Transformation of enriched mammary cell populations with polyomavirus middle T antigen influences tumor subtype and metastatic potential

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Abstract

Introduction: Breast cancer exhibits significant molecular, histological, and pathological diversity. Factors that impact this heterogeneity are poorly understood; however, transformation of distinct normal cell populations of the breast may generate different tumor phenotypes. Our previous study demonstrated that the polyomavirus middle T antigen (PyMT) oncogene can establish diverse tumor subtypes when broadly expressed within mouse mammary epithelial cells. In the present study, we assessed the molecular, histological, and metastatic outcomes in distinct mammary cell populations transformed with the PyMT gene.

Methods: Isolated mouse mammary epithelial cells were transduced with a lentivirus encoding PyMT during an overnight infection and then sorted into hormone receptor–positive luminal (CD133+), hormone receptor–negative luminal (CD133−), basal, and stem cell populations using the cell surface markers CD24, CD49f, and CD133. Each population was subsequently transplanted into syngeneic cleared mouse mammary fat pads to generate tumors. Tumors were classified by histology, estrogen receptor status, molecular subtype, and metastatic potential to investigate whether transformation of different enriched populations affects tumor phenotype.

Results: Although enriched mammary epithelial cell populations showed no difference in either the ability to form tumors or tumor latency, differences in prevalence of solid adenocarcinomas and squamous, papillary, and sebaceous-like tumors were observed. In particular, squamous metaplasia was observed more frequently in tumors derived from basal and stem cells than in luminal cells. Interestingly, both molecularly basal and luminal tumors developed from luminal CD133+, basal, and stem cell populations; however, luminal CD133− cells gave rise exclusively to molecularly basal tumors. Tumors arising from the luminal CD133−, basal, and stem cell populations were highly metastatic; however, luminal CD133+ cells generated tumors that were significantly less metastatic, possibly due to an inability of these tumor cells to escape the primary tumor site.

Conclusions: Expression of PyMT within different mammary cell populations influences tumor histology, molecular subtype, and metastatic potential. The data demonstrate that luminal CD133+ cells give rise to less metastatic tumors, luminal CD133− cells preferentially establish basal tumors, and the cell of origin for squamous metaplasia likely resides in the basal and stem cell populations.

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Introduction

The classification of breast cancer into several distinct molecular and histological subtypes can provide information to help guide patient therapy and predict outcome [1, 2]. Tumors that retain histological and molecular attributes of normal tissue are considered well differentiated and are generally less aggressive and correlate with better patient prognosis. In contrast, the loss of normal tissue structure and the dysregulation of genes involved in modulating growth and differentiation indicate transition of the disease into a more advanced stage [3]. A better understanding of tumor etiology and processes that control the transition between early and advanced states of breast cancer may improve strategies for detection, treatment, and prevention of the disease.

How a particular cell responds to a transforming event, its susceptibility for malignant progression, and its role in establishing a tumor’s histological and molecular fate are poorly understood. The mammary gland is a complex tissue composed of two distinct cell lineages, the luminal epithelium and myoepithelium, with each lineage encompassing a hierarchy of cells at various states of differentiation [4, 5]. When a normal cell is transformed, preexisting signaling networks intrinsic to that particular cell type may become dysregulated and contribute to tumor growth and progression. For example, tumors classified as a hormone-receptor positive subtype express estrogen receptor (ER or Esr1) and are generally dependent on estrogen for growth, recapitulating characteristics of a subset of normal, luminal cells found in the breast [6]. Molecular similarities are also observed between other normal cell populations and cancer subtypes. Mammary stem cells have a gene expression signature similar to spinloid and claudin-low tumors [7-9], whereas the molecular signature of normal luminal progenitors is associated with basal-like breast cancer [8]. Even differentiated mammary epithelial cells (MECs) share molecular features with a cancer subtype. Tumors histologically classified as lipid-rich carcinoma of the breast express metabolic and differentiation markers observed in alveolar cells, the milk-producing cells of the mammary gland [10-12]. These molecular associations suggest that tumors can arise from different cell populations and maintain signaling networks of their cell of origin.

Although transforming events within a cell may initiate and drive the process of tumor progression, these events do not necessarily establish a tumor’s histological or molecular fate. Data derived from The Cancer Genome Atlas demonstrate that major oncogenic drivers and tumor-associated mutations are broadly represented among breast cancer subtypes, suggesting that diverse mechanisms contribute to a tumor’s phenotype [13]. Such mechanisms may include cell-intrinsic networks acquired from the cell of origin. In support of this, several studies have demonstrated that normal cell types can respond to the same oncogenic pathway in unique ways. Ince et al. used different cell culture conditions to enrich for either luminal-like or myoepithelial-like human breast cell populations [14]. Both populations were subsequently transformed with a common set of oncogenic drivers. When transplanted, each precursor cell population generated a distinct tumor phenotype, with notable differences in tumor histology and metastasis. The myoepithelial-like cells established tumors similar to squamous carcinoma of the breast, whereas the luminal-like cells generated papillary adenocarcinomas [14]. Other studies with mouse mammary tumor models have also supported a role for the cell of origin in controlling tumor fate. Differences in tumor phenotype were observed when a Brca1 mutation was induced in different normal cell populations. Conditional Brca1 loss of function targeted to the basal compartment using Keratin-14-Cre Brca1^{10/10} p53^{+/−} mice established tumors that were primarily adenoid squamous carcinomas and adenomyoepitheliomas. In contrast, disruption of Brca1 in luminal progenitors using Blg-Cre Brca1^{10/10} p53^{+/−} mice resulted in ductal carcinomas with a basal-like molecular subtype [15]. These studies suggest that intrinsic differences between cell populations may influence the histopathology of the tumors they generate.

The polyomavirus middle T antigen (PyMT) oncogene has been used extensively in mice to model breast cancer [16, 17]. In these models, PyMT drives transformation of MECs by signaling through several pathways, including Src, Ras, and phosphoinositide 3-kinase [18-21], resulting in a phenotype similar to ErbB2/Neu-induced tumors [7, 22, 23]. Transgenic mice that express PyMT under control of the promoter derived from the long terminal repeat (LTR) of the mouse mammary tumor virus (MMTV) develop mammary tumors that undergo progressive transition from precancerous lesions to late-stage malignant tumors and exhibit a high frequency of metastasis [16, 24, 25]. Tumor progression is marked by a loss of both myoepithelial cells and ER+ luminal cells [24], and a concomitant expansion of cells expressing the luminal progenitor marker CD61 [26]. Through intrinsic gene set analysis and hierarchical clustering of gene expression profiles, MMTV-PyMT tumors have been classified within the luminal subgroup [7, 9]. In addition, a close association between the molecular signature derived from luminal progenitors and MMTV-PyMT tumors has been described [9]. Similar to other mouse models that function through ErbB/Ras signaling proteins, the predominant tumor histology in the MMTV-PyMT
model is solid adenocarcinoma. However, varied histopathology is observed, with approximately 30% of tumors having papillary, glandular, or acinar features and 10% exhibiting either adenosquamous, pilar, or type P histology [22].

