**Introduction**

Despite the fact that overall five years survival of breast cancer has recently improved to 98% in the United States (1), the survival of patients with metastatic disease remains low at only 23%, which accounts for approximately 40,000 deaths annually (2,3). It has been estimated that 20-30% of women with early stage breast cancer will eventually develop metastatic disease. The lung is the second most common anatomic site of first exclusive distant metastasis of breast cancer (4), thus advances in the understanding and management of lung metastases are expected to have a large impact on breast cancer survival (5). Despite the central role of mouse models in breast cancer research, these models have not been critically evaluated for their appropriateness for the study of lung metastasis (2-15,16-31).

A commonly employed murine model to study breast cancer lung metastasis entails injection of cancer cells via the tail vein (TV) to implant cells in the lung, and thus produce lung tumors (TVt) (32-34). Advocates of TV argue that it is an easy and quick method to form metastatic TVt lesions, especially with cell lines that take long periods of time to metastasize, if at all. However, others have argued that this model may not adequately mimic human metastatic breast cancer because it does not follow the biological steps that a primary tumor must take to produce a distant metastatic tumor (33,34), and it ignores the cross-talk between primary and metastatic lesions (32,35-38). Indeed, it has been reported that lung metastatic tumors (LMet), which progressed biologically from...
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a primary tumor generated by orthotopic implantation (OS) of breast cancer cells have a different morphology compared to TVt tumors. Although there has been debate in the literature regarding advantages and disadvantages of TV implantations versus orthotopic implantation, evaluation by gene expression profiling of tumors from these models or of the metastatic lesions which they produce has not previously been examined (39).

Using genome-wide gene expression microarrays, we have now found that primary OS tumors and their lung metastatic lesions have differentially expressed genetic profiles, but lung metastatic lesions produced by TV or OS have similar profiles.

Materials and methods

Virginia Commonwealth University Institutional Animal Care and Use Committee (IACUC) approval was obtained for all experiments. Female Balb/c mice, 12 weeks of age, weighing approximately 20 g were obtained from Harlan Laboratories (Frederick, MD). The 4T1-luc2, adenocarcinoma cell line derived from the mammary glands of Balb/c mice and genetically manipulated to overexpress the firefly luciferase gene was obtained from Caliper Life Sciences (Hopkinton, MA). The cells were cultured in RPMI media, suspended at a concentration of 1×10^6 cells/100 µL, and 10 µL of this solution were then injected, unless otherwise stated.

All cell implantations were performed under isoflurane anesthesia using sterile technique. Orthotopic implantation under direct vision (OS; Figure 1A): a 5 mm incision was made medial to the nipple, and a cotton swab was used to expose the mammary gland. The cells were implanted directly into the mammary gland under direct vision, using x10 microscopic magnification, and the wound was closed. Subcutaneous implantation (Sq): under anesthesia the skin was tented up and 10 µL of the 4T1-luc2 cells were implanted into the subcutaneous space. Tail vein injection (TV; Figure 1B): 100 µL of 1×10^5 cells/100 µL were injected into the median tail vein.

Xenogen’s IVIS® 200 and Living Image® software (Caliper Life Sciences, Hopkinton, MA) were used to quantify the photon/sec emitted by 4T1-luc2 cells which reflects tumor burden after 200 µL of luciferin (Fisher Scientific, Inc.) was injected intraperitoneally. The quantification of photons emitted allowed for the quantification of tumor burden and cancer progression in vivo. To compare tumor growth and survival, 16 Balb/c mice in 2 experimental groups were used: OS and TV.

