Bone metastasis-related signaling pathways in breast cancers stratified by estrogen receptor status

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Abstract

Background: Breast cancer bone metastasis (BCBM)-specific genes have been reported without considering biological differences based on estrogen receptor (ER) status. The aims of this study were to identify BCBM-specific genes using our patient dataset and validate previously reported BCBM-specific genes, and to determine whether ER-status-related biological differences matter in identification of BCBM-specific genes.

Methods: We used Affymetrix GeneChips to analyze 365 primary human epidermal growth factor receptor 2 (HER2)-negative invasive breast cancer specimens. Genes that were differentially expressed between patients who developed bone metastasis and those who developed non-bone metastasis were identified using Cox proportional hazards model, and differential expression of gene sets was assessed using gene set analysis. We performed gene set analysis to determine whether biological function associated with bone metastasis were different by ER status using 2,246 functionally annotated gene sets assembled from Gene Ontology data base.

Results: Among 16,712 probe sets, 592 were overexpressed in the bone metastasis cohort compared to the non-bone-metastasis cohort (false discovery rate ≤ 0.05). However, no BCBM-specific genes met our significance tests when the cancers were stratified by ER status. In ER-positive and ER-negative breast cancers, 151 and 125 gene sets, respectively, were overexpressed for BCBM and the majority of BCBM-related pathways were different. Of significant gene sets, only 13 gene sets were overlapped between ER-positive and -negative cohorts.

Conclusion: ER-positive and ER-negative breast cancers have different biological pathways in BCBM development. We have yet to explore BCBM-related biomarkers and targets considering the biological features associated with BCBM depending on the ER status.

Key words: Breast cancer, bone metastasis, gene expression, gene sets, estrogen receptor status.

Introduction

Bone is the first site of metastasis in 26% to 50% of breast cancer patients [1-4]. However, the mechanism of breast cancer bone metastasis (BCBM) has not been clarified. In the past decade, microarray analysis has enabled successful prediction of patients who have a high risk of recurrence [5, 6], but the sites of metastasis are not predictable. If we could identify patients likely to develop BCBM in the future, we might be able to assess the effect of a bisphosphonate in the adjuvant setting to reduce the incidence of bone
metastases for these high risk population.

Smid et al. [7] have studied genes associated with breast cancer metastatic to bone. Using 107 primary breast cancer samples, they found 69 genes significantly overexpressed in patients whose relapses manifested in bone compared to those whose relapses manifested in other sites (non-bone). In that study, there was a significant difference in estrogen receptor (ER) positivity between patients with bone metastasis and those with non-bone metastasis (82.6% vs 60.5%, P = 0.019). This difference indicates that the significant difference in gene expression between the patient groups may have been strongly affected by ER status, which is the only clinical marker associated with metastasis to bone [8].

This significant difference in ER status between bone and non-bone metastases has not been considered in previous studies identifying BCBM-specific genes. None of the previously published gene sets have demonstrated that there are different biological features in ER-positive and ER-negative breast cancers that develop BCBM. Therefore, we hypothesized that ER-positive and ER-negative breast cancers have a different biological pathway in development of BCBM.

The first aim of this study was to identify BCBM-specific genes using our patient dataset and validate previously reported BCBM-specific genes. The second aim was to determine whether the biological function associated with bone metastasis were different by ER status in identification of BCBM-specific genes by gene set analysis.

Materials and methods

Patient data and tumor samples

Tumor samples were collected prospectively from 365 patients with primary invasive breast cancer who underwent surgery from November 1999 to August 2008 at The University of Texas MD Anderson Cancer Center. We excluded patients who were diagnosed with ductal carcinoma in situ, metaplastic carcinoma, sarcoma, or clinical stage IV breast cancer. Patients with human epidermal growth factor receptor 2 (HER2)-positive breast cancer were also excluded because whether they received trastuzumab may have been affected occurrence of metastasis. Participation was voluntary, and all patients gave written informed consent. The study was performed under institutional review board approval.

Tumor samples were obtained from fine-needle aspiration (FNA) of the primary tumor before initiation of any systemic therapy. The following clinical data were collected from MD Anderson’s Breast Cancer Management System database.

Metastases were confirmed by imaging and/or bone biopsy. The time to metastasis and site of first metastasis were recorded. We classified the samples into three cohorts: bone as the first metastatic site, non-bone as the first metastatic site, and no metastasis (no mets). The bone cohort consisted of patients whose first metastases manifested in bone or in bone and other sites at the same time. The non-bone cohort consisted of patients whose first metastases manifested only in other sites.

