP-1 (Fig. 4B right plot). We also observed an increase of CXCR4 expression in response to all cytokines, while CXCR7 increased expression only in response to IL-8 and MCP-1 (Fig. 4C right plots). However, the T47D invasion assay was not as clear, since G-CSF, GM-CSF and MCP-1 were able to induce invasion but IL-8 was...
Figure 1. Aggressive breast cancer cells promote mesenchymal and invasive features in non-aggressive cells. (A) Analysis of the basal expression levels of epithelial to mesenchymal (EMT) markers E-cadherin (green) and vimentin (red), nuclei (DAPI, blue) by immunofluorescence (IF) staining. (B) Invasion assays. (C) MCF-7 cells were cultured with CM from HA-BrC cell lines HS578T and MDA-MB-231 for 72 h. As controls, MCF-7 cells were cultured with NA-BrC CMs or were cultured in their regular media (unstimulated). Left panels show representative IF images of EMT markers and optical images of invasion assays. Right panels show plots of the integrated optical density (IOD) of E-cadherin levels and the number of invasive cells. (D) T47D cells were cultured with CM from HA-BrC cell lines HS578T and MDA-MB-231 for 72 h. As controls, T47D cells were cultured with NA-BrC CMs or were cultured in their regular media (unstimulated). Left panels show representative IF images of EMT markers and optical images of invasion assays. Right panels show plots of the integrated optical density (IOD) of E-cadherin levels and the number of invasive cells. Representative images are shown. Data represent the mean ± SEM (standard error of the mean) from 3 independent experiments; ***P<0.001. Scale bars indicate 100 µm. Magnification of x400 for IF images and x100 for optical images.
Figure 2. The CXCL12-CXCR4/CXCR7 axis in the inducible-invasive phenotype. (A) Levels of CXCL12 expressed in pg/ml in the CMs of the BrC cell lines. (B) MCF-7 and T47D cells were cultured with CM from HA-BrC cell lines and controls, and expression of chemokine receptors was analyzed by flow cytometry. (C) CXCL12 or (D) CMs as chemotacticants. (E) CMs from the HA BrC cells were used as chemotacticants and 0.5 µg/ml of an anti-CXCL12 neutralizing antibody was added to the CMs. Invasive cells were quantified after 24 h (magnification of x100). The integrated optical density (IOD) values of invading cells were plotted. Data represent the mean ± SEM from 3 independent experiments; *P<0.05, **P<0.01 and ***P<0.001. Scale bars indicate 100 µm.
and of MMP-1 and MMP-2 metalloproteinases in sera of BrC patients (see Table I for the clinical characteristics of patients) and healthy controls, finding detectable levels of all analytes tested. Since we could only obtain five sera of each BrC subtype no statistical analysis was performed (Fig. 4F).

CD44, Oct4 and Sox2 are upregulated in induced-invasive cells. The hyaluronan receptor (also known as CD44) has been widely associated with a CSC-like phenotype, and Oct-4 and Sox-2 are essential transcription factors involved in maintenance of pluripotency during embryogenesis, and their induced-expression has been evidenced in several types of cancer upon acquisition of metastatic potential and EMT (29,30). When we evaluated the expression of these CSC markers on the BrC cell lines we found that CD44 positive cells were not (MCF-7=0% of positive cells) or rarely represented (T47D=2.83%) on NA-BrC cells, while they were highly represented on HS578T (90.8%) and MDA-MB-231 (96.1%) HA-BrC cells (Fig. 5A). A similar observation was made for Oct-4 and Sox-2 (Fig. 5B). We addressed whether the inducible-invasive trait correlated with acquisition of these stem markers. We found that both the frequency of CD44 positive cells and the MFI of CD44 expression were induced upon treatment with the HA-BrC CMs in both MCF-7 and T47D NA-BrC cells (Fig. 5C and D, respectively). T47D cells were the most sensitive cells to stimulation as more than half of the cells expressed CD44 upon treatment with the MDA-MB-231 CM. We also observed a significantly increased expression of Oct-4 and Sox-2 in MCF-7 cells upon stimulation with the CM from HA-BrC cell lines (Fig. 5E). T47D cells significantly changed the expression of Sox-2 upon stimulation, while the levels of Oct-4 remained low (Fig. 5F).