These data demonstrate that the MMTV-PyMT model can establish tumors with both a luminal phenotype and diverse histopathology. It is unknown whether these characteristics are a result of the activity of the MMTV LTR within a particular cell type or through unique transforming activity of the PyMT oncogene. Because the MMTV LTR is widely active in mammary epithelium and drives expression at early stages of postnatal development [27–29], it is difficult to identify the cell of origin for tumors in this model. In order to assess how cellular context affects tumor progression, the PyMT oncogene has been targeted to various MEC populations by uncoupling expression of the oncogene from the MMTV LTR. For example, virus-based approaches have been used to express PyMT either ubiquitously in all mammary cell populations or specifically within distinct cell types, and these studies have shown that tumor histology and molecular subtype can vary as a result of the targeting approach [12, 30, 31]. The restricted expression of PyMT in the keratin 6 (K6) mammary cell population resulted in tumors with predominantly papillary and cystic histology and a distinct normal-like molecular subtype, representing a phenotype dissimilar to what has been observed in the MMTV-PyMT mouse model [27]. In contrast, non-cell-type-specific expression of PyMT in MECs using the ubiquitous elongation factor 1 alpha (EF1α) promoter generated tumors with diverse histology, including a high frequency of adenosquamous carcinomas and the occurrence of a unique lipid-rich carcinoma [12]. In addition, this model produced tumors that classified within both luminal and basal molecular subgroups [12]. In the present study, we extended upon these studies by evaluating the latency, histopathology, molecular subgroup, and metastatic potential of tumors derived from four different fluorescence-activated cell sorting (FACS)-enriched MEC populations. The results suggest that the originating cell population influences several tumor characteristics and further implicates a cell residing predominantly in the basal and stem cell compartment as the cellular origin for squamous metaplasia.

Methods
Mice
FVB/NJ mice were obtained from The Jackson Laboratory (Bar Harbor, ME, USA) and maintained in a pathogen-free facility. The University of Utah Institutional Animal Care and Use Committee approved mouse handling and procedures.

Generation of mouse mammary tumors
MECs were collected from 8–10-week-old FVB/NJ mice as described previously [12]. Then, freshly isolated MECs were infected with EF1α-PyMT-ZsGreen lentivirus overnight at 37 °C as described previously [12]. Following infection, cells were washed five times with Hanks’ balanced salt solution (HBSS; Gibco/Thermo Fisher Scientific, Grand Island, NY, USA) and incubated with 0.05% trypsin-ethylenediaminetetraacetic acid (EDTA; Gibco/Thermo Fisher Scientific, Grand Island, NY, USA) to isolate single cells. Trypsin was inactivated with MEC media [12], and cell clumps were removed by straining MECs through a 40-μm cell strainer (Falcon; Fisher Scientific, Pittsburgh, PA, USA). MECs were then resuspended in wash buffer (HBSS + 2% fetal bovine serum; HyClone Laboratories/GE Healthcare, Logan, UT, USA) and kept on ice for antibody staining and FACS.

Staining consisted of six tubes: (1) no antibody control, (2) CD24-V450 control, (3) CD49f-phycoerythrin (CD49f-PE) control, (4) CD133-allophycocyanin (CD133-APC) control, (5) 7-amino-actinomycin D (7-AAD) control, and (6) CD24-V450/CD49f-PE/CD133-APC/7-AAD sample. During antibody staining, control tubes contained 5 × 10⁶ cells and the sample tube contained 20 × 10⁶ cells resuspended in 200 μl of wash buffer. All antibodies were obtained from BD Pharmingen (San Diego, CA, USA) and were used at a 1:100 dilution. After the primary antibodies were added, cells were incubated on ice for 15 minutes. Following incubation, cells were washed with 1 ml of wash buffer and centrifuged at 1000 × g for 2 minutes. Stained MECs were then resuspended in wash buffer and sorted into luminal CD133+, luminal CD133−, basal, and stem cell populations as described previously [32–35] on a BD FACSARia Cell Sorter using FACSDiva version 6.1.3 software for analysis (BD Biosciences, San Jose, CA, USA). Isolated MEC populations were kept on ice until transplantation.

For each transplantation, 1 × 10⁵ untransduced and unsorted MECs were mixed with 2 × 10⁷ transduced luminal CD133+, luminal CD133−, basal, or 5 × 10⁵ stem enriched MECs. MECs were then resuspended in 10 μl of Matrigel (BD Biosciences, San Jose, CA, USA) by transplantation, and the Matrigel cell mixture was injected into the fourth cleared inguinal mammary fat pad of 3-week-old FVB/NJ mice. Only a single fat pad was injected per mouse. Tumor growth was monitored, and tumors were collected upon reaching 2 cm in diameter. Once tumors were harvested, viable cells were collected using the same protocol for MEC isolation and then frozen in freeze media as described previously [12]. Portions of the tumors were also flash-frozen for RNA isolation using a Qiagen RNeasy kit (Qiagen, Valencia, CA, USA), and additional tumor fragments were processed for paraffin embedding.
Infection, cell sorting, and transplantation experiments were performed over two rounds. Each time, 10 transplants were performed per sorted MEC population, for a total 20 transplants per group.

Antibody staining and histology
Portions of transduced and FACS-sorted MECs were used to quantify basal and luminal cell enrichment. For each isolated population, 1 x 10^6 MECs resuspended in 200 μl of wash buffer were centrifuged (Cytospin 4 Cytocentrifuge; Thermo Scientific, Hudson, NH, USA) onto slides (Shandon Cytoslide; Thermo Scientific, Hudson, NH, USA) at 900 rpm for 10 minutes. Cytospun cells were then incubated with fixative solution (4 % paraformaldehyde in phosphate-buffered saline [PBS]) for 15 minutes and washed five times with PBS for 5 minutes each. Fixed cells were permeabilized for 10 minutes with 0.2 % Triton X-100 (Sigma-Aldrich, St. Louis, MO, USA) in PBS, washed with 1 % bovine serum albumin (BSA; EMD Millipore, Billerica, MA, USA) in PBS, and blocked with 1 % BSA in PBS for 10 minutes. Cells were then incubated with primary antibodies against keratin 14 (K14, 1:400 dilution, rabbit, PRB-P-100; Covance, Princeton, NJ, USA) and keratin 8 (K8, 1:50 dilution, rat, TROMA-I; Developmental Studies Hybridoma Bank, Iowa City, IA, USA) for 1 h at room temperature. Following incubation, slides were washed with 1 % BSA in PBS and stained with 4',6-diamidino-2-phenylindole dihydrochloride and secondary antibodies Alexa Fluor 594 chicken anti-rat immunoglobulin G (IgG, 1:1000 dilution; Invitrogen/Life Technologies, Carlsbad, CA, USA) and Alexa Fluor 488 goat anti-rabbit IgG (1:1000 dilution; Invitrogen/Life Technologies, Carlsbad, CA, USA).