Gene expression profiling of 4T1 cells in vitro, OS and Sq tumors, and metastatic tumors

Ten days after OS or Sq (8 Balb/c mice per group, one implantation site each), the tumors were harvested and snap-frozen at –80 °C. Day ten after implantation was chosen based upon our previous study (36). RNA Extraction: Snap-frozen tissues were used for histopathological scoring after hematoxylin and eosin (H&E) staining of standard features performed on frozen sections adjacent, above and below the tissue used for RNA isolation. All samples contained more than 70% tumor. Total RNA was extracted and the quality evaluated using a method of sample processing established previously in our laboratory (40). Total RNA was extracted from multiple 10-µm thick frozen tissue sections using the MagMAX™-96 for Microarrays Total RNA Isolation Kit (Invitrogen™ Life Technologies, Carlsbad, CA), in an automated fashion using the magnetic particle processor MagMAX™ Express. RNA purity was judged by spectrophotometry at 260, 270, and 280 nm. RNA integrity was assessed by running 1 µL of every sample in RNA 6000 Nano LabChips® on the 2100 Bioanalyzer (Agilent Technologies, Foster City, CA). Gene expression microarray analyses: The Affymetrix® protocol (Affymetrix, Santa Clara, CA) has been previously described (40). Briefly, starting from 500 ng of total RNA, cDNA synthesis and cRNA labelling were performed using the GeneChip® 3’ IVT Express Kit (Affymetrix). Ten µg of fragmented cRNA were hybridized on the GeneChip® Mouse Genome 430A 2.0 array for 16 hrs at 60 rpm in a 45 °C hybridization oven. This array provides a comprehensive coverage of the transcribed murine genome by
including over 22,600 probe sets that analyze the expression level of over 14,000 well-characterized mouse transcripts. The arrays were washed and stained with streptavidin phycoerythrin (SAPE; Molecular Probes, Eugene, OR) in the Affymetrix fluidics workstation. Every chip was scanned at a high resolution, with pixel locations ranging from 2.5 µm down to 0.51 µm, by the Affymetrix GeneChip® Scanner 3000 according to the GeneChip® Expression Analysis Technical Manual procedures (Affymetrix). After scanning, the raw intensities for every probe were stored in electronic files (in .DAT and .CEL formats) by the GeneChip® Operating Software (GCOS v1.4) (Affymetrix). The overall quality of each array was assessed by monitoring the 3’/5′ ratios for a housekeeping gene (GAPDH) and the percentage of “Present” genes (%P) where arrays exhibiting GAPDH 3′/5′<3.0 and %P>40% were considered good quality arrays.

Statistical methods

For photon emission, Student’s t-test statistical analysis was utilized. For the microarray data analysis, standard statistical methods were utilized, including Robust Multi-array Analysis (RMA) method (41), hierarchical cluster analyses, and false discovery rates (FDR) as previously reported (42).

Results

Implantation methods

To evaluate how appropriate the commonly used mouse models are for studying breast cancer lung metastasis, we standardized our methodology based on the most commonly used implantation methods in the literature. OS is implantation of 4T1-luc2 cells into the mammary gland after exposing it via a small incision (Figure 1A), and TV is injection of the cells directly into the middle TV (Figure 1B).

TV obviates the biologic sequence of lung metastasis arising from a primary tumor and can produce tail injection site tumors

Bioluminescent technology allows detection of 4T1-luc2 cells in any part of the body due to the whole body scan. Figure 2 demonstrates bioluminescence imaging of lung metastasis 10 days after OS implantation (LMet). LMet tumors were obscured by OS tumor in the chest mammary gland (Figure 2A), which can be visualized in situ after removal of OS tumor (Figure 2B) or ex vivo (Figure 2C). Note that lung metastasis arising from OS (LMet) produced discrete tumors (Figure 2B,C). The cancer cells metastasized to the lung 7 to 10 days after OS implantation, which is consistent with our previous results (36,38). In contrast, lung implantation of 4T1-luc2 cells (TVt) was immediately confirmed after TV injection in the tail (Figure 2D). Note that TVt produced disseminated 4T1-luc2 cell implants throughout the lungs without discrete tumors. Additionally, we occasionally observed development of “tail tumor” at the injection site, which was easily detected with bioluminescent whole body scan.