A more quantitative analysis of the frequency of Sox-2 expressing cells by flow cytometry confirmed the results obtained with the immunofluorescence analyses (Fig. 5G). Since the formation of tumourspheres is a model of CSC seeding and expansion (31), we addressed the BrC cell lines basal and induced potential of formation of tumourspheres. We observed that all the cell lines exhibited similar capacities of formation of tumourspheres (Fig. 5H), the only difference found was the morphology of the spheres, with HA-BrC cells forming less adherent spheres that resemble cell aggregates, typical of aggressive cell lines with low expression of adhesion proteins (32). When NA-BrC cells were treated with the CM of HA-BrC cells, the frequency and morphology of the tumourspheres remained unchanged; however, spheres of increased size were observed (Fig. 5I). Taken together all these data show that the inducible-invasive trait correlates with acquisition of a CSC-like phenotype defined by expression of CD44, Oct-4 and Sox-2 stemness markers, and also with the formation of larger tumourspheres in low adherent plates.

Discussion

The original conception of cancer initiation and progression highlighted the importance of accumulation of genetic mutations, with tumors exhibiting greater genetic heterogeneity also exhibiting an increased risk to display more aggressive features. More recent studies have also highlighted the capacity of the tumor to communicate with non-tumor cells within the tumor

| Serum from patients | Histological subtype | TNM staging | Clinical stage | Molecular classification |
|---------------------|---------------------|-------------|----------------|-------------------------|
| UIVC-IDC-13         | IDC                 | T2, N0, M0  | IIA            | Triple-negative         |
| UIVC-IDC-14         | IDC                 | T1, N0, M0  | I              | Luminal B               |
| UIVC-IDC-15         | IDC                 | T1, N0, M0  | I              | Luminal A               |
| UIVC-IDC-16         | IDC                 | T1, N0, M0  | I              | Triple-negative         |
| UIVC-IDC-17         | IDC                 | WD          | WD             | Her-2                   |
| UIVC-IDC-18         | IDC                 | WD          | WD             | Luminal A               |
| UIVC-IDC-19         | IDC                 | T2, N0, M0  | IIA            | Triple-negative         |
| UIVC-IDC-20         | IDC                 | WD          | WD             | Luminal B               |
| UIVC-IDC-21         | IDC                 | WD          | WD             | Her-2                   |
| UIVC-IDC-22         | IDC                 | T1, N0, M0  | I              | Luminal A               |
| UIVC-IDC-23         | IDC                 | T1, N1, M0  | IIA            | Luminal A               |
| UIVC-IDC-24         | IDC                 | T1, N0, M0  | I              | Luminal A               |
| UIVC-IDC-25         | IDC                 | T2, N0, M0  | IIA            | Triple-negative         |
| UIVC-IDC-26         | IDC                 | T2, N0, M0  | IIA            | Her-2                   |
| UIVC-IDC-27         | IDC                 | T2, N0, M0  | IIA            | Luminal A               |
| UIVC-IDC-28         | IDC                 | T2, N1, M0  | IIB            | Her-2                   |
| UIVC-IDC-29         | IDC                 | T1, N0, M0  | I              | Luminal A               |
| UIVC-IDC-30         | IDC                 | T3, N0, M0  | IIB            | Luminal A               |
| UIVC-IDC-31         | IDC                 | T2, N0, M0  | IIA            | Luminal A               |
| UIVC-IDC-32         | IDC                 | T1, N0, M0  | I              | Her-2                   |
| UIVC-IDC-33         | IDC                 | T3, N1, M0  | IIIA           | Luminal A               |
| UIVC-IDC-34         | IDC                 | T2, N0, M0  | IIA            | Luminal B               |
| UIVC-LC-2           | ILC                 | T3, N1, M0  | IIIA           | Triple-negative         |
| UIVC-LC-3           | ILC                 | T2, N1, M0  | IIB            | Luminal A               |
| UIVC-MC-2           | MC                  | T1, N1, M0  | IIIA           | Luminal B               |
| UIVC-MC-3           | MC                  | T3, N1, M0  | IIIA           | Luminal B               |
| UIVC-MC-4           | MC                  | T2, N0, M0  | IIA            | Her-2                   |
| UIVC-MC-5           | MC                  | T2, N0, M0  | IIA            | Luminal A               |

