Genetic Ancestry-dependent Differences in Breast Cancer-induced Field Defects in the Tumor-adjacent Normal Breast

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Abstract

Purpose: Genetic ancestry influences evolutionary pathways of cancers. However, whether ancestry influences cancer-induced field defects is unknown. The goal of this study was to utilize ancestry-mapped true normal breast tissues as controls to identify cancer-induced field defects in normal tissue adjacent to breast tumors (NATs) in women of African American (AA) and European (EA) ancestry.

Experimental Design: A tissue microarray comprising breast tissues of ancestry-mapped 100 age-matched healthy women from the Komen Tissue Bank (KTB) at Indiana University (Indianapolis, IN) and tumor-NAT pairs from 100 women (300 samples total) was analyzed for the levels of ZEB1, an oncogenic transcription factor that is central to cellular fate, mature luminal cell–enriched estrogen receptor alpha (ERα), GATA3, FOXA1, and for immune cell composition.

Results: ZEB1+ cells, which were localized surrounding the ductal structures of the normal breast, were enriched in the KTB-normal of AA compared with KTB-normal of EA women. In contrast, in EA women, both NATs and tumors compared with KTB-normal contained higher levels of ZEB1+ cells. FOXA1 levels were lower in NATs compared with KTB-normal in AA but not in EA women. We also noted variations in the levels of GATA3, CD8+ T cells, PD1+ immune cells, and PDL1+ cell but not CD68+ macrophages in NATs of AA and EA women. ERα levels did not change in any of our analyses, pointing to the specificity of ancestry-dependent variations.

Conclusions: Genetic ancestry–mapped tissues from healthy individuals are required for proper assessment and development of cancer-induced field defects as early cancer detection markers. This finding is significant in light of recent discoveries of influence of genetic ancestry on both normal biology and tumor evolution.

Introduction

Recent data demonstrating a correlation between lymph node positivity at the time of detection, and the probability of disease recurrence even decades postdetection, only solidifies the principle that detection of breast cancer prior to lymph node metastasis can appreciably improve clinical outcomes (1). Although the last decade witnessed significant improvements in imaging technologies including 3D-mammography, false negatives remain a significant concern (2). One way to overcome these false negatives is to complement radiologic techniques with molecular assays that measure "transcriptomic and epigenetic field effect" of tumors on adjacent "normal" (NAT) tissues. Tschendoff and colleagues demonstrated tumor-induced epigenetic field defects in NATs specifically targeting transcription factor–binding sites specifying chromatin architecture and stem cell differentiation pathways (3). These include Wnt and FGF signaling networks. Unfortunately, the Tumor Genome Atlas (TCGA) of breast cancer utilized reduction mammaplasty or NATs as their controls in transcriptome analyses (4). These are often substituted for "normal" controls in comparative analyses with breast cancers. This limitation was highlighted in another study, which compared TCGA "normal" breast transcriptome with the transcriptome of epithelial cells from the breast of healthy women. Significant differences were noted between these 2 sources of normal tissues (5). Reduction mammaplasty samples are also histologically abnormal compared with breast tissues from healthy women (6).

While molecular markers of cancers, particularly gene expression signatures, are traditionally developed by comparing gene expression between available "normal" and cancer tissues, the possibility of genetic ancestry of samples having an impact on gene expression under normal and abnormal conditions is rarely taken into consideration. The effects of genetic ancestry on tumor evolution and gene expression are just beginning to be
Translational Relevance

Breast cancer diagnosis prior to lymph node metastasis can appreciably improve clinical outcomes. While radiologic techniques have improved early diagnosis, molecular markers that can complement radiologic techniques are needed to improve specificity. This study aimed to investigate how both genetic ancestry and appropriate control tissues influence detection of cancer-induced changes in the breast. We show that alterations in ZEB1+ cells in tissues surrounding tumors are observed predominantly in women of European (EA) ancestry, whereas FOXA1+ cells were altered in normal tissues adjacent to tumors of women of African American (AA) ancestry. Immune cell activation in tumors, as well as surrounding tissue showed genetic ancestry-dependent variations as evident from differences in PD1+ and PDL1+ cells in the normal tissue adjacent to tumors of women of AA and EA ancestry. Thus, biomarker discovery needs to consider not only sample size and statistical methods but also genetic ancestry and true normal control tissues.