Paraffin-embedded tumor samples were processed and subjected to hematoxylin and eosin (H&E) staining, ESR1 immunohistochemistry (IHC), and cytokeratin staining as described previously [12]. The following primary antibodies were used: K14 (1:400 dilution, rabbit, PRB-P-100; Covance), K8 (1:50 dilution, rat, TROMA-I; Developmental Studies Hybridoma Bank, Iowa City, IA, USA), and ESR1 (1:200 dilution, MC-20, sc-542; Santa Cruz Biotechnology, Santa Cruz, CA, USA). Secondary antibodies included Alexa Fluor 594 chicken anti-rat IgG (1:1000 dilution; Invitrogen/Life Technologies, Carlsbad, CA, USA), Alexa Fluor 488 goat anti-rabbit IgG (1:1000 dilution; Invitrogen/Life Technologies, Carlsbad, CA, USA), and biotin-SP-conjugated protein (1:1000 dilution; Jackson ImmunoResearch Laboratories, West Grove, PA, USA).

All immunofluorescence imaging was performed on an Olympus IX81 microscope (Olympus, Tokyo, Japan) using a Hamamatsu Photonics ORCA-ER camera (Hamamatsu Photonics, Hamamatsu City, Japan). Fluorescence image recording and processing were performed using SlideBook 64 version 5.0.0.24 software (Intelligent Imaging Innovations, Denver, CO, USA). Slides processed for IHC and H&E staining were imaged on an Olympus BX50 microscope with a Canon EOS Rebel XSI camera using EOS imaging software (Canon, Melville, NY, USA). Any changes in contrast and brightness were performed using Photoshop CS4 software (Adobe Systems, San Jose, CA, USA) on entire images to enhance appearance without altering image content.

In vitro and in vivo assessment of tumor growth with estrogen receptor inhibition
For in vitro dose–response studies, tumor cells and MECs were isolated using the same tissue dissociation protocol as described above. Single cells were suspended in Matrigel (BD Biosciences, San Jose, CA, USA) and 10 μl of a cell/Matrigel mixture was plated per well in a 96-well plate (Costar, Corning Life Sciences, Oneonta, NY, USA). 4-Hydroxytamoxifen (Sigma-Aldrich, St. Louis, MO, USA) was dissolved in ethanol and serially diluted in MEC media. Cells were dosed with 100 μl of media containing drug or vehicle control for 48 h. Media with drug or vehicle control was refreshed every 24 h. Each drug concentration was tested in triplicate. Cell viability was measured using the ATPlite assay (PerkinElmer, Waltham, MA, USA) according to the manufacturer’s instructions and normalized to vehicle control.

Tumor growth dependence on estrogen was tested in vivo by conducting an additional primary cell infection, FACS, and transplantation experiments as described above. Ovaries were removed from all mice that received cell transplants, as described previously [36]. Ten surgeries were performed per infected and sorted population.

Microarray analysis
Flash-frozen tumors were randomly selected from each of the tumor groups for RNA extraction and microarray analysis. Total RNA was isolated using the Qiagen RNeasy kit.

To perform supervised hierarchical clustering, all steps of microarray processing, data filtering and normalization, and analysis were performed as described previously [12]. Batch adjustment was performed in two batches. A dataset generated by Herschkowitz et al. was treated as one batch (Gene Expression Omnibus [GEO] accession number [GEO:GSE3165]) [7], and data generated at the Huntsman Cancer Institute (HCI) was treated as a second batch. The HCI microarray dataset has been deposited in the National Center for Biotechnology Information GEO database under accession number [GEO:GSE64453].

To perform unsupervised hierarchical clustering, quantile-normalized, log-scaled microarray intensity data were hierarchically clustered in R using Ward’s method. The resulting dendrogram revealed a marked batch effect related to the date on which the arrays were
processed. Each class of samples (luminal CD133−, luminal CD133+, basal, and stem) were adjusted for batch effect separately using the ComBat procedure [37] and then recombined. Differential expression analysis comparing the luminal CD133− samples with the remaining samples identified 1111 microarray probes representing 1046 unique genes showing at least two-fold differential expression at an adjusted \( p \) value <0.05 (\( p \) value by \( t \) test with adjustment using the Benjamini-Hochberg method).

**Assessment of tumor metastasis**

As described above, transplantation experiments were performed twice for each transduced and enriched MEC population. Lung metastasis for the first round was assessed by H&E staining. Lung tissue processing and staining were performed as described above. Paraffin-embedded lungs were serially sectioned at 10 \( \mu \)m, and every fifth slide was stained and examined for metastasis. For the second round of transplants, lung metastases were analyzed by fluorescence imaging after lungs were flattened between two glass slides. Slides were imaged as described above, and numbers of unique metastatic sites and tumor areas were quantified using ImageJ software (National Institutes of Health, Bethesda, MD, USA). Prevalence of lung metastases and numbers of metastatic foci were consistent over two rounds of transplants.

To quantify circulating tumor cell (CTC) numbers, fresh whole blood was collected by cardiac puncture immediately after mice were killed according to University of Utah–approved Institutional Animal Care and Use Committee procedures. CTCs were isolated for FACS analysis as described previously from mice bearing primary tumors, as well as from no-tumor control mice [38]. CTCs expressing ZsGreen were detected by analyzing cells using a FACScan cytometer (BD Biosciences, San Jose, CA, USA), and results were quantified using FlowJo software (Tree Star, Ashland, OR, USA). Owing to low numbers of CTCs present within isolated whole blood, the ZsGreen-positive threshold was set at 0.05 % of no-tumor control background signal. This threshold was then used as a baseline for detecting CTCs in tumor-bearing mice. All CTC values were then normalized to no-tumor control background signal.

Tail vein injections were performed to assess the ability of tumor cells to colonize the lungs after introduction into the bloodstream. Single cells were isolated from primary tumors using the same procedure employed for MEC isolation. Cells were then resuspended in HBSS at 10 \( \times \) 10^6 cells/ml. A quantity of the HBSS/cell mixture (250 \( \mu \)l; 2.5 \( \times \) 10^5 cells) was injected into the lateral tail veins of 8–12-week-old FVB/NJ mice. Cells isolated from individual tumors were injected into five mice each. Twenty days postinjection, mice were killed and tumor lung foci numbers were quantified by fluorescence imaging with ImageJ software.

**Analysis of PyMT expression in transduced mammary epithelial cells**

Freshly isolated MECs were infected in suspension overnight and then transferred to adherent culture in MEC media the following day. Separate plates of cells were analyzed for ZsGreen and PyMT expression expression each day for 5 days. Protein lysates were prepared for simple Western blot analysis to detect PyMT expression, and cells were analyzed for ZsGreen expression by flow cytometry. For cytometry, cells were washed with PBS, lifted with 0.25 % trypsin-EDTA (Gibco/Thermo Fisher Scientific), fixed in 2 % paraformaldehyde, and resuspended in wash buffer on ice. Single cells were isolated by straining them through a 40-\( \mu \)m cell strainer (Falcon; Fisher Scientific) and immediately analyzed with a BD LSR II flow cytometer (BD Biosciences, San Jose, CA, USA) for ZsGreen expression.