IDC, invasive ductal carcinoma; ILC, invasive lobular carcinoma; MC, mixed (lobular/ductal) carcinoma; WD, without data.
Figure 3. The inducible-invasive phenotype is TGF-β independent. (A) The concentration expressed in pgs/ml of TGF-β was measured in the CM from all BrC cell lines and data were plotted. The inducible-invasive phenotype of MCF-7 and T47D cells was activated with HA-BrC CMs in the presence of 2 µg/ml of neutralizing anti-TGF-β. After 72 h of culture, (B) EMT markers were analyzed by IF and IODs of E-cadherin expression were quantified and plotted. (C) Invasion assays were performed. Left panels show representative images of invading cells and right plots show the number of invading cells. (D) Analysis of EMT markers and invasion assays of NA-BrC cell lines treated with increasing concentrations of TGF-β. Representative images are shown. Data represents the mean ± SEM from 3 independent experiments; ***P<0.001; ns, non-significant. Scale bars indicate 100 µm, magnification of x400 for IF and x100 for invasion assays.
Figure 4. (A-D) The role of pro-inflammatory mediators in the inducible-invasive phenotype. (A) Milliplex assays were performed to determine the concentration of pro-inflammatory mediators and metalloproteinases (expressed in pg/ml) in all the CMs; only analytes exhibiting significant differences between the CMs of NA- and HA-BrC cells are shown. MCF-7 and T47D cells were cultured with 100 ng/ml of any of the following: G-CSF, GM-CSF, IL-8 or MCP-1 for 72 h. (B) Analysis of the EMT marker E-cadherin by IF. (C) Plots of the analysis of CXCR4 and CXCR7 chemokine receptor expression by flow cytometry. Invasion assays using CXCL12 (D) as chemoattractant. Representative images and plots of resulting data are shown. Data represent the mean ± SEM from 3 independent experiments; *P<0.05, **P<0.01 and ***P<0.001. In the panels of T47D cells of (D), IL-8 was significantly different than the other cytokines (*P<0.05). Scale bars indicate 100 µm and magnification, x100.
microenvironment as the source of aggression (5). In this study, we evidenced a potential mechanism of paracrine communication between tumor cells that results in lateral transmission of aggressive features. The induced aggressive characteristics consist of acquisition of a partial EMT phenotype with loss of E-cadherin but without gain of vimentin expression, gain of CSC markers Cd44, Sox-2 and Oct-4, increased expression of CXCL12 receptors CXCR4 and CXCR7, and increased invasiveness in response to CXCL12. Altogether, these mechanisms of tumor communication may facilitate the appearance of new clones with novel functions, further extending the clonal heterogeneity that increases the tumor aggressive potential without relying on genetic irreversible mutations.