Statistical analysis

Patient characteristics by type of first metastasis were summarized with medians and ranges for age and follow-up time and with frequency and percentage for all other characteristics. Bone metastasis-free survival was defined as the time interval from the diagnosis of primary breast cancer until the diagnosis of bone metastasis or the last follow-up date. Non-bone metastasis-free survival was defined as the time interval from the diagnosis of primary breast cancer until the diagnosis of non-bone metastasis or the last follow-up date. We compared the bone and non-bone cohorts according to clinical and tumor characteristics of patients using Fisher’s exact test. All statistical tests were two-sided.

Gene expression data

These datasets are referred to as the MicroArray Quality Control Phase II (MAQC-II) Project (GSE16716) that has been published. We extracted a part of the dataset which included HER2-negative patients treated with weekly paclitaxel (T) (80 mg/m2) for 12 treatments, followed by 5-fluorouracil (500 mg/m2), doxorubicin (50 mg/m2), and cyclophosphamide (500 mg/m2) for four treatments, given once every 21 days (BAC), with available clinical data. Samples were placed in RNAlater (Ambion, Austin, TX) storage reagent and stored at ~80°C until gene expression analysis. As described by Hess et al., all gene expression data were generated using Affymetrix U133A gene chips (Affymetrix, Inc, Santa Clara, CA) [9]. Gene expression data were normalized with the MA55 algorithm, mean centered to 600, and log2 transformed before further analysis. Probe sets with the lowest 25% mean expression value were removed from all higher level analyses to reduce noise from low expressed probe sets. This left 16,712 probe sets for analysis. Tumors with normalized ESR1 mRNA expression (probe set 205225_at) >10.18 were considered ER-positive [10]. Tumors with normalized HER2 mRNA expression (probe set 216836_s_at) >12.54 were considered HER2-positive [10]. All statistical analysis was performed with BRB-ArrayTools version 3.8.1 α software and R.

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Identification of differentially expressed genes and gene sets

We compared gene expression data among the bone, non-bone, and no-mets cohorts without any further stratification by clinical or molecular class using the Cox proportional hazards model to test whether expression of a particular gene significantly influences development of metastasis. We also performed similar class comparisons separately for ER-positive and ER-negative cancers. False discovery rates (FDRs) were calculated to adjust for the multiple comparisons using BRB-ArrayTools. The FDR was calculated with the significance analysis of microarrays (SAM) tool as the median number of false-positive genes from permutation testing divided by the number of nominally significant genes defined from the unperturbed data [11]. We mapped significantly differentially expressed genes to functional pathways using Ingenuity pathway analysis (IPA) software (http://www.ingenuity.com) [12, 13].

We next examined differential expression of prior defined gene sets using gene set analysis (GSA) [12, 13] to assess whether members of a set of genes that corresponded to a particular biological pathway tended to occur toward the top or the bottom of a rank-ordered gene list (rank ordered by differential expression among the bone, non-bone, and no-Mets cohorts) [14]. We tested 2,246 different, functionally annotated gene sets from the Gene Ontology (GO) database that collectively represent most known biological pathways in eukaryotic cells. Gene sets with a minimum of 10 genes and maximum of 100 genes were selected for inclusion in this analysis. We used the Efron and Tibshirani GSA method that employs “maxmean” statistics and is implemented by BRB-ArrayTools [15, 16]. We determined whether gene sets were differentially expressed among cohorts, with statistical significance being determined by a permutation test (n = 1,000). The null hypothesis was that the average degree of differential expression of members of a given gene set among the bone, non-bone, and no mets cohorts is the same as expected from a random set of genes of similar size.

Results

Patient characteristics

Clinicopathological characteristics of the 365 patients are summarized in Table 1. The bone metastasis cohort consisted of 34 patients (9.3%); the non-bone metastasis cohort, 32 patients (8.8%); and the no-mets cohort, 299 patients (81.9%).