**Real-time quantitative PCR analysis**

RNA was isolated from freshly sorted MECs or flash-frozen tumors using the Qiagen RNeasy Mini Kit. For cDNA synthesis from tumor RNA, 1 \( \mu \)g of total RNA was reverse-transcribed using the iScript Reverse Transcription Supermix for real-time quantitative PCR (RT-qPCR) synthesis kit (Bio-Rad Laboratories, Hercules, CA, USA) according to the manufacturer’s instructions. For sorted MECs, cDNA was synthesized from 20 ng of total RNA via reverse transcription, followed by preamplification with pooled PrimeTime assays (Integrated DNA Technologies, Coralville, IA, USA) for genes of interest using the Qiagen RT² PreAmp cDNA Synthesis Kit. Genes investigated were K8 (assay identification: Mm.PT.58.6862465), K14 (Mm.PT.58.43652691), progesterone receptor (Pgr; Mm.PT.58.10254276), Esr1 (Mm.PT.58.8025728), p63 (Mm.PT.58.13970687), Slug (Mm.PT.58.43645779), c-Kit (Mm.PT.56a.33701407), and glyceraldehyde 3-phosphate dehydrogenase (Gapdh; Mm.PT.39a.1) for reference. RT-qPCRs were performed in 20-\( \mu \)l volumes using 2x SYBR Green PCR Master Mix (Applied Biosystems, Foster City, CA, USA) in a Roche LightCycler96 system (Roche Applied Science, Indianapolis, IN, USA). The primers for PyMT and the reference gene, Rplp0, have been described previously [12]. Each sample was normalized to the reference gene, sorted MEC cycle threshold change (\( \Delta C_T \)) values were compared with a control population of unsorted MECs, and relative fold induction level for each sorted population/tumor was calculated using the 2^(-\Delta \Delta C_T) method [39].

**Simple Western blot analysis**

Using a PowerGen 125 sawtooth homogenizer (Fisher Scientific), protein lysates were prepared from flash-frozen tumors using Pierce IP Lysis Buffer (25 mM Tris-HCl, pH 7.4, 150 mM NaCl, 1 % Nonidet P-40, 1 mM EDTA, 5 % glycerol; Pierce Biotechnology/Thermo Scientific,
Results

FACS-enriched mammary cell populations expressing PyMT develop tumors with equivalent latency

The degree of tumor heterogeneity in PyMT-based mouse models is associated largely with stage of progression [24], mouse strain [40–42], and the approach used to target the oncogene to mammary epithelium [12, 27, 30, 31]. To extend upon these studies, we asked whether PyMT oncogenesis within specific luminal, basal, and stem cell populations would affect tumor latency, pathology, metastasis, and molecular subgroup. The experimental design for this analysis was to transduce primary MECs overnight with the EF1α-PyMT-ZsGreen lentivirus [12], followed by FACS enrichment of specific cell populations and transplantation with an excess of uninfected and unsorted MECs (Fig. 1a). This technique was designed to ensure that primary cells would be exposed to similar conditions during ex vivo procedures and to provide comparable cellular environments for tumor development.

In order to determine whether expression of the lentiviral transgenes would affect the cell sorting procedure, we assessed ZsGreen fluorescence and PyMT protein levels in MECs for 5 days following transduction with the EF1α-PyMT-ZsGreen lentivirus (Additional file 1, panel a). Only 2 % of cells were ZsGreen-positive at 24 h after transduction, whereas 30 % were positive at day 5. In addition, PyMT protein expression was nearly undetectable until day 3 (Additional file 1, panels b and c). These data suggest that the lentiviral genes would not affect cell sorting within 24 h after transduction.

Freshly isolated primary MECs were infected with the EF1α-PyMT-ZsGreen lentivirus overnight in suspension [12] and then stained and sorted for the cell surface proteins CD24, CD49f, and CD133, which are markers known to delineate luminal, basal, and stem cell populations [34, 35]. Infected cells were not gated on ZsGreen, owing to the low level of expression at 24 h posttransduction. In accordance with published studies, luminal, basal, and stem cells were isolated based on their expression of CD24, CD49f, and CD133, which are markers known to delineate luminal, basal, and stem cell populations [34, 35]. The luminal cell population was further separated into hormone receptor–positive and hormone receptor–negative fractions by expression of CD133 (hereinafter called luminal CD133+ and luminal CD133−, respectively) (Fig. 1b) [34]. We next verified that enriched populations contained the expected keratin markers by evaluating each population for the expression of luminal cell-specific K8 and basal cell-specific K14 by immunofluorescence (Fig. 1c) [43]. As expected, the representative population markers were enriched in the appropriate groups (Fig. 1d) [32–35]. Interestingly, approximately 15 % of luminal CD133+ cells and 33 % of enriched stem cells were positive for both K8 and K14.

Statistical analysis

All data analyses were performed in using GraphPad Prism 6.0d software (GraphPad Software, La Jolla, CA, USA). For each analysis, specific statistical tests are indicated in the figure legends.
Fig. 1 (See legend on next page.)
within the luminal mRNA was variable in tumor samples (Additional file 2, panel b). Moreover, the short tumor latency suggests that transformation is rapid and may be independent of acquired mutations, which is also consistent with the observation that three-dimensionally cultured MECs infected with a PyMT-expressing lentivirus exhibit a transformed phenotype within 2 weeks [12].

Expression of PyMT mRNA and AKT, SRC, and ERK proteins was analyzed in 31 tumors to assess the variability of the PyMT signaling pathway between tumors. Similar to previous findings [12], the expression of PyMT mRNA was variable in tumor samples (Additional file 3, panel a). However, the average expression of PyMT in tumors was not significantly different between tumors arising from the four cellular origins (Fig. 1g) [12]. Furthermore, protein analysis demonstrated that AKT, ERK, and SRC activation was not significantly different between tumors arising from the four cellular origins (Fig. 1h and Additional file 3, panels b–d). These data suggest that PyMT expression and signaling are similar in tumors arising from different MEC populations.

Enriched mammary epithelial cell populations establish tumors with broad histopathology

We next analyzed the histology of tumors originating from each of the enriched MEC populations. We classified tumors by H&E staining and cytokeratin expression and identified the following histologies: acinar, papillary, solid adenocarcinoma, squamous, lipid-rich, and sebaceous-like (Fig. 2a–f, Table 1). The frequency of a specific histology was quantified by estimating its area in two or three different sections per tumor (Fig. 2g–k). Data are presented as the total percentage area of a histological feature in each tumor (Fig. 2g–k).
and as the predominant tumor histology of each tumor (Additional file 4). Tumors that produced lung metastases and those used for microarray analysis are also indicated.