The clonal evolution model of tumors points out that distinct genetic clones exhibit differential fitness, and only those clones with specific advantages and under particular selective pressures will be maintained, favoring the progression of disease in a kind of Darwinian evolution competition (33). In 2014, Marusyk et al proposed that interactions between rare and affluent tumor clones favored the emergence of clones with novel phenotypes and functions allowing the tumor to adapt to microenvironmental changes (34). Other studies support intra-clonal communication and cooperation, particularly among metastatic and non-metastatic clones (6,7,11), adding another layer of complexity to the origin and evolution of tumors. The plasticity of the tumor cell has been extensively studied, with the EMT at the center of this plasticity. More recent evidence support that cancer cells undergoing EMT also increase expression of stem markers, and tumors in which the EMT/stemness programs are active, are also more invasive and metastatic denoting cancers with the worst clinical outcomes (16,17,24,35,36). How the EMT and stemness programs support tumor heterogeneity and intra-clonal coexistence to facilitate tumor maintenance remains as one of the most challenging puzzles in cancer biology. Interestingly, although TGF-β is one of the best characterized EMT triggers (15), we could not find any evidence of a TGF-β participation in the inducible invasive stem-like phenotype. To our knowledge, Mani et al in 2008 were the first to describe a strong correlation between EMT and stemness (16), laying the bases for a novel understanding of tumor plasticity and tumor aggression. Today, mounting evidence supports that association; for instance, BrCs with a high density of CD44+ positive cells are specifically associated with reduced disease-free survival (37). Expression of Oct-4 and Sox-2 is also associated with poor clinical outcomes in BrC patients (38,39). While we were working in this study, Mukherjee et al reported that CSCs with different capacities of migration co-exist within primary tumors and in MCF-7 mammospheres. Low migrating inner core CSCs have the capacity to induce migratory properties into outer core non-CSCs through paracrine secretion of EGF, TGF-β1, VEGF and IL-6 (11). In agreement with our study, induced non-CSCs showed increased expression of stemness markers CD44, Oct-4 and Sox-2, and invasion was marked by upregulation of CXCR4. We observed an increased expression of CXCR4 and CXCR7 on induced-NA-BrC cell lines, which promoted cell migration in response to CXCL12, particularly in induced MCF-7 cells. In the case of T47D there is not a clear correlation that could be explained because contrary to MCF-7 cells, T47D cells also upregulate CXCR3 and CCR7 expression, as it can be observed in Fig. 2B. Still, the CXCR4/CXCR7/CXCL12
Figure 5. (A-D) The induced-invasive phenotype correlates with acquisition of stemness markers. Analysis of the basal expression levels of stemness markers: CD44 by flow cytometry (A), and of Oct-4 and Sox-2 by immunofluorescence (IF) (B). Representative images are shown. Scale bars indicate 100 µm (B). Magnification, x400. Analysis of the expression levels of CD44 after induction of the invasive phenotype in MCF-7 (C) and T47D (D) cells. The upper panel shows Cd44 expression and the lower panel shows plots of the frequency of CD44+ cells and the CD44 MFI. Data represent the mean ± SEM from 3 independent experiments, representative images are shown. *P<0.01 and **P<0.001.
Figure 5. Continued. (E-I) The induced-invasive phenotype correlates with acquisition of stemness markers. Analysis of Oct-4 and Sox-2 IODs is shown for MCF-7 (E) and T47D (F) cells. (G) Sox-2 was also examined by FACS; the upper panels show representative images of cell density plots, while the frequency of Sox-2 positive cells and the MFI of Sox-2 expression are graphed below. (H) The intrinsic sphere forming efficiency of the BrC cell lines was analyzed in ultra-low attachment plates, and (I) after induction of the invasive phenotype in MCF-7 and T47D cells. Plots of the frequency and size of the tumorspheres are shown. Data represent the mean ± SEM from 3 independent experiments, representative images are shown. *P<0.05, **P<0.01 and ***P<0.001. Scale bars indicate 50 µm for MCF-7 and T47D, and 100 µm for HS578T and MDA-MB-231 tumorspheres (H and I). Magnification, x400.
CSC (HA-clone):  
Mesenchymal features  
Positive to stemness markers  
Highly invasive  

Non-CSC (NA-clone):  
Highly heterogeneous  
Epithelial features  
Negative to stemness markers  
Non-invasive  