Histological analysis using H&E stain showed that OS lung metastases produced isolated LMet tumors (Figure 3A,B), in contrast to the TVt lung tumors in which the cancer cells colonized the lungs along the normal parenchymal architecture of the lung without forming isolated tumors (Figure 3C,D).

Genomic expression profiles were different between 4T1-luc2 cells in culture dish, Sq and OS primary tumors, and OS lung metastases (LMet), but not different between lung metastases after TV (TVt) and OS (LMet)

To evaluate the genomic profiles of the tumors produced by these methods, genome wide microarray analysis was performed of the 4T1 cells in culture, Sq and OS tumors as well as of the TVt and LMet metastases. Ten thousand three hundred fifty probe sets (45.7% of transcriptome) were significantly differentially
expressed between 4T1 cells in culture compared to OS and Sq tumors, TVt and LMet metastases (FDR≤1%; P<0.0001). In addition, between Sq and OS tumors, 700 probe sets were differentially expressed (FDR≤15%; P<0.01). Furthermore, 1,247 probe sets were differentially expressed (1.5-fold; P<0.01) between OS primary tumors and LMet metastases, without any difference within each group. We also found that the same 1,247 probe sets were differentially expressed (1.5-fold; P<0.01) between OS primary tumors and TVt lung metastases without any difference within each group. Finally, there was no significant difference in gene expression between TVt and LMet metastases (Figure 4).

Discussion

Breast cancer drug development is an expensive and inefficient process, and use of animal models for the screening of novel agents is a major component of it. However, the literature has not critically examined how appropriate these models are for the study of breast cancer metastasis (6-16,17-31). In translational research it is important to consider the degree to which breast cancer metastasis models provide clinically relevant endpoints (6,8-10,30,31,43-52). The most commonly used animal models for screening for anti-cancer drug development are Sq for local breast cancer and TV for breast cancer lung metastasis. Regardless of the model employed to evaluate the efficacy of novel therapeutics, it is critical to understand the limitations of each method.

It is important to appreciate to what extent the animal model produces clinically relevant endpoints that are translatable to human breast cancer. As we have demonstrated, TV does indeed produce TVt lung metastasis quickly after injection (Figure 2), but there are important limitations to consider. First, TVt obviates the biologic progression from primary tumor to distant lung metastasis and it will not evaluate the efficacy of the therapeutic agents that target the process of cancer progression.

Figure 3. H&E stain histological analysis of lung tumors. Low magnification (×200) (A) and high magnification (×400) (B) of LMet metastasis after OS implantation show an isolated lung tumor. Low magnification (×200) (C) and high magnification (×400) (D) of TVt tumor after TV implantation demonstrate disseminated colonization of 4T1-luc2 cells throughout the lungs along the parenchymal architecture without forming isolated tumors.
to metastasis. Second, TV produces TVt lung metastasis without the presence of a primary tumor in the mammary gland. Human metastatic breast cancer patients are commonly treated with systemic therapy before the primary breast cancer is removed, which will not be modeled by TV which lacks a primary tumor. Situations where patients will be treated for metastatic breast cancer without a primary tumor include cases after urgent palliative mastectomy due to bleeding or ulceration of the primary tumor in the setting of metastatic disease, as well as cases of recurrence in the lung after mastectomy. Both situations present biological systems which are very different than the mere intravenous injection of cancer cells resulting in lung implantation. By producing lung metastasis without a primary tumor, TV ignores the cross-talk between primary and metastatic lesions (32,35-38). This element of breast cancer biology is not only important from a biological or basic science perspective, but it also has important implications in the clinical management of human breast cancer. In fact, there are ongoing clinical trials to evaluate the effect on metastatic progression and overall survival of mastectomy in patients with metastatic breast cancer (38). The third factor is the morphology of the TVt lung metastases (Figures 2,3). TVt lung metastases are diffusely disseminated throughout the lung because the cancer cells colonize the lung via hematogenous embolization. In contrast, OS produce discrete LMet metastatic tumors which progressed to the lung along the primary-tumor-to-distant-metastasis progression pathway. In humans, breast cancer forms lung metastases as discrete lung nodules which progress by pathways more analogous to LMet than TVt. Understanding this difference may have implications in terms of the bioavailability and drug delivery of therapeutics in the solid tumor versus the disseminated cells, the relationships of the nodules to the blood and lymphatic vessels in the lung, and the effects on mortality. In fact, mortality in TV typically follows a sudden death due to a thromboembolic phenomenon, rather than mortality via a more gradual process due to cancer progression and overall tumor burden, which occurs in LMet and human breast cancer (6). This factor cannot be underscored enough because the final translatable clinical endpoint, i.e. survival, to screen for the efficacy of novel therapeutics before entering clinical trials varies significantly between these methods.

