Metastasis Dormancy in Estrogen Receptor–Positive Breast Cancer

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Abstract

About 20% to 40% of patients with breast cancer eventually develop recurrences in distant organs, which are often not detected until years to decades after the primary tumor diagnosis. This phenomenon is especially pronounced in estrogen receptor–positive (ER⁺) breast cancer, suggesting that ER⁺ cancer cells may stay dormant for a protracted period of time, despite adjuvant therapies. Multiple mechanisms have been proposed to explain how cancer cells survive and remain in dormancy, and how they become reactivated and exit dormancy. These mechanisms include angiogenic switch, immunosurveillance, and interaction with extracellular matrix and stromal cells. How to eradicate or suppress these dormant cancer cells remains a major clinical issue because of the lack of knowledge about the biologic and clinical nature of these cells. Herein, we review the clinical manifestation of metastasis dormancy in ER⁺ tumors, the current biologic insights regarding tumor dormancy obtained from various experimental models, and the clinical challenges to predict, detect, and treat dormant metastases. We also discuss future research directions toward a better understanding of the biologic mechanisms and clinical management of ER⁺ dormant metastasis.

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Introduction

Nearly all breast cancer–related deaths are caused by metastases rather than the primary tumor. Different subtypes of breast cancer exhibit distinct metastasis behaviors in terms of the temporal kinetics and anatomic sites. Estrogen receptor–positive (ER⁺) breast cancer, in particular, predominantly recurs in bone, often years to occasionally decades after the diagnosis of the primary tumor. This protracted latency suggests a dormant stage of metastasis progression wherein cancer cells either stay quiescent or proliferate very slowly. Although late recurrences (>5 years) occur in more than 50% of patients, our knowledge about tumor dormancy is extremely limited. This is largely due to the fact that dormant cancer cells are rare and difficult to detect and isolate from clinical specimens. Moreover, ideal animal models that can fully recapitulate the natural history of dormant tumors are also lacking. Despite these difficulties, several hypotheses have been proposed as major mechanisms underlying tumor dormancy. The common theme of these mechanisms is the cross-talk between tumor cells and the microenvironment they encounter. Processes and factors that have been implicated in dormancy include angiogenesis (1, 2), immunosurveillance (3–5), and a wide variety of microenvironmental cues such as extracellular matrix (ECM), growth factors, and cytokines (Fig. 1). Although illuminating, very few, if any, studies have been conducted using ER⁺ breast cancer models, despite the fact that metastasis dormancy is most common in this tumor subtype. In this review, we describe the clinical manifestation of ER⁺ dormant metastasis. We then discuss the urgent need, possible solutions, and the conceptual challenges faced by basic and clinical scientists who want to study metastasis dormancy in ER⁺ breast cancer. Several other excellent reviews of tumor dormancy are also recommended (2, 4–9).

Clinical manifestation of dormancy of ER⁺ breast cancer

Tumor dormancy is defined as clinically undetectable microscopic metastases that eventually progress to overt cancer after a long period of time. In breast cancer, metastases usually become evident asynchronously with the primary tumor, and they show variable lengths of time to become clinically apparent. This lag time depends to some extent on the initial tumor volume or stage at first diagnosis. At one end of the spectrum, neglected locally advanced cancers are often diagnosed with metastases already evident or appearing soon. At the other end, metastases of low-stage breast cancer may occur many years after the diagnosis. Other factors also relate to the time until metastases are identified. Such factors as the rate of proliferation or the ER status of the tumor are also related to the time to distant recurrence. Indeed, late recurrences, sometimes decades after the initial primary diagnosis, indicating a long
dormant period, present a significant clinical challenge mainly for ER\(^+\) breast cancer. More than half of the recurrences of ER\(^+\) tumors occur 5 years or longer after diagnosis and surgical removal of the primary tumor, and some patients suffer recurrence after more than 20 years (10–12). This is in sharp contrast to ER\(^-\)/C0 tumors, for which the recurrence rate peaks at around 2 years but diminishes to a low rate after 5 years (13). Current prognostic markers often focus on and are reasonably good at predicting early recurrences within 5 years (14–17), but the risk of late recurrences remains poorly predictable. Is late recurrence merely a reflection of very slowly proliferating ER\(^+\) cancer cells lying in distant sites or due to cancer cells actually entering a period of total dormancy in which they stop proliferating until they are activated to grow again by some as yet unknown stimulus. We know that many patients with early breast cancer harbor micrometastases in their bone marrow. Although the presence of these cells predicts a slightly worse prognosis, the majority of the patients with these disseminated cells never suffer a recurrence, at least within the time of follow-up on the studies (7, 18).