Figure 6. Working model. BrC is a heterogeneous disease with extensive intra-tumoral clonal diversity in which paracrine communication between different tumor clones influences the aggressive behavior of the tumor. Pro-inflammatory mediators, such as MCP-1, IL-8, GM-CSF and G-CSF, secreted by highly aggressive (HA) tumor clones have the potential to activate signaling pathways associated with aggressive behavior in non-aggressive (NA) clones. These paracrine intra-clonal communication results in the formation of populations with particular transcriptional profiles favoring processes related to the partial epithelial to mesenchymal transition (loss of E-cadherin but no gain of vimentin), stemness and invasion. Induced cells with acquired EMT/stemness like and invasion potential concur with upregulation of chemokine receptors, responding to HA-BrC (or other tumor stromal cells) secreted chemokines. In addition, paracrine secretion of metalloproteinases (MMPs) would facilitate the invasion of induced-clones through degradation of the extracellular matrix. This mechanism may be critical to shape the tumor clinical outcome influencing metastasis, chemoresistance and disease relapse.

axis is widely documented as a potent promoter of invasion and metastasis in several types of cancer, and most likely also has an important role in the promotion of invasion of the induced-NA BrC cells (40,41).

Tumors with high density of CSCs are characterized by high expression of inflammatory mediators (42,43). We found that HA-BrC cell lines secret high levels of G-CSF, GM-CSF, MCP-1 and IL-8. Of them, MCP-1 and IL-8 were potent inducers of invasiveness of MCF-7 cells, and G-CSF, GM-CSF, IL-8 and MCP-1 of T47D cells. MCP-1 and IL-8 are considered critical pro-tumoral cytokines mainly because of their capacity to shape the TME by attracting immune cell populations (44,45). We have also observed that aggressive tumor cells are particularly proficient to attract monocytes/macrophages through secretion of GM-CSF and MCP-1, and monocyte/macrophage co-cultivation with tumor cells further increase secretion of IL-8 and IL-1β (46). Importantly, this study supports the capacity of tumor cells to secrete pro-inflammatory cytokines promoting stemness and invasion independent of other cellular components of the tumor stroma. In agreement, other studies support that autocrine overexpression of MCP-1 and IL-8 promotes tumor cell proliferation, migration, chemoresistance, metastasis and disease relapse (27,28,47). In patients with breast and prostate cancers MCP-1 overexpression correlates with poor prognosis (48,49), and autocrine regulation of MCP-1 is associated with EMT, immunosuppression and metastasis (50). Blocking MCP-1 in triple-negative BrC decreased the frequency and self-renewal potential of breast CSCs (51). Other studies support an association between IL-8, MCP-1 and MMPs, and induction of CSCs and metastasis (24,51,52). Of note, we also observed high levels of MMP-1 and MMP-2 secreted by HA-BrC cells.

In conclusion, our results support a model in which in a heterogeneous disease like cancer, highly aggressive tumor clones communicate with less aggressive clones through paracrine mediators, such as IL-8, MCP-1, G-CSF, GM-CSF and MMPs, transferring aggressive features in line with a CSC-like phenotype and increased potential for invasion (Fig. 6). Although TGF-β remains the best-understood signal accountable for tumor cell plasticity, we could not find TGF-β participation in the inducible-invasive phenotype. Further studies should look into these pro-inflammatory
factors in carefully staged BrC patients, comparing metastatic tumors against non-metastatic tumors or comparing other relevant clinical parameters, such as resistance to treatment, disease relapse and overall survival. We were unable to obtain the clinical data in our series of patients. Almost all the patients included in this study were classified as stage I and II, with only three patients in stage III and none in stage IV. Understanding the mechanisms guiding intra-tumoral heterogeneity, cell plasticity and tumor aggressiveness will provide better targets for diagnosis, prognosis and therapeutic strategies.

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