Role of plasmacytoid dendritic cells in breast cancer bone dissemination

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Osteolytic bone metastases are common in breast cancer (BCa). Approximately, 70% of patients dying from BCa show evidence of bone metastasis at postmortem examinations. The presence of such bone lesions usually signifies serious morbidity and a grave prognosis. Despite the complications deriving from bone metastasis, the therapies for metastatic BCa patients are limited and are not aimed at controlling the disease. Therefore, developing new strategies to control bone metastasis and to improve patient survival is an absolute necessity, which requires a deeper understanding of the molecular mechanisms involved in BCa metastatic dissemination.

As the primary tumor disseminates to the bone, it triggers the production of osteolytic cytokines and growth factors that—altogether—(1) result in osteoclast activation, (2) promote the growth of tumor cells and (3) facilitate the establishment of an immunosuppressive microenvironment. Moreover, the products of bone cells are critical for the normal development of the hematopoietic and immune systems. Thus, understanding the influence and interaction of metastasizing cancer cells with cells of the skeletal system and on cells of the immune system will provide clues for the design of preventive and therapeutic strategies for osteolytic bone metastasis.

In a pre-clinical mouse model of metastatic BCa, we observed high numbers of plasmacytoid dendritic cells (pDC) in the bone, which continued to increase as the tumor growth progressed (Fig. 1). Increased pDC infiltration at both primary and the metastatic sites has been reported also in BCa patients, but the significance of these findings was unclear. Besides BCa, lung cancer and multiple myeloma, which primarily affects the skeleton, have been associated with an increased bone infiltration by pDC. This indicates that pDC may exert an important role in the establishment of bone metastases. But the question remains what role, if any, do these cells play?

pDC can induce immunosuppression through a variety of mechanisms. In BCa, pDC promote tumor progression via the expression of ICOS-ligand and also as a result of CD40/CD40L interactions, which allow for the accumulation of immunosuppressive CD4+ T cells and hence limit the number and function of cytotoxic CD8+ T cells. These soluble factors induce indeed the osteoclastogenesis, either directly or indirectly. The role of pDC-generated soluble RANKL, which is critical for the osteoclast-mediated bone resorption, hence helping metastatic cells to grow. A recent publication has shown that pDC isolated from the bone marrow of rats express high levels of RANKL. This observation adds a further facet to the role of pDC in bone metastasis, whereby pDC-generated soluble RANKL may directly induce osteoclastogenesis by acting on bone marrow osteoclast progenitors. Using a murine BCa model, we have recently identified that, besides immunosuppressive T cell populations, myeloid-derived suppressor cells (MDSC) accumulated in high numbers together with pDC during BCa bone dissemination. Furthermore, MDSC in the cancer-bone microenvironment were found to function as novel osteoclast progenitors. Based on these findings, one could speculate that pDC-generated RANKL may directly act upon MDSC, inducing their differentiation into osteoclasts and thus promoting bone destruction and local BCa growth.
Although the above mentioned observations pointed to a possible role for pDC in promoting bone metastasis, a more direct and substantiated evidence was necessary. This led us to deplete pDC in vivo using an anti-PDCA-1 antibody, which causes a selective and effective pDC depletion. Our data clearly show that pDC-depleted mice fail to develop BCa bone metastasis and also that the overall tumor growth is dramatically reduced in pDC-depleted mice as compared with their normal counterparts.2 Further evidence in support of this observation was established in IFNα receptor-deficient (Ifnar−/−) mice, which lack functional pDC and also fail to develop BCa-derived bone metastasis. pDC-depleted mice exhibited low levels of osteoclastogenesis-promoting cytokines and growth factors. Reduced tumor burdens in these animals were the result of a skew in the immune response toward a Th1 profile, resulting in increased levels of cytotoxic CD8+ T cells as well as in an overall decrease of immunosuppressive cells.

Taken together, our data suggest that pDC may play a key role in the establishment of BCa bone metastasis. This novel function of pDC makes them a viable target for the development of novel therapeutic strategies. Both in humans and mice, pDC express the DC immunoreceptor (DCIR), which is a putative C-type lectin receptor (CLR). DCIR-mediated antigen uptake by human pDC leads to efficient antigen presentation and results into the induction of a memory T cell response.6 Besides DCIR, human pDC also express sialic acid binding Ig-like lectin H (Siglec-H). Antigen presentation to pDC via Siglec-H induces a Th1/Th17 polarization of CD4+ T cells without skew toward a Th2 or regulatory T (Treg) profile.7 Therefore, targeting DCIR and Siglec-H may be useful in the treatment of BCa. BCa-associated pDC are irresponsive, meaning that they fail to produce Type I IFNs, to TLR9 agonists such as CpG-A. Nevertheless, the therapeutic activation of pDC with imiquimod (a TLR7 agonist) has been shown to result in pDC-dependent Type I IFN.
production. Hence, several approaches may be used for the development of better therapeutic strategies against metastatic BCa and could possibly be extended to the treatment of other carcinomas associated with osteolytic bone pathology.

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Disclosure of Potential Conflicts of Interest
No potential conflicts of interest were disclosed.