In the era of targeted therapeutics, it is important to consider the genomic profiles of the tumors and lung metastases produced. The differential expression of over 10,000 genes between cells in culture and the tumors and lung metastases of the same cell line reinforces the importance of in vivo screening of the efficacy of novel therapeutics (Figure 4). In addition, the differential expression of so many genes between OS primary tumors and LMet metastases also reinforces the importance of
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not just evaluating the efficacy in primary tumors. Furthermore, the differential expression of so many genes between Sq and OS tumors implies that investigators should understand the differences in the potential target expressions of the tumors their model produces. Finally, there was no significant difference in the genomic expression profiles of TVt versus LMet metastases, which is an important strength of the TV model, especially in light of the limitations stated above. Based upon our findings, we cannot help but speculate that tumor microenvironments have a significant role in the gene expression profiles.

The choice of the appropriate model for breast cancer research relies upon the underlying hypothesis being tested. Understanding the limitations of these models, beyond the differences in genetic profile, is therefore of great importance. The benefit of Sq is that it produces tumors which can be followed locally for therapeutic effect, but they do not metastasize to the lung. The strength of the OS model is that it utilizes the biologic progression from primary tumor to distant metastasis, and it allows for testing hypotheses on primary-metastatic tumor interactions. It is also beneficial to evaluate the effect of tumor microenvironment especially when syngeneic cells are used. The weakness of this model is that some cell lines, especially utilizing xenograft models where human cell lines are implanted into mice, do not readily metastasize to the lung. TV has the strength in that it will implant cell lines immediately into the lungs without relying on metastasis from primary lesions that will significantly shorten the duration of the experiment, but it has no primary lesion for questions related to primary tumor-metastatic lesion interactions. In addition, TV colonizes the lung with cancer cells, rather than producing isolated lung “tumors”. This is an advantage for researchers since it allows them to quantify the amount of lung metastasis utilizing colony formation assay, which is an established commonly used method for 4T1 cells. On the other hand, TV often causes sudden death via thromboembolism, instead of by cancer progression, which makes it an unstable model to assess survival. As is the case for any animal model, TV should be utilized with its advantages and limitations in mind in order to translate the findings to human breast cancer lung metastasis. Strengths and weaknesses of these models are summarized in Table 1.

In summary, our study suggests that primary tumors and their lung metastatic lesions have differentially expressed genetic profiles, although there are no differences between metastatic lesions produced by TV injection versus orthotopic implantation of the same cancer cells. Although the TV injection method is limited because it evaluates lung metastasis without a primary tumor, ignores the biologic progression from primary tumor to metastatic lesions, and produces lung metastasis with a different morphology and mechanism of mortality than the orthotopic implantation method and human cancer, it does produce lung metastases with similar genomic profiles as lung metastases after orthotopic implantation. Utilizing these animal metastatic breast cancer models with an understanding of their limitations is expected to improve the efficiency of breast cancer drug development and the advancement of breast cancer research.

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References