We compared the bone and non-bone cohorts according to ER status. One hundred eight seven of the 214 with ER-positive tumor (87.3%) received hormonal therapy. There was a trend of high occurrence of ER-positive breast cancer among patients with bone metastasis compared to those with non-bone metastasis (55.9% vs 31.3%, P = 0.052). This trend was similar to that shown in Smid et al.’s study [7]. Other characteristics of the patients did not differ significantly between patients with bone metastasis and those with non-bone metastasis.

| Characteristics | All patients | No mets | Bone | Non-bone | P value* (bone vs non-bone) |
|-----------------|-------------|---------|------|----------|---------------------------|
| No. of patients | 365 | 100% | 299 | 100% | 34 | 100% | 32 | 100% |
| Median age (years) | 51 | 51 | 52 | 51 |
| Range (years) | 24-85 | 25-82 | 24-85 | 29-73 |
| Clinical stage | | | | |
| 0 | 1 | 0.3% | 0 | 0.0% | 1 | 2.9% | 0 | 0.0% | 1 |
| I | 18 | 4.9% | 18 | 6.0% | 0 | 0.0% | 0 | 0.0% |
| II | 204 | 55.9% | 180 | 60.2% | 12 | 35.3% | 12 | 37.5% |
| III | 142 | 38.9% | 101 | 33.8% | 21 | 61.8% | 20 | 62.5% |
| Tumor grade | | | | |
| Poor | 230 | 63.0% | 183 | 61.2% | 21 | 61.8% | 26 | 81.3% | 0.057 |
| Good to moderate | 134 | 36.7% | 116 | 38.8% | 13 | 38.2% | 5 | 15.6% |
| Unknown | 1 | 0.3% | 0 | 0.0% | 0 | 0.0% | 1 | 3.1% |
| ER status | | | | |
| Positive | 214 | 58.6% | 185 | 61.9% | 19 | 55.9% | 10 | 31.3% | 0.052 |
| Negative | 151 | 41.4% | 114 | 38.1% | 15 | 44.1% | 22 | 68.8% |
| Neoadjuvant chemotherapy | | | | |
| Received | 338 | 92.6% | 277 | 92.6% | 30 | 88.2% | 31 | 96.9% | 0.357 |
| Not received | 27 | 7.4% | 22 | 7.4% | 4 | 11.8% | 1 | 3.1% |
| Adjuvant chemotherapy | | | | |
| Received | 56 | 15.3% | 45 | 15.1% | 5 | 14.7% | 6 | 18.8% | 0.748 |
| Not received | 309 | 84.7% | 254 | 84.9% | 29 | 85.3% | 26 | 81.3% |

ER = estrogen receptor. *Fisher’s exact test.
Site-specific relapse-free survival according to ER status

Next, we examined the relapse-free survival of patients according to bone metastasis or non-bone metastasis as a first metastatic site in ER-positive and ER-negative breast cancers using the Cox proportional hazards model. When we evaluated bone-metastasis-free survival, there was no significant difference in survival times between patients with ER-positive and ER-negative breast cancers. However, when we evaluated non-bone-metastasis-free survival, patients with ER-negative breast cancer had shorter metastasis-free survival than did patients with ER-positive breast cancer. These results indicated that bone and non-bone metastasis have a distinct clinical course relative to ER status, which might come from distinct biological differences related to ER status. Therefore, we sought to assess the gene expression associated with BCBM in ER-positive and ER-negative breast cancers separately.

Gene expression associated with BCBM

To determine the differences between bone metastasis and non-bone metastasis, individual gene (i.e., probe set)-level analysis was performed using the Cox proportional hazards model for 14 genes that have been reported to have an important role in development of BCBM [17-32]. In 66 patients with metastasis, without any stratification for ER status, 6 probe sets—BMP1, BMP10, TNFRSF11A, TGFβ2, CXCR4, and ESR1—were significantly overexpressed in the bone metastasis cohort compared to the non-bone metastasis cohort, with FDR ≤ 0.05. However, when this analysis was done in ER-positive and ER-negative breast cancers separately, no gene (FDR ≤ 0.05) was found to be significantly overexpressed in both patient populations. This result may suggest that previously reported BCBM-specific genes might be capturing gene differences based on ER status, not BCBM status.

Next, to examine possible differences between BCBM and non-bone metastasis, we performed the same analysis using all 16,712 probe sets. In 66 patients with metastasis without any stratification by ER status, 592 probe sets were overexpressed in the bone metastasis cohort compared to the non-bone metastasis cohort with FDR ≤ 0.05 (Table 2). However, no BCBM-specific genes were found when ER-positive and ER-negative breast cancers were analyzed separately.

Table 2. Gene sets overexpressed in the bone-metastasis cohort compared with the non-bone-metastasis cohort

| FDR       | All (n=66) | ER positive (n=29) | ER negative (n=37) |
|-----------|-----------|-------------------|-------------------|
| ≤ 0.3     | 6,427     | 3,617             | 0                 |
| ≤ 0.1     | 2,785     | 0                 | 0                 |
| ≤ 0.05    | 592       | 0                 | 0                 |
| ≤ 0.01    | 2         | 0                 | 0                 |

FDR = false discovery rate; ER = estrogen receptor.