Systemic therapy given to patients with early breast cancer after surgery in an attempt to eradicate micrometastases (adjuvant therapy) could also influence the time it takes for a patient to show a recurrence by slowing cell proliferation or by killing the majority but not all of the metastatic cells, thereby delaying their clinical appearance. Adjuvant chemotherapy is effective in reducing recurrences within the first 5 years but ineffective in preventing late recurrences between 5 and 15 years (1). These data suggest that late recurrences arise from residual disease that is treatment resistant.

Further insights may be obtained from studies of adjuvant endocrine therapies in patients with ER\(^+\) tumors. These studies have shown that longer and longer durations of treatment are better at reducing recurrence and prolonging survival than shorter durations. Early clinical trials indicated that 5 years of tamoxifen treatment was more effective than 1 or 2 years (1). Recent studies of prolonging adjuvant tamoxifen from 5 to 10 years show a reduction in recurrence and death between years 10 and 15, suggesting that longer treatment continued to suppress proliferation of micrometastases still viable after just 5 years of treatment (see Table 1 and references therein). These results indicate that inhibition of ER signaling may suppress the exit of cancer cells from dormancy or growth inhibition but may not kill them. Whether ER\(^+\) micrometastases are ever totally eradicated by treatment will require very long follow-up of patients to ascertain.

**Biologic Insights into ER\(^+\) Metastasis Dormancy**

**Biologic models of metastasis dormancy**

Before this discussion, it is necessary to define the key parameters of an experimental model for ER\(^+\) metastasis dormancy. Such models need to have several key features: (i) recapitulation of several characteristics of human ER\(^+\) tumors, including estrogen dependence, growth inhibition by antiestrogen strategies, as well as the potential to develop resistance to these treatments; (ii) recapitulation of the natural progression of ER\(^+\) tumors, including tumorigenesis, local invasion and intravasation, and the temporal kinetics and anatomical site of metastasis (predominantly bone); and (iii) opportunities to investigate the roles of major cell types that may be involved in dormancy. In subsequent
paragraphs, we go over the major models and techniques that have been used in breast cancer and point out their strengths and weaknesses for dormancy research. It needs to be noted here that although the abovementioned properties are highly desirable, models lacking these features may still generate useful information. For instance, late recurrences are not exclusively ER$, and the mechanistic insights obtained from ER$ models may also be relevant to ER$ diseases.

The most widely used breast cancer models are human breast cancer cell lines that are largely derived from pleural effusions of patients with advanced breast cancer. Recent genomic studies support the validity of cell lines as breast cancer models by showing the common genomic/gene expression profiles shared between primary tumors and cancer models by showing the common genomic/gene expression profiles shared between primary tumors and cohorts of cell lines (19). In particular, ER$ breast cancer cell lines have been essential for the elucidation of steroid hormone–stimulated signaling and resistance mechanisms of antihormonal therapies (20). When transplanted into the mammary glands of immunodeficient mice, ER$ cancer cells can generate orthotopic xenograft tumors, which are typically dependent on estradiol supplementation. However, such tumors rarely give rise to spontaneous metastases during the relatively short lifespan of the mouse and therefore fall short of serving as models of metastasis dormancy. As an alternative approach, direct introduction of ER$ cancer cells via intracardiac injection delivers them throughout the entire arterial circulation and can result in colonization in multiple organs, including bone. Other limitations of xenograft models include the lack of an intact immune system in immunodeficient mice, the use of already fully transformed and metastatic cancer cells, and the absence of potential influence from the primary tumor (21). Despite these caveats, important discoveries pertinent to metastasis dormancy were made using this approach. In one example, a subpopulation of MDA-MB-231 cells that outgrew after a long dormancy in animals was compared with the parental population. VCAM1 was identified as a key molecule that could serve as a chemoattractant for osteoclast precursors, which in turn trigger the proliferation and exit from dormancy of cancer cells (Fig. 2A; ref. 22). The role of VCAM1 was verified in several other models, including MCF-7 cells (22). Therefore, the observation may be relevant to ER$ breast cancer.

Murine cell lines derived from mouse mammary tumors have also been widely used to investigate breast cancer. As opposed to xenograft models, these models provide the advantage of using immunocompetent syngeneic mice. Considering the growing evidence for the role of immune cells in tumor progression (23), this is a potential major advantage for such models. The disadvantage of the vast majority of murine cell line models is the lack of estrogen dependence. Nevertheless, important progress has been made in understanding metastasis dormancy using murine cell lines. One recent study using this model revealed the bone morphogenetic protein (BMP) regulator Coco as a mediator driving the outgrowth of micrometastases into macrometastases in lungs (Fig. 2B; ref. 24). Another study identified Irf7 as a suppressor of spontaneous bone metastases, apparently by regulating the adaptive immune system (25). How relevant these findings are to ER$ human breast cancer remains to be tested.

Transgenic and other genetically engineered mouse models (GEMM) have also been developed to enhance our understanding of mechanisms driving tumor progression and metastases. These models offer the obvious strengths of capturing the full course of tumor progression from a premalignant stage to the terminal metastatic stage. They also preserve the native interactions between cancer cells and their microenvironment at every stage. Like murine cell lines, however, GEMMs are rarely ER$
and estrogen dependent and, therefore, cannot be directly used for studies of ER⁺ breast cancer dormancy. Moreover, with few exceptions, GEMMs, unlike ER⁺ breast cancers, predominantly metastasize to lungs but not bones within the lifespan of the tumor-bearing mice (26). Despite these weaknesses, several groups have created elegant models that exhibit relevant features, including local and metastasis dormancy. In a series of studies, deinduction of an inducible MMTV-Neu oncogene resulted in tumor regression. The regressed tumor, however, still contains viable tumor cells that remain dormant but can be activated by several processes and factors, including expression of Snail, a driver of epithelial–mesenchymal transition (EMT). This finding provided a connection between dormancy and a cellular program that was later linked to traits of cancer progenitor (stem) cells (27, 28). A more recent study showed that Ron receptor kinase suppresses T cells and promotes the outgrowth of micrometastases in lungs, highlighting the important role of the immune system in tumor dormancy (29). Again, whether these findings can be applied to human ER⁺ tumors is not known.

In addition to these models, patient-derived primary xenografts generated by directly transplanting pieces of surgically removed tumors to immunocompromised mice preserve histopathologic features, genomic and transcriptomic similarities, and pharmacologic responses similar to the original tumors in the patient. A minority of these primary xenografts maintain ER expression and ER dependence. Importantly, many such xenografts spontaneously spread to the bone marrow and therefore provide relevant sources of quiescent disseminated tumor cells (DTC; refs. 30–32).

In summary, although significant progress has been made in our understanding of metastasis dormancy, individual biologic models cannot cover every essence of this process, especially for ER⁺ breast tumors. To overcome this barrier, researchers may need to combine different types of models and create novel ones. Like many other biologic questions, the establishment of better models may benefit from our deeper understanding of the objectives. One possible solution is to divide the dormancy process into separate steps, conquer each of them separately, and finally synthesize our knowledge to construct an ideal experimental model. In the next section, we divide the problem of dormancy into a few conceptual components, summarize our current understanding of each, and propose future directions.