Several trends became apparent. First, histologies were not associated with ERK, AKT, or SRC phosphorylation levels, suggesting that histopathology is uncoupled from

Fig. 2 Analysis and prevalence of histology in tumors derived from mammary epithelial cell (MEC) populations. a through f Representative images of hematoxylin and eosin and cytokeratin staining of tumor histologies: acinar (a), papillary (b), solid adenocarcinoma (c), squamous (d), lipid rich (e), and sebaceous-like (f). Immunofluorescence staining was performed for basal keratin 14 (K14; red) and luminal keratin 8 (K8; green) (scale bar = 100 μm). ZsGreen fluorescence was not detected in the processed sections. Histological area per tumor was derived from luminal CD133+ cells (g), luminal CD133− cells (h), basal cells (i), and stem cells (j). Boxes above each column indicate tumors that were used for microarray analysis. Red boxes = basal subgroup; green boxes = luminal subgroup. Black circles mark tumors that were metastatic. k Average area of histology per MEC group (unpaired t test; n = number of tumors). l Representative images of estrogen receptor (ESR1) staining, including negative (left panel) and positive staining (right panel) (scale bar = 50 μm; n = number of tumors). m Quantification of ESR1 staining per MEC group (two proportion z test). n Quantification of ESR1 staining per histological specimen (two proportion z test). *p < 0.05, **p < 0.01, ***p < 0.0005, ****p < 0.0001
and no difference in tumor latency was observed when transduced MECs were transplanted into ovariectomized mice (Additional file 2). Similar to these results, both the expression of ESR1 in tumor cells and lack of estrogen-dependent growth were observed when PyMT was targeted to mammary glands through intraductal injection of a modified avian retroviral vector [31]. Thus, ESR1 expression and estrogen-dependent growth appear uncoupled in virus-based PyMT tumor models.

### Luminal CD133− mammary epithelial cells give rise to an exclusively basal subgroup mammary tumors

We next tested whether MEC populations influenced the molecular classification of the tumors they generate. Tumors derived from each MEC population were classified by hierarchical analysis of microarray gene expression data. This classification method has previously been used to cluster tumors from a variety of mouse models into basal and luminal subgroups [7]. We analyzed between 6 and 11 tumors generated from each enriched MEC population. An intrinsic 669-gene set consisting of genes differentially expressed in mouse tumors representing basal and luminal subgroups was used to determine the molecular classification [7], and gene set expression data were hierarchically clustered with 12 mouse models of breast cancer. On the basis of this analysis, we found that luminal CD133−, basal cell, and stem cell enriched MECs were able to give rise to tumors of both basal and luminal subgroups (Fig. 3), with no correlation to tumor histology or ESR1 status (data not shown). However, protein analysis of AKT, SRC, and ERK did reveal a difference in AKT phosphorylation in the basal subgroup (Additional file 5, panel b). Interestingly, the luminal CD133− population, which is enriched in luminal progenitors [34], exclusively generated tumors with a broad but distinct spectrum of histological subtypes. ESR1 is a standard clinical marker used to guide a patient's course of treatment and predict clinical outcome [50]. ESR1 IHC staining was used to determine if any enriched MEC subgroups developed tumors with extensive ESR1-positive (ER+) cells, as defined by strong nuclear staining (Fig. 2l) in 20 % or more of tumor cells. Surprisingly, all of the enriched MEC populations gave rise to ER+ tumors, and luminal CD133+ cells generated the highest proportion of ER+ tumors (Fig. 2m). Most ER+ tumors exhibited either papillary or acinar histology, whereas few squamous tumors stained positive for ESR1 (Fig. 2n). These observations are consistent with previous reports that ER+ status appears to be associated with well-differentiated tumor histologies in PyMT oncogene-driven tumors and is less prevalent in tumors that are poorly differentiated [24]. Although ER+ tumors were observed, their hormone dependence was not apparent. Cells isolated from both ER+ and ER− tumors exhibited similar in vitro sensitivity to tamoxifen, and no differences in tumor latency were observed when transduced MECs were transplanted into ovariectomized mice (Additional file 2). Similar to these results, both the expression of ESR1 in tumor cells and lack of estrogen-dependent growth were observed when PyMT was targeted to mammary glands through intraductal injection of a modified avian retroviral vector [31]. Thus, ESR1 expression and estrogen-dependent growth appear uncoupled in virus-based PyMT tumor models.
established tumors within the basal subgroup. To determine the significance of this observation, we compared the distribution of basal and luminal subgroups generated from each cell population with the distribution we observed when unsorted MECs were transduced with the EF1α-PyMT-ZsGreen lentivirus [8]. Compared with unsorted MECs, the basal-restricted distribution of tumors derived from luminal CD133− cells was statistically significant (p = 0.017) (Additional file 6: Table S1). In contrast, all other cell populations generated tumors of both basal and luminal subgroups at a frequency similar to that of unsorted MECs. These data demonstrate that the stem, basal, and luminal CD133+ MEC populations can establish tumors that classify within either molecular subgroup, whereas CD133− luminal cells preferentially establish tumors of the basal subgroup.

Unsupervised hierarchical clustering of tumors generated from enriched MEC populations revealed two major hierarchical tumor clusters, one composed of only CD133− luminal tumors and another composed of tumors generated from all MEC populations (Additional file 7). Differential expression analysis comparing these two major hierarchical clusters identified 1046 unique genes showing at least twofold differential expression at an adjusted p value <0.05 (Additional file 8: Table S2 and Additional file 9: Table S3). One gene of interest, Sox11, was upregulated in tumors derived from luminal CD133− MECs. Interestingly, Sox11 is a transcriptional regulator upregulated in BRCA1-mutant breast cancers and has been shown to promote cell survival and proliferation in breast cancer cells [51].

Luminal CD133+ cells give rise to tumors that are significantly less metastatic than other tumor groups

We investigated whether the enriched MEC populations expressing PyMT could generate tumors with different metastatic capacities. Metastasis was measured in mice with similar tumor latency, tumor size (2 cm), and tumor burden (single tumor). Similar to other PyMT-driven models, primary tumors generated from each MEC population metastasized to the lungs (Fig. 4a, b) [16]. Metastases to other organs were not detected. The metastatic burden in lungs was quantified by whole mount fluorescent imaging and H&E staining of serial lung sections from mice with similar tumor size and latency. Interestingly, although no significant differences in tumor prevalence or progression were observed between the four tumor-initiating populations (Fig. 1d), tumors derived from the luminal CD133+ cell population were significantly less metastatic than all other tumor groups (Fig. 4c). Furthermore, these cells formed significantly fewer lung tumor foci than the other tumor groups (Fig. 4d); however, there was no difference in the average area per metastatic focus (Fig. 4e).

To determine whether tumors with specific histologies have different metastatic propensities, we examined the numbers of metastases per tumor histology type (Additional file 10, panels a–d). We did not find an association between metastatic propensities and histology (Additional file 11). In addition, an association was not observed between PyMT downstream activity and number of metastases (Additional file 5, panel c). Taken together, these data suggest that metastatic properties of a tumor may be influenced by the tumor’s cellular precursor.

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**Fig. 3** Tumor microarray gene expression profiling. Tumors were analyzed by microarray gene expression profiling and hierarchically clustered with mouse mammary tumor models using an intrinsic gene set identified by Herschkowitz et al. [7]. Vertical lines indicate individual tumors. Each enriched mammary epithelial cell population is indicated by a different color: green = luminal CD133+; blue = luminal CD133−; red = basal; black = stem cells. Mouse mammary tumor models that generate molecularly luminal tumors are shown in dark blue, and those predominantly within the basal subgroup are displayed in dark red. Normal mouse mammary tissue is represented in brown (n = number of tumors). DMBA 7,12-dimethylbenz[a]anthracene, MMTV mouse mammary tumor virus, PyMT polyomavirus middle T antigen.
Metastasis progresses through several stages: Cells leave the primary site and enter the bloodstream or lymphatics, survive within the circulatory system, exit the circulatory system, and colonize a secondary site [52].

We next tested whether tumors derived from CD133+ cells were deficient in one or more steps of the metastatic process. First, we quantified CTCs in mice bearing primary tumors, which provides a measure of tumor cell invasion and intravasation into the bloodstream. CTCs expressing ZsGreen, a fluorescent protein coexpressed with PyMT, were quantified in whole blood isolated from tumor-bearing mice by FACS and normalized to both the volume of collected blood and background fluorescence observed in non-tumor-bearing mice. Whole blood from...
mice with luminal CD133+ cell transplants had significantly fewer CTCs than the other tumor groups (Fig. 4f). Next, using a non-spontaneous metastasis assay, metastatic colony formation was assessed following tail vein injection of isolated tumor cells to evaluate their ability to extravasate from the bloodstream and colonize secondary sites. When injected into the tail vein, tumor cells derived from both luminal CD133+ and luminal CD133− populations were able to colonize and proliferate within the lung, regardless of whether the donor tumor was metastatic (Fig. 4g and Additional file 12). These data suggest that luminal CD133+ cells give rise to tumors with a limited ability to escape the primary cancer and intravasate into the bloodstream.

Discussion

The PyMT oncogene has a broad capability to establish a variety of tumor histologies and subtypes. Members of the Li laboratory developed a model wherein a modified avian leukosis sarcoma virus expressing PyMT (RCAS-PyMT) was used to infect transgenic mice engineered to express tva, the RCAS (replication competent avian sarcoma leukosis virus LTR splice acceptor) receptor, on different mammary cell types [53]. Intraductal infection with RCAS-PyMT was performed in mice expressing tva under the control of several promoters, including the MMTV LTR and K6 promoter [27, 31]. When tumors generated by the different models were compared, dissimilar histological and molecular phenotypes were observed. MMTV-tva/RCAS-PyMT tumors were acinar and composed of luminal and myoepithelial cells, a phenotype that contrasted with the papillary tumors generated by K6-tva/RCAS-PyMT. These researchers also showed that the molecular profiles of tumors generated by MMTV-tva/RCAS-PyMT, K6-tva/RCAS-PyMT, and MMTV-PyMT were not similar, with the K6-tva tumors having a unique profile similar to that of a normal-like breast cancer subtype [27]. In an expanded analysis of molecular profiles from the RCAS-PyMT and MMTV-PyMT models, Hollern and Andrechek further demonstrated significant diversity, particularly among tumors derived from different mouse strains [40]. Taken together, these data demonstrate that the PyMT oncogene has the capacity to generate several tumor subtypes and that differences in the cellular context of the oncogene can affect tumor heterogeneity.

We have similarly observed a variety of histological and molecular subtypes in tumors induced by the PyMT oncogene. Previously, we showed that broad expression of the oncogene in mammary epithelium, using the EF1α-PyMT-ZsGreen lentivirus, generated late-stage tumors consisting of both luminal and myoepithelial cells, with a majority of tumors having acinar and solid histology [12], which are similar features of tumors established by the MMTV-tva/RCAS-PyMT model [53]. However, a notable difference between these models was the appearance of squamous metaplasia, which was observed in 30% of the EF1α-PyMT-ZsGreen tumors, but not in MMTV-tva/RCAS-PyMT tumors [12, 27]. This tumor subtype is rare in the MMTV-PyMT model, with a reported frequency of only 4–8% [22, 40]. The data in our present study expand upon those previous studies by demonstrating that squamous metaplasia was observed more frequently when basal and stem cell populations were infected with the EF1α-PyMT-ZsGreen lentivirus. Similar data were obtained by Keller et al. from studies performed with normal human breast cells. They showed that transduction of CD10-enriched human basal cells with oncocgenic lentiviruses resulted in ER− and metastatic tumors with squamous differentiation. In contrast, epithelial cell adhesion molecule–positive (Epcam+) luminal cells generated ductal carcinomas with luminal features [54]. In addition, Ince et al. showed that transformation of myoepithelial-like human breast cells generated squamous carcinomas of the breast, whereas luminal-like cells gave rise to adenocarcinomas [14]. Collectively, these studies provide strong support that a cell within the basal lineage, either a myoepithelial cell or a stem cell, is the origin of squamous metaplasia in mammary tumors.

The ability to fractionate primary mouse mammary cells into luminal, basal, and stem cell populations, and their receptiveness to ex vivo manipulation and outgrowth following transplantation, provides a unique method to study how cellular context affects oncogenesis [12]. In the present study, we investigated whether tumor latency, histology, metastasis, and molecular subtype are altered when the PyMT oncogene is targeted to distinct FACS-enriched mammary cell populations. The data demonstrate that each MEC population is able to generate tumors at similar latency and with broad pathology. However, some cell populations preferentially established distinct pathologies. Most striking was that luminal CD133+ cells gave rise to a higher proportion of tumors with papillary histology and ESR1 expression and the lowest proportion of tumors with squamous metaplasia. Consistent with their well-differentiated pathology, tumors from luminal CD133+ cells also produced fewer CTCs and metastases. An opposing phenotype was observed in tumors generated by enriched stem cells. Notably, these cells generated tumors that were ER−, exhibited squamous metaplasia, and produced more CTCs and metastases than luminal CD133+ cells. Thus, luminal CD133+ cells had a propensity to produce well-differentiated tumors, whereas poorly differentiated and metaplastic tumors were more commonly observed from enriched stem cells.
Although the MMTV LTR is expressed primarily in luminal cells, it is also active broadly in basal, stem, and luminal cell populations of the mammary gland [27, 29]. In the MMTV-PyMT model, the oncogene may target a multipotent progenitor because clonal cell lines derived from precancersous lesions can establish cell types expressing markers for both luminal and myoepithelial populations [25, 55]. Histological and cellular heterogeneity are also observed in vivo in the MMTV-PyMT model, particularly during precancersous development. Early lesions are composed of both hormone receptor-positive and hormone receptor-negative cells, with sporadic myoepithelial coverage [24]. As the tumors develop to malignant lesions, they exhibit a loss of both hormone receptor-expressing cells and myoepithelial cells [24] and an expansion of cells double-positive for CD61 and CD29 [26], markers that are coexpressed on normal luminal progenitors. Some similarities between MMTV-PyMT tumors and those derived from the luminal CD133− tumors are evident. In particular, the histopathology of tumors from this population was most similar to that of tumors in the MMTV-PyMT mouse model. Solid adenocarcinomas are seen in 60 % of MMTV-PyMT tumors [22], and this was the dominant histology in approximately 40 % of tumors derived from the luminal CD133− cell population. However, tumors established by luminal CD133− cells were classified as a basal subgroup, whereas MMTV-PyMT tumors clustered within the luminal subgroup [7]. The reason for this discrepancy is unclear, but it may be a result of differences in the developmental stage of cells targeted in each model. The MMTV LTR is active at prepubertal stages of mammary gland development [29], which suggests that immature mammary epithelium is the origin for MMTV-PyMT tumors. In contrast, the primary MECs used for FACS enrichment in our study were derived from postpubertal mice. These cells would have had been exposed to the maturation effects of systemic hormones, which may alter their lineage potential. Recent studies have shown that oncogenic stress can significantly increase the plasticity and multilineage potential of differentiated MECs [56–58]. Both the MMTV-tva/RCAS-PyMT and EF1α-PyMT-ZsGreen models generate heterogeneous tumors consisting of myoepithelial and luminal cell lineages, which contrasts with the predominantly luminal cell type that is observed in late-stage MMTV-PyMT tumors [59]. These data suggest that tumor cells generated by the MMTV-PyMT and virus-based models have differences in their lineage plasticity, particularly at more advanced stages of progression, which may influence the molecular classification of tumors. Further studies are necessary to assess the effect of oncogenic stress on the lineage plasticity of MECs at pre- and postpubertal stages of development.

Using the method described by Herschkowitz et al., we categorized tumors derived from the sorted cell populations and 12 different mouse models of breast cancer into 2 general molecular subgroups: basal and luminal [7, 60]. These data demonstrate that both luminal and basal tumor subgroups can arise from enriched luminal CD133+, stem, and basal cell populations. However, we also show that transformation of CD133− luminal cells, which are enriched for luminal progenitors [34], generated tumors of the basal subgroup. This finding suggests luminal progenitors preferentially establish basal rather than luminal tumors. Consistent with this, several recent observations attribute basal-like breast cancer to a luminal population. Lim et al. demonstrated that the molecular profile of untransformed luminal progenitors most closely resembles basal-like breast cancers [8]. In addition, transformation of human EpCAM+/CD10−/CD49f− luminal progenitors derived from reduction mammoplasties established tumors with features similar to basal-like breast cancer, including reduced ESR1 and greater K14 expression than tumors derived from differentiated luminal cells [54]. Furthermore, targeting Brca1 loss of function to luminal cells in mice generated tumors with basal-like features that closely resembled those observed in patients carrying the BRCA1 mutation. However, the same loss of function in basal cells generated adenomyoepitheliomas [10, 15]. Taken together, these data support luminal progenitors as a cellular origin of basal-like breast cancer.

There are a number of challenges to studying how the cell of origin influences tumor progression in breast cancer. These include the complexity of normal mammary cell types from which a tumor may arise [60] and technical difficulties in precisely targeting oncogenes to specific cell types. As such, the data from this study should be considered in the context of limitations of the approach used to target expression of PyMT to distinct cell populations. For example, the cells that were transplanted represent enriched populations but not purified cell types. Thus, a tumor phenotype cannot be credited to a specific cell type. In addition, specific cell populations may have a transduction bias, as has been observed between cultured luminal and myoepithelial cells, which may limit the diversity of cell types targeted by the lentivirus [61]. Transduction of primary MECs with lentiviruses and the subsequent expression of PyMT and ZsGreen may also alter the sorting process. However, maximum expression of lentiviral transgenes generally takes several days. ZsGreen expression was observed in only about 2 % of cells 24 h after transduction, and PyMT protein expression was low at this time. Because cell sorting and transplantation were performed within 24 h after the cells were exposed to virus, it is unlikely
that expression of PyMT would have any significant influence on the populations before the sort. In addition, the study was designed to ensure each sorted population was exposed to similar conditions during the ex vivo procedures. Primary cells were collected, transduced, and sorted by FACS from a pool, and each population was cotransplanted with 5–20-fold more uninfected, unsorted MECs, which would provide comparable in vivo environments for tumor development. Thus, the main variable was the cell population used to generate the tumors.

**Conclusions**

We report that differentiated luminal (CD133+), luminal progenitor (CD133−), basal, and stem cell enriched populations have the capacity to give rise to mammary tumors at equivalent frequency and latency when transformed with PyMT oncogene. However, mammary cell populations can produce tumors with differences in histopathology, molecular classification, and metastatic potential. Basal and stem cell enriched populations predominantly established poorly differentiated squamous tumors, whereas differentiated luminal cells gave rise to a high proportion of ER+ papillary tumors. We also demonstrate that the luminal progenitor enriched population specifically gives rise to tumors that cluster within the basal molecular subgroup. Taken together, these data demonstrate that targeted expression of PyMT in different cell types affects tumor histology, molecular subtype, and metastatic potential. In addition, the data provide support that the cellular origin of squamous metaplasia resides within the basal or mammary stem cell population.

**Additional files**

**Additional file 1: Figure S1.** Postinfection PyMT expression in cultured total MECs. a Flow cytometric analysis of percentage of ZsGreen-positive cells for 5 days post-ET1a-PyMT-ZsGreen infection (10,000 events, mean ± SD, n = 3). b Quantification from simple Western blot analysis of PyMT-HA expression over time in infected MECs and uninfected MEC controls for 5 days following infection. PyMT-HA peak area was normalized to GAPDH peak area. 293 T cells are shown as positive and negative controls for PyMT-HA expression. c Multiplex simple Western blot stained for hemagglutinin (HA) tag and GAPDH. Protein lysates were prepared each day for 5 days following infection and PyMT-HA (60 kDa) with 293 T positive and negative controls. HA tag and GAPDH signals are shown with different exposure times. (TIFF 1824 kb)

**Additional file 2: Figure S2.** Hormone independence of ET1a-PyMT-ZsGreen-transduced MEC tumors. a Half-maximal inhibitory concentration values of wild-type MECs and cells isolated from luminal CD133+, luminal CD133−, basal, and stem cell tumors treated for 48 h with 4-hydroxytamoxifen (4-OHT). No significant differences among the viability of tumor cells and normal MECs were observed after 4-OHT treatment (t test). b Assessment of tumor growth response to ovariectomy. Transduced primary cells were transplanted into ovariectomized mice to test if lack of hormone signaling would result in reduced tumor growth from luminal CD133+ cells. No differences in tumor latencies were observed among tumor groups (n = number of mice). (TIFF 1112 kb)

**Additional file 3: Figure S3.** PyMT expression and activity in tumors. a RT-qPCR quantification of relative mRNA expression levels in individual MMTV-PyMT, luminal CD133+ cell, luminal CD133− cell, basal cell, and stem cell tumors normalized to the β-actin housekeeping gene. Analysis was performed in triplicate for each tumor sample. b Western blot quantification of relative ratio of pAKT to AKT protein expression in individual MMTV-PyMT, luminal CD133+ cell, luminal CD133− cell, basal cell, and stem cell tumors normalized to β-actin. c Western blot quantification of relative ratio of pERK to ERK protein expression in individual MMTV-PyMT, luminal CD133+ cell, luminal CD133− cell, basal cell, and stem cell tumors normalized to β-actin. d Western blot quantification of relative ratio of pSRC to SRC protein expression in individual MMTV-PyMT, luminal CD133+ cell, luminal CD133− cell, basal cell, and stem cell tumors normalized to β-actin. (TIFF 2407 kb)

**Additional file 4: Figure S4.** Distribution of basal and luminal tumor subtypes generated by EF1α-PyMT-ZsGreen lentiviral transduction of each sorted cell population as representing at least 50% of the tumor area. Tumors classified as mixed did not have a dominant histological type. Tumors derived from stem cells were significantly more squamous than tumors derived from luminal CD133+ and CD133− cells (two proportion z test, n = number of tumors). (TIFF 833 kb)

**Additional file 5: Figure S5.** PyMT activity comparison with tumor histology, molecular subtype, and metastasis. a Western blot quantification of relative ratio of pAKT to AKT, pERK to ERK, and pSRC to SRC protein expression in enriched population tumors segregated based on tumor histology and normalized to β-actin. No significant differences in protein expression were detected among tumor groups (t test, n = number of tumors). b Western blot quantification of relative ratio of pAKT to AKT, pERK to ERK, and pSRC to SRC protein expression in enriched population tumors segregated based on tumor molecular subtype and normalized to β-actin. Molecularly basal tumors exhibited significantly higher levels of pERK/ERK levels than luminal tumors (t test, p value shown, n = number of tumors). c Western blot quantification of relative ratio of pAKT to AKT, pERK to ERK, and pSRC to SRC protein expression in enriched population tumors segregated based on tumor metastasis and normalized to β-actin. No significant differences in protein expression were detected among tumor groups (t test, n = number of tumors). (TIFF 1046 kb)

**Additional file 6: Table S1.** Distribution of basal and luminal subtypes. Distribution of basal and luminal tumor subtypes generated by ET1a-PyMT-ZsGreen lentiviral transduction of each sorted cell population as compared with the distribution obtained from ET1a-PyMT-ZsGreen-transduced but unsorted MECs [8], using Fisher’s exact test. (DOCX 14 kb)

**Additional file 7: Figure S6.** Unsupervised hierarchical clustering of tumors generated from enriched MEC populations. Tumor cell of origin, histology, metastatic propensity, and molecular subtype are indicated below the dendrogram cluster. (TIFF 1486 kb)

**Additional file 8: Table S2.** 100 unique differentially expressed genes in the luminal CD133+–tumor cluster compared with the mixed tumor cluster. Group 1 is the mixed tumor cluster, and group 2 is the luminal CD133+–tumor cluster. A t test was performed on Benjamini–Hochberg-corrected, log-transformed data. Table includes gene name, ratio, p value, and direction. (XLSX 14 kb)

**Additional file 9: Table S3.** All differentially expressed genes in the luminal CD133+–tumor cluster compared with the mixed tumor cluster. Group 1 is the mixed tumor cluster, group 2 is the luminal CD133+–tumor cluster. A t test was performed on Benjamini–Hochberg-corrected, log-transformed data. Table includes gene name, ratio, p value, and direction. (XLSX 73 kb)

**Additional file 10: Figure S8.** Number of metastases per tumor histology. Histological area per tumor derived from luminal CD133+ cells (a), luminal CD133− cells (b), basal cells (c), and stem cells (d), as well as the number of metastatic foci detected in the lungs of the mice bearing each of the tumors (shown as red values above each tumor). (TIFF 1562 kb)

**Additional file 11: Figure S9.** Quantification of the number of metastases per tumor histology. A histology was assigned to a tumor when the histology comprised at least 50% of the tumor area. Tumors
that mixed histologies were classified as mixed. No statistically
significant differences were noted in the metastatic properties among histological
tumor types were observed (Mann–Whitney U test). (TIFF 654 kb)

**Additional file 12: Figure S7.** Enriched MEC populations metastasize
to lung. Representative fluorescence images of lungs from mice that received
tail vein injections of metastatic luminal CD133+ tumor cells (tumor 1), nonmetastatic luminal CD133+ tumor cells (tumor 2), or
metastatic luminal CD133− tumor cells (tumors 3 and 4). Lungs were examined 3 weeks after the tail vein injection. Fluorescent black-and-white
lung images were false-colored green in ImageJ software and overlaid on a
black background. (TIFF 2284 kb)

**Abbreviations**

7-AAD: 7-Amino-actinomycine D; APC: Allophycocyanine; BSA: Bovine serum albumin; CTC: Circulating tumor cell; DMBA: 7,12-Dimethylbenz[a]anthracene;
EDTA: Ethylenediaminetetraacetic acid; EF1α: Elongation factor 1 alpha; EpCAM: Epithelial cell adhesion molecule; ER and Er1: Estrogen receptor; Er−: Estrogen receptor−negative; Er+ and Er1: Estrogen receptor positive; ERK/ 1,2: Extracellular signal-regulated kinase 1/2; FACS: Fluorescence-activated
cell sorting; GAPDH: Glyceraldehyde 3-phosphate dehydrogenase; GEO: Gene Expression Omnibus; HA: Hemagglutinin; H&E: Hematoxylin and eosin; HBSS: Hanks’ balanced salt solution; HCl: Huntsman Cancer Institute;
IgG: Immunoglobulin G; IHC: Immunohistochemistry; KB: Keratin 8; K14: Keratin 14; LTR: Long terminal repeat; MEC: Mammary epithelial cell;
MMTV: Mouse mammary tumor virus; PBS: Phosphate-buffered saline; PR and Pgr: Progesterone receptor;PyMT: Polyomavirus middle T antigen;
RCAS: Replication competent avian sarcoma leukosis virus long terminal repeat splice acceptor; RT-qPCR: Real-time quantitative PCR.

**Competing interests**

The authors declare that they have no competing interests.

**Authors’ contributions**

DD participated in study design and performed experiments, analyzed data,
and drafted the manuscript. BAS participated in study design, experiments,
data analysis, and edited the manuscript. MM and KPG designed and
performed RT-qPCR and protein analysis experiments and edited the
manuscript. HAE performed tail vein injections and lung metastasis
quantification and contributed to editing of the manuscript. BDW participated in study design and coordination and helped draft and edit the manuscript. All
authors read and approved the final manuscript.

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