Gene set analysis

Validation of previously published BCBM-specific gene sets

Because no statistical differences in any individual gene expression level between bone and non-bone metastases were shown among ER-positive or -negative patients, we determined to, using gene set enrichment analysis, detect consistent but relatively small-scale differences in the expression of gene sets that belong to functional pathways [12, 13]. We tested five gene sets that were previously reported to be associated with BCBM from breast cancer [7, 23, 32-34]. Four of the five BCBM-specific gene sets were not validated in our patients for their predictive value for occurrence of BCBM. Only the BCBM-specific gene set from the study by Kang et al. [33], which was detected using the breast cancer cell line MDA-MB-231, was significantly overexpressed in the bone cohort (Table 3). Also, when we compared the bone and non-bone cohorts by ER status separately, in ER-positive patients, only that gene set was significantly overexpressed in the bone cohort. None of the five sets were overexpressed in ER-negative patients.

Table 3. Comparison of overexpression of previously reported BCBM-associated gene sets between the bone-metastasis and non-bone-metastasis cohorts when stratified by ER status

| Gene sets       | Number of genes | All P value (direction) | ER positive P value (direction) | ER negative P value (direction) |
|-----------------|----------------|------------------------|--------------------------------|--------------------------------|
| Kang et al [33] | 15             | 0.013 (Bone)           | 0.028 (Bone)                   | 0.259 (Bone)                   |
| Wilson et al [34]| 31             | 0.164 (Others)         | 0.238 (Bone)                   | 0.397 (Others)                 |
| Nannuru et al [32] | 23             | 0.448 (Bone)           | 0.378 (Bone)                   | 0.13 (Bone)                    |
| Zhang et al [23] | 36             | 0.193 (Bone)           | 0.225 (Bone)                   | 0.391 (Others)                 |
| Smid et al [7]  | 86             | 0.001 (Others)         | 0.029 (Others)                 | 0.166 (Others)                 |

*Efron-Tibshirani’s GSA test.
Identification of biological difference between ER status

The outcome that we could not identify BCBM-specific genes when patients were stratified by ER status indicated the possibility of the presence of a biological difference between ER status in development of BCBM. Therefore, for the second aim of our study, we sought to detect, using GSA, differences in biological features related to developing BCBM between ER-positive and ER-negative breast cancers. We also determined 2,246 functionally annotated gene sets assembled from Gene Ontology (http://www.geneontology.org/) to examine possible differences between bone and non-bone cohorts over a broad range of biological processes (Table 4). At a $P$ value of $\leq 0.05$ by Efron-Tibshirani test, 151 gene sets in ER-positive breast cancer and 125 gene sets in ER-negative breast cancer were overexpressed in the bone metastasis cohort compared to the non-bone metastasis cohort. Only 13 gene sets were common to both ER-positive and ER-negative breast cancers. These results demonstrated that the majority of bone-metastasis-related pathways were differently expressed in ER-positive (138/151, 91.4%) and ER-negative breast cancers (112/125, 89.6%).

Table 4. Number of gene sets overexpressed in the bone metastasis cohort when stratified by ER status

| $P$ value | ER positive | ER negative | Common |
|-----------|-------------|-------------|--------|
| 0.1       | 313         | 261         | 49     |
| 0.05      | 151         | 125         | 13     |
| 0.01      | 22          | 21          | 0      |

ER = estrogen receptor.

To show the biological process, we mapped the top 10 canonical pathways significantly overexpressed for the bone metastasis cohort in ER-positive breast cancers and in ER-negative breast cancers using the top 20 gene sets overexpressed in ER-positive breast cancers and ER-negative breast cancers by Ingenuity pathway analysis, separately (Figure 1 and 2). Cellular growth and proliferation, as well as intracellular and second messenger signaling pathways were overexpressed for the bone cohort in ER-positive breast cancer (Figure 1). Nuclear receptor signaling and cytokine signaling-related pathways were overexpressed in ER-negative breast cancer (Figure 2). Kang et al. [33] have reported that transforming growth factor β (TGF-β) signaling plays an important role for BCBM; in our study, the TGF-β signaling pathway was the 2nd most highly overexpressed pathway for the bone cohort in ER-negative breast cancer.

Figure 1. The top 10 canonical pathways by significantly overexpressed genes associated with bone metastasis in ER-positive breast cancers assessed by Ingenuity pathway analysis software results. The bars show logarithms of $P$ values that were calculated by Fisher’s exact test. The line indicates the ratio for the percentage of genes in a pathway that were found in our uploaded list. The threshold indicates the line at $P = 0.05$. 