**Biologic mechanisms underlying metastasis dormancy**

We envision that the dormancy process is composed conceptually of several components, including cell survival mechanisms that constantly sustain the viability of cancer cells, self-renewal mechanisms that maintain tumorigenesis capacity, and activation/suppression mechanisms that restore/prevent aggressive outgrowth. Very importantly, all of these mechanisms are likely to involve cross-talk between dormant cells and their microenvironment. We focus on bone as the host tissue of dormant cancer cells because it is the most frequent metastatic site of ER⁺ breast cancer and is the major reservoir of DTCs.

Upon arrival in the bone marrow, cancer cells need to extravasate from the blood circulation and enter the area...
close to the interface between the bone marrow and bone matrix. The extravasation process is not expected to be a major hurdle for cancer cells because of the sinusoid structures in the bone marrow, which are extensively fenestrated with 1-μm-wide pores. The cancer cells need to exploit the foreign microenvironment for survival. This can be viewed as a Darwinian selection process during which only a small fraction will succeed. Our previous study suggested that Src is a survival mediator specifically in bone colonization in that it potentiates PI3K/AKT activation in cancer cells by CXCL12 and IGF1, both of which abound in the bone marrow microenvironment. Although the cell models we used were ER−, it is noteworthy that most ER+ tumors also exhibit high Src activity, in part due to the interaction between ER and Src (33). Thus, our results provide one possible mechanism underlying the survival of cancer cells in bones. This mechanism links the intrinsic survival machinery to unique features of the bone microenvironment. Further investigations are needed to confirm this mechanism or to discover additional ones because the survival mediators represent ideal therapeutic targets to permanently eradicate dormant cancer cells. As a result, short-term intervention could generate long-lasting effects, thereby alleviating concerns about side effects and prohibitive cost of treatment.

It has long been suspected that dormant cancer cells are also tumor-initiating cells or cancer stem cells, although direct evidence is still lacking. Conceptually, if a dormant cell maintains the potential to eventually reinitiate a tumor, it should be a tumor-initiating cell by definition. However, are dormant tumor-initiating cells that ultimately initiate bone metastases the same population of tumor-initiating cell that maintains the tumorigenesis potential of the primary tumor? Many recent studies suggest that the potential of initiating a tumor can be dynamically acquired or lost and therefore represents a cellular status rather than a cellular entity (34–36). In this scenario, the tumor-initiating potential needs to be suppressed to ensure dormancy or activated to exit dormancy. A recent study revealed that BMP2 signaling in the lung microenvironment inhibits tumor-initiating potential. The acquisition of Coco expression terminates metastasis dormancy and drives lung metastasis outgrowth (Fig. 2B; ref. 24). In another study, cancer cells disseminated to lungs were shown to induce the expression of Periostin (POSTN) in fibroblasts, which reciprocally increases WNT signaling and promotes tumor-initiating potential and tumor outgrowth (37). Similarly, lung micrometastases can also produce tenasin C (TNC), an ECM protein, that facilitates maintenance of tumor initiating potential by activating WNT and Notch pathways in cancer cells (38). The above-mentioned molecules and the underlying pathways may represent therapeutic targets to diminish tumor-initiating potential of dormant cancer cells. However, whether similar mechanisms are operative in bone and for ER+ breast cancer cells remains unknown.

The termination of dormancy not only needs the restoration of tumor-initiating potential but also requires strong proliferative signals, which may already be in place but actively suppressed by the host (e.g., immunosurveillance) or need to be acquired de novo. Several recent studies provide interesting examples of this process. In one study, dormant cancer cells are proposed to activate osteoclast progenitors via soluble VCAM1 as a nonconventional chemoattractant. The activated osteoclasts in turn initiate the osteolytic vicious cycle and trigger the outgrowth of cancer cells (Fig. 2A; ref. 22). In a second study, the authors showed that cancer cells in lungs need to form filopodia-like protrusions with ECM to grow out. The formation of these structures activates the integrin/FAK/ERK signaling cascades and drives cancer cell proliferation (39). The perivascular niche may also play a role in regulating the maintenance and exit of dormancy. Ghajar and colleagues showed that endothelial cells secrete thrombospondin 1 to suppress cancer cell proliferation, and this suppression is lost in sprouting neovascularature, which leads to outgrowth (40). Again, whether and how these processes are similar in ER+ tumors in bones after long-term dormancy remains to be investigated.

In summary, although insights have been gained into the dormancy process, it remains unexplored whether these discoveries can be applied to ER+ models under a dormant setting in bone. A key piece of information missing in our understanding of ER+ metastasis is the biology of bone micrometastases. It is well established that in the advanced stage, bone metastases are typically osteolytic and driven by a vicious cycle between cancer cells and osteoclasts. However, it is also evident that many patients do not exhibit symptoms of osteolyis for many years before bone relapses. How cancer cells exist as microscopic lesions and how they crossstalk with the bone microenvironment in the preosteolytic stage remains completely elusive. DTCs may represent cancer cells in this stage, but direct evidence is lacking. Several questions need to be addressed regarding preosteolytic micrometastases (Fig. 3). First, do they have preferential microenvironmental niches in bone and bone marrow? Some recent studies using prostate cancer models suggest that cancer cells tend to lodge in the same niches as hematopoietic stem cells (HSC; ref. 41). However, the definition of the HSC niche is itself a biologic question under intensive study. Second, are these micrometastases truly quiescent or are they slowly proliferating (Fig. 1)? Third, are these micrometastases dormant or indolent because they are enriched with tumor-initiating cells or because they are restricted by the bone microenvironment. Finally, are there biologic pathways that specifically mediate their survival and proliferation? The answers to these questions will shed light on the dormancy behavior of ER+ breast cancer, and they warrant intensive investigation.

Clinical Challenges of ER+ Dormancy

Detection of micrometastases

The current detection of metastases in the clinic primarily relies on the appearance of symptoms caused by macrometastases. As mentioned above, bone metastases cause osteolysis in their advanced stage, which is usually manifested by skeletal-related events such as bone pain, fractures,
and hypercalcemia. At the time of diagnosis, the vicious cycle between cancer cells and osteoclasts has already started, and the dormancy stage has been terminated. Some metastases are asymptomatic and are detected by tests such as X-rays, computed tomography (CT), or positron emission tomography with fluorodeoxyglucose (FDG-PET). However, microscopic metastases composed of only a few cancer cells are below the detection threshold of these technologies. Thus, we have not been able to identify micrometastases in the clinic unless they are discovered “accidentally” in a patient undergoing a bone marrow biopsy for research or an unrelated purpose. We may be approaching the ability to detect submillimeter or even single-cell metastases owing to the development of several new technologies. For instance, high-resolution hyperpolarized 3He MRI can detect micrometastases that are 300 μm in diameter in preclinical models (42). Circulating tumor cells (CTC) and tumor DNA in patient peripheral blood may represent useful surrogates of micrometastases in distant organs. Recently, therapeutic and biologic insights have been obtained by monitoring and characterizing CTCs (43) and their released DNAs (44) in small cohorts of patients. Molecular characterization of CTCs from the same blood sample has shown heterogeneous subpopulations with different transcriptional profiles and hormone receptor phenotypes (43, 45–47). Heterogeneity in ER expression of CTCs coexisting within a single blood draw was observed in our study (Fig. 4). The exact relation between CTCs and dormant micrometastases remains to be experimentally elucidated. DTCs in the bone marrow have been considered as dormant bone metastases and mark a poorer prognosis (7, 48, 49). However, the correlation between these cells and bone relapses has not been firmly established. In some studies, patients with detectable DTCs and those with CTCs only partially overlap and exhibit different prognoses (50–52). Further investigations are needed to provide a deep understanding of the biologic nature of DTCs and CTCs and to answer the question of whether they represent or can be used as surrogates for dormant micrometastases.

Prediction of late recurrences

Early relapses within 5 years can be predicted with reasonable accuracy using histopathologic features and/or gene expression profiles. Current successful predictors share the commonality of having some components reflecting proliferation, suggesting that early relapses may be caused by the rapid proliferation of residual tumor cells (14–17). Late recurrences beyond 5 years, however, apparently are not associated with proliferation markers and have been difficult to predict (53). The ratio of two individual mRNAs, HOXB13/IL17BR, has been shown to be a prognostic factor in ER+ and node-negative patients, which is the major population giving rise to late recurrences (54, 55). Gene expression signatures that are retrospectively associated with late recurrences have recently been derived by comparing the gene expression profiles of primary tumors of early versus late recurrences (56, 57), or using dormant cancer cells in experimental systems (58). Whether these signatures can prospectively predict late recurrences remains to be tested. The difficulty of predicting recurrences beyond 5 years may be anticipated for biologic reasons. The use of information derived from the primary tumor to predict recurrences is based on the assumption that the
traits driving recurrences are encoded in the bulk of primary tumors. While this assumption has been validated for early recurrences in numerous studies, it might need to be reconsidered in the case of late recurrences. The fact that dormant micrometastases stay in distant organs for many years suggests a long evolutionary process of these cells after their departure from the primary tumor. During this time, independent genetic and epigenetic traits may arise and drive the recurrences that will not be present in the original primary tumors (59). Thus, it may be necessary to examine the metastases from patients with late relapses and compare them with the primary tumor and early metastases to decipher the genetic or epigenetic alterations that lead to late recurrences. Other factors may also confound the prediction of late recurrences. Some late recurrences may exceed the detection threshold but are asymptomatic and undetected till the patients are deceased. These cases may therefore be wrongly excluded from the late recurrence category. On the other hand, patients who succumb to early recurrences may harbor dormant metastases at different sites, which would have progressed further had the patient survived. Therefore, it remains a challenge to accurately classify primary tumors as to their propensity for late recurrences.

**Treatment and prevention of late recurrences**

As mentioned earlier, roughly 20% to 40% of patients with ER\(^+\) breast cancer eventually develop distant metastases and half of these events occur 5 years or later after diagnosis of the primary tumor. However, the proportion of patients carrying dormant cells may be much higher if in some patients these cells never progress to become macrometastases during the patient’s lifetime, and even if they do, the resultant slowly growing metastases may remain asymptomatic and undetected. A recent example of a patient cared for by one of the authors (C.K. Osborne) who was found on a CT scan ordered for a kidney stone to have a 5-cm asymptomatic osteolytic lesion in the pelvis proven on biopsy to be an ER\(^+\) breast cancer metastasis developing 22 years after her primary tumor diagnosis illustrates this possibility. Even proposing to evaluate asymptomatic patients looking for occult metastasis presents practical and ethical dilemmas. With the rapid development of imaging and other detection technologies, it is foreseeable that we will be able to detect submillimeter or even single cancer cells in the future. However, key questions remain: (i) How many of the detected dormant metastases will progress to threaten patients’ lives; (ii) How can we distinguish the dormant metastases that are likely to progress from those that are not; and (iii) What therapeutic strategies can be used to eradicate them or prevent their progression? The answers to these questions require us to directly investigate dormant cancer cells. Thus, there are urgent needs to establish biologic models of dormant metastases in the laboratory and obtain patient-derived dormant cancer cells from the clinic. These efforts will open several diagnostic and therapeutic avenues. First, characterization of dormant cancer cells may lead to identification of highly specific serum markers of dormant cancer cells, including DNA, RNA, proteins, and metabolites, that are released to the circulation. Second, identification and investigation of survival pathways that sustain dormant cancer cells may lead to therapies that eradicate dormant cancer cells. Finally, elucidation of how dormant cancer cells progress to resume aggressive growth may help the prevention of overt metastases and facilitate the prediction of which dormant metastases need to be proactively managed in the clinic. Our current knowledge is far from sufficient to achieve these goals. Therefore, preclinical and
clinical studies on dormant micrometastases are imperative—
ously needed to promote our understanding of these critical
issues and ultimately to prevent late recurrences and further
improve the outcome of patients with breast cancer.

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