recognized (7). This observation is highly relevant in the context of known differences in cancer incidence and/or outcome based on genetic ancestry. For example, women of African American ancestry (AA) suffer higher mortality from the aggressive breast cancer subtype, triple-negative breast cancer (TNBC), than women of European ancestry (EA; ref. 8). In contrast, breast cancer in Hispanic and Native American women is less prevalent and these women have better outcomes (9, 10). Whether the worse outcome in AA women is due to an increased incidence of TNBC or unique biological factors that promote aggressive biology is an important but unresolved challenge in cancer disparities research. Dietze and colleagues (8) recently highlighted that key molecular pathways, including Aurora A-PLK, EZH2, and Wnt-stem cell and Nat colleagues (8) recently highlighted that key molecular pathways, including Aurora A-PLK, EZH2, and Wnt-stem cell signaling networks, are significant upregulated in TNBCs of AA women compared with TNBCs of EA women. The review (8) further emphasized that it remains unknown whether genomic aberrations unique to TNBCs in AA women result in activation of these signaling pathways in tumors or whether the basal activity of these pathways in normal AA women’s breasts is inherently different compared with EA women’s breasts. It remains possible that normal breast biology varies based on genetic ancestry. Evidence for this possibility comes from a recent discovery of breast cancer protective alleles in Latinas (11). SNPs in the protective allele are located on gene regulatory regions affecting the expression of genes linked to differentiation. Our own studies have discovered enrichment of a unique population of cells in the normal breast of AA women (12). Furthermore, a breast cancer susceptibility locus in AA women, potentially altering the expression levels of miRNA mir-3065, has recently been described (13).

Here, we took advantage of genetic ancestry–mapped true normal breast tissues to identify differences between true normal and NATs. These differences can potentially be developed into the earliest markers of breast cancer initiation. A tissue microarray (TMA) comprising breast tissues from clinically normal breast, NATs, and tumors were analyzed for markers that are expressed in cells with stem or mature luminal cell properties. We also examined the TMA for CD8+ T cells, CD68+ macrophages, PD1+ immune cells, and PDL1+ epithelial cells to determine whether immune cell composition of tumors and NATs in AA women differ from those of EA women.

Materials and Methods

Generation of TMA
Breast core biopsies from healthy women donated to the Komen Tissue Bank (KTB) at Indiana University (Indianapolis, IN) and surgical material left over after pathologic assessment as a part of a treatment protocol were obtained after informed written consent from the subjects. All experiments were carried out in accordance with the approved guidelines of the Indiana University Institutional Review Board (Indianapolis, IN). International Ethical Guidelines for Biomedical Research Involving Human Subjects were followed. We created a TMA comprising healthy breast tissue from the KTB (KTB-normal), matched normal adjacent to tumor (NAT), and tumor tissue of approximately 50 each of AA and Caucasian women (total ~300 samples). KTB-normal tissues were age- and race-matched to NATs/tumors. Body mass index (BMI) of AA women who donated tissues to KTB was 32.3 ± 9, whereas it was 28.3 ± 8.5 in case of Caucasian women. Each sample was spotted in duplicate in cases of NATs and tumors.

IHC and statistical analyses
TMAs were analyzed for ZEB1, MSRB3, estrogen receptor alpha (ERα), FOXA1, and GATA3 expression. All IHC was done in a Clinical Laboratory Improvement Amendments (CLIA)-certified histopathology laboratory and evaluated by 3 pathologists in a blinded manner. Quantitative measurements were done using the automated Aperio Imaging system and analysis was done using an FDA-approved algorithm. Positivity and H-scores were scored and statistically analyzed as described previously (14, 15). With respect to PD-L1, a tumor proportion score (TPS) was conducted alongside a pathology hand score. The TPS describes the ratio of positive viable tumor cells against all viable tumor cells. The PD-L1 TPS followed the prescribed FDA reading: a negative score having <1% positive staining on the cell membrane, a positive score having 1%–49% tumor cells partially or completely expressing PD-L1 on the cell membrane, and a strong positive having ≥50% tumor cells partially or completely expressing PD-L1 on the cell membrane at a stronger intensity (16, 17). Data were analyzed in 3 different ways: (i) expression differences between AA and EA KTB-normal; (ii) expression differences between KTB-normal and NATs; and (iii) expression differences between NATs and tumors. The statistical software SAS version 9.4 was used to complete the statistical analyses with P < 0.05 considered significant. Nonparametric Wilcoxon rank-sum tests were used for unpaired analyses, as positivity and H-scores were not normally distributed, whereas nonparametric Wilcoxon signed-rank tests were used for paired analyses. The following antibodies were used: CD8 (Dako IR623), CD68 KP1 (Dako IR609), ER clone:EP1 (Dako IR 084), FOXA1 (Santa Cruz Biotechnology sc-6553), GATA3 (Santa Cruz Biotechnology sc-268), MSRB3 (HPA014432, rabbit polyclonal, Sigma), PD1 (Cell Marque Corporation 315M-98), PDL1 (Keytruda; clone 22c3, Dako IHC 22c3), and ZEB1 (3G6, catalog No. 14-9741-82, bBioscience).
Results