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Discussion

We demonstrated that there are no unique genes that predict BCBM when we stratified the ER status. This is partly due to the major biological difference between ER-positive and ER-negative groups in development of BCBM, as we hypothesized. Notably, many genes were overexpressed in the bone metastasis cohort as a whole, without stratification by ER positivity, as described in previous studies. However, when gene-expression-level analysis was performed with ER status stratification, there were no bone-metastasis-specific genes. Therefore, these results suggest that ER-positive and ER-negative breast cancers have a biological difference in their mechanism to develop BCBM.

When we validated five previously reported bone-associated gene sets, one showed significant association with BCBM in ER-positive breast cancer in our patients. Kang et al. identified that set of 122 bone-metastasis-specific genes from tumor cells of BCBM of mice injected with MDA-MB-231 cells [33]. The top 11 genes overexpressed in BCBM in the study were validated as significant in ER-positive but not in ER-negative breast cancer in our validation analysis.

From 107 primary breast tumor samples, Smid et al. [7] found 69 genes significantly differentially expressed between patients who experienced relapse to bone and those who experienced relapse to non-bone sites. Combining their 69-gene list with that of Kang et al. [7, 23, 32-34] revealed that the fibroblast growth factor receptor-p42/44 mitogen-activated protein kinase (FGFR-MAP) K signaling pathway was in preference of tumor cells that metastasize to bone in an animal model system. They developed a classifier of 31 genes, which in an independent validation set correctly predicted all tumors spreading to bone with a specificity of 50%. They suggested that this model could predict BCBM more accurately than ER status. However, we could not identify a BCBM-specific gene signature by gene-expression-level analysis using our dataset in either ER-positive or ER-negative breast cancers because of the very subtle differences in the expression of genes.

Kang et al. [33] also reported that the TGF-β signaling pathway is associated with occurrence of BCBM, inducing overexpression of interleukin-11 (IL-11) and CTGF, which were the most highly overexpressed genes in osteolytic metastasis. TGF-β not only facilitates tumorigenesis and producing osteoclasts [35, 36] but also causes osteolytic change by inducing expression of PTHrP [31] and IL-11 [37], RANKL [27], and matrix metalloproteinases [25, 29]. Furthermore, TGF-β stimulates angiogenesis [38] and induces epithelial-to-mesenchymal transition [39, 40]. Similar to Kang et al.’s description [33], the TGF-β signaling pathway was significantly overexpressed for our bone cohort in ER-negative breast cancer compared to our non-bone cohort (P < 0.05).

There are some limitations of exploratory research using microarray analysis for prediction of BCBM. Only a few genes that were detected to be associated with BCBM overlapped among previous studies [7, 23, 32-34]. In addition, many genes relating to proteins such as PTHrP, IL-8, insulin-like growth factor (IGF-1), and platelet-derived growth factor (PDGF), which have been reported as important factors in development of BCBM, were not identified.
as bone-metastasis-specific genes in any studies [35, 36]. The possible reasons for the differences in gene signatures among the studies may be due to the various backgrounds of the experiments [41]. For example, tumor samples in various studies were taken from a breast cancer cell line [33], mouse model [32, 34], or human samples of primary tumor (including stage IV cancers) [7] or of metastatic sites [23]. No previous studies analyzed the gene expression in ER-positive and -negative breast cancers separately. Samples were collected by FNA or dissection [7, 23], analyzed for late-onset BCBM [23] or BCBM occurring at any time course, and compared between the tumor-bone interface and tumor itself [32], or among bone metastasis, non-bone metastasis, and no metastasis. These various experimental backgrounds may have induced the different results. Reproducibility of microarray analyses may be another issue [42].

We demonstrated no bone-metastasis-specific genes for early-onset BCBM. But because of the short median follow-up time and small number of events, we still need to determine whether such a specific gene can predict for late onset of BCBM. Because the number of patients with ER-positive breast cancer who develop metastasis will continue to increase 5 years after treatment [43, 44], we need a definitive study with a longer follow-up time and a larger number of events to assess the biological difference in developing BCBM between ER-positive and ER-negative breast cancers.

Our results indicate that the biological differences between ER-positive and ER-negative breast cancers affect development of BCBM. Therefore, we need to assess the biological features associated with BCBM depending on the ER status.

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Competing Interests

The authors have declared that no competing interest exists.

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