Can we predict and prevent specific sites of metastases in breast cancer patients?

Fernando Salvador†3, Anna Bellmunt†3 & Roger R Gomis*1,2

1Oncology Program, Institute for Research in Biomedicine (IRB Barcelona), The Barcelona Institute of Science & Technology, Barcelona, Spain
2Institució Catalana de Recerca i Estudis Avançats (ICREA), Barcelona, Spain
*Author for correspondence: Tel.: +34 934039959; roger.gomis@irbbarcelona.org
†Authors contributed equally

Despite improvements in breast cancer therapies, cancer cells frequently spread to distant organs years or decades after primary tumor surgery and adjuvant treatment. This expansion, known as metastasis, can bring about fatal consequences. Traditionally, the risk of metastasis has been predicted by prognostic factors such as tumor size, axillary lymph node status and histological grade. More recently, genomic tests have also been used for this purpose. The presence of ER, PR and ERBB2 gene amplification are currently key markers in the characterization of breast tumor type that drive the selection of specific therapies [1]. ER-positive tumors are more prone to metastasize into the bone, whereas ER-negative tumors preferentially spread to visceral organs such as lung, liver and brain [2]. However, the reliability of these markers is limited. In this regard, substantial efforts have been made to find new markers that predict the most probable target organ of metastasis, with the aim to improve diagnosis and develop organ-specific treatments for breast cancer metastatic patients.

Metastasis is an inefficient process through which cancer cells must overcome several hurdles to establish a secondary lesion in a distant site. These steps involve intravasation into the blood stream, extravasation into a distant tissue and colonization of the target organ. Colonization involves cancer cell–host tissue interactions, evasion of the immune system, activation of cytokine signaling and extracellular matrix modifications that allow tumor cells to complete metastatic growth. This set of activities supports the establishment of a new metastasis that mimics the formation of a new and independent tumor entity [3]. Interestingly, recent evidence supports the notion that modifications of the microenvironment of distant organs are required prior to the tumor cells reaching the metastatic site. The preparation of a ‘premetastatic niche’ suitable for the reception and growth of metastatic cells may be needed [4]. These lines of evidence collectively validate the initial ‘seed and soil’ hypothesis promulgated by Steven Paget in the 19th century, suggesting that the local microenvironment of a specific tissue is more accessible, fitted and hence permissive than others for the establishment and colonization of a given tumor cell [5].

During the past 15 years, several studies have identified sets of genes whose expression is associated with metastasis – some in a tissue-specific manner. Unfortunately, most of these genes have failed to provide new diagnostic tools to stratify patients on the basis of risk of distant relapse and tissue-specific metastasis. The absence of primary tumor sample cohorts for which clinical annotations of site-specific time to metastasis are available, as well as feasibility issues regarding the collection of metastatic biopsies, has become a major limitation. These limitations, in turn, are magnified when the prognostic/predictive power of genes associated with metastasis is restricted to the primary tumor. Many tissue-specific metastasis genes may be gained or lost at the distant site where tissue-specific functions are needed. Contrary, genes whose expression changes at the primary site and that are associated with metastasis may confer both a specific advantage for growth at the primary site and beyond once disseminated to specific sites. Alternatively, these genes can be ascribed to clones within the heterogeneity of primary tumor populations [6]. These clones have site-specific advantages with respect to settling at the distant site that they have migrated to and where they are expanded and dominant.

Several genes contribute to the lung metastasis signature in primary tumors [7]. Elevated expression of EREG, PTGS2, MMP1 and cytokine ANGPTL4 increases breast cancer cell extravasation in lung capillaries [8,9]. Lung-tropic breast cancer cells also express VCAM, which interacts with macrophages and enhances cell survival by activating PI3K–AKT signaling [10]. Similarly, the downregulation of RARRES3 facilitates the adhesion of breast cancer cells to the extracellular matrix proteins of lung parenchyma and suppresses tumor cell differentiation, thereby favoring metastasis initiation [11]. All these genes, beyond their predictive value in the clinical setting, are potential candidates for the prevention and therapeutic intervention of metastasis. But
it is unclear when, where and how they could be effective.

A large subset of breast cancer patients suffers from bone metastasis as the first site of relapse, and for these individuals the disease is largely confined to bone during the course of the disease [12]. Due to the fenestrated endothelia, bone marrow sinusoids are more permissive to the homing of tumor cells than the capillaries of other tissues. In the bone, breast cancer cells can take advantage of several factors secreted by bone matrix cells, such as chemokine CXCL12, to activate a survival-signaling pathway. In fact, tumor cells with elevated SRC signaling activity and high levels of CXCR4, a receptor for CXCL12, are preferentially favored by survival signals in the bone marrow, and the expression of this receptor is associated with breast cancer bone relapse [13–15]. To generate an overt metastasis, a variety of factors such as PTHRP, TNF-α and IL-6/11 are secreted by tumor cells, stimulating the production of RANKL from osteoblasts. RANKL activates its receptor RANK to promote osteoclast differentiation. Osteoclast activation is a hallmark of osteolytic development and promotes bone resorption. The osteolytic process causes the release of bone matrix growth factors into the microenvironment (i.e., TGF-β), thus stimulating tumor cell growth. This ‘vicious cycle’ gives rise to aggressive tumor cells in bone metastasis [16,17]. The expression of these factors correlates with poor prognosis and bone relapse in some breast cancer patients [18,19]; however, it fails to predict the risk of bone metastasis in early stage tumors [20].

The identification of new predictive molecular biomarkers in nonadvanced tumors is of emerging clinical interest. Bisphosphonates and Denosumab have proven effective in the management of the morbidity of skeletal-related events morbidity. However, these treatments do not improve disease progression or overall survival rates [21]. Interestingly, recent evidence showed that 16q23 genomic gain in early stage primary tumors is associated with a high risk of developing bone metastasis and with poor overall survival. The MAF gene was identified as the genetic driver of the 16q23 region. In fact, breast cancer patients with high MAF expression (mRNA and protein) have a higher cumulative risk of metastasis to bone but not to other organs. Moreover, functional validation and mechanistic studies showed that MAF acts as a transcription factor to control the expression of a gene program, including functions such as migration, adhesion and tumor cell–stroma interaction. Among these genes, PTHrP was identified as an important element for MAF-driven bone metastasis [22]. This novel finding opens up new therapeutic strategies in breast cancer. Bone microenvironment modifying agents such as bisphosphonates and the anti-RANK ligand antibody Denosumab have the theoretical potential to prevent bone metastasis, albeit data from clinical trials are as yet inconclusive in unselected patient populations [21,23]. The identification of a biomarker that predicts bone-specific metastasis in breast tumors in a timely manner has raised the possibility of including such agents in the adjuvant setting to effectively prevent dissemination and bone metastases in MAF-expressing breast cancer patients [22].

While breast cancer metastasis to the liver and brain is less frequent than to the bone, the former have the worst outcome. Similarly to the bone, the hepatic endothelium is permissive for cancer cell extravasation. In this process, adhesion molecules are involved in the establishment of metastasis. Claudin-2 plays a key role in mediating the interaction between hepatocytes and cancer cells promoting the activation of metastatic signaling pathways. Indeed, Claudin-2 expression is considered a poor prognosis factor that mediates breast cancer relapse to the liver [24,25].

In contrast to bone and liver, the brain is the most difficult organ to access by breast cancer cells due to the presence of the blood–brain barrier (BBB). Consequently, most brain metastasis mediators are adhesion-, extravasation- and survival-related genes [26,27]. On the other hand, the presence of the BBB also limits drug delivery, thus impeding effective brain metastasis treatment. Recently, it has been suggested that patients with the HER2-enriched breast cancer subtype treated with Trastuzumab develop a higher risk of metastasis to the brain compared with other organs. New small drugs that penetrate the BBB, including lapatinib, are being used in the advance setting treatment [28].

Recent years have witnessed a significant improvement in breast cancer therapy directed at reducing primary tumor growth; however, distant metastasis has emerged as a new problem. Current therapies, mainly aimed at the primary tumor, are not as effective at preventing and controlling metastasis to distant organs. The metastasis gene signatures from primary tumors identified in the last decade provide relevant information about the mechanisms underlying metastasis mechanisms, and tissue specificity. This information may eventually allow the identification of patients who can benefit from the inclusion of therapies seeking to prevent tissue-specific metastasis. The integration of these predictive markers in routine clinical practice opens up new avenues
in an era of personalized medicine. In addition, the development of organ-specific metastatic animal models would contribute to establishing preclinical systems to functionally validate metastasis biomarkers, thus providing an invaluable tool with which to study organ-specific metastasis and to develop new therapies.

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