**ZEB1**

Cells are enriched in the normal breasts of AA women compared with EA women.

In the mouse mammary gland, PROCR<sup>+</sup>/EpCAM<sup>+</sup> cells are purported to function as multipotent stem cells (18). In our previous study focused on evaluating ethnicity-dependent differences in the normal breast, we observed specific enrichment of PROCR<sup>+</sup>/EpCAM<sup>+</sup> cells in cultured normal breast epithelial cells from biopsies of healthy AA women compared with EA women (12). These cells are enriched for the expression of stemness-related transcription factor ZEB1 and have enhanced Wnt pathway activity compared with PROCR<sup>+</sup>/EpCAM<sup>+</sup> cells (12). ZEB1 has recently been demonstrated to limit oncogenic p53-driven DNA damage response in stem cells and thus increase the stem cells' intrinsic susceptibility to malignant transformation (19). ZEB1 cells coexpress the methionine sulfoxidase reducer (MSRB3), which protects against DNA damage (19). These observations raised the possibility that PROCR<sup>+</sup>/ZEB1<sup>+</sup> cells are naturally present at higher levels in the normal breasts of AA women and that failure to consider natural variation in gene expression pattern, influenced at least partially by genetic ancestry, could have an impact on identifying cancer-induced field effect on the adjacent normal breast. Measuring PROCR itself in the breast tissue is complicated because there are 4 haplotypes of PROCR due to SNPs and only 1 among them is a cell surface protein (20). Because ZEB1 expression is enriched in PROCR<sup>+</sup>/EpCAM<sup>+</sup> cells, we used ZEB1 as a surrogate marker for PROCR<sup>+</sup>/EpCAM<sup>+</sup> cells in unmanipulated breast tissues.

Normal breast tissue from biopsies of healthy AA women and those from AA and EA women are shown in Fig. 1A and statistical analyses are presented in Table 1. Descriptive statistics of ethnicity, age, menstrual status, pregnancy and breastfeeding history, hormone replacement therapy, and numerical values are presented in Table 1. While ZEB1 levels of KTB-normal tissues of AA women were modestly higher than in EA women, NATs of AA women had lower FOXA1 expression compared with KTB-normal tissues (Fig. 2B). Thus, tumors through their field effect may decrease FOXA1 in the surrounding breast tissues of AA women.

GATA3 levels are higher in KTB-normal of EA compared with AA women

We also examined expression levels of GATA3 to determine whether hormonal signaling networks show genetic ancestry-dependent variation. Consistent with this possibility, GATA3 H-score and positivity were higher in KTB-normal tissues of EA women compared with those of AA women (Table 1; Supplementary Fig. S2B). Furthermore, GATA3 is a likely candidate for cancer-induced field defects in EA women as its levels were significantly lower in NATs of EA but not AA women compared with their KTB-normal counterparts (Supplementary Fig. S2C).

ERα<sup>+</sup> cells remain stable

ERα<sup>+</sup> cells in the normal breast are considered to be highly differentiated nonproliferative cells and control proliferation of ERα<sup>−</sup> cells through paracrine mechanisms (31). Representative ERα staining pattern is shown in Fig. 3A and statistical analyses are presented in Fig. 3B-D and Table 1. Neither KTB-normal tissues nor NATs showed genetic ancestry-dependent differences in ERα levels. The results are not only relevant but also reassure that our TMA detects only specific changes.

FOXA1 expression is lower in NATs of only AA women

FOXA1 serves as a pioneer factor that controls chromatin access of various nuclear receptors including ERα and controls the expression of genes enriched in luminal cells compared with basal cells (22-24). FOXA1 along with another pioneer factor GATA3 and ERα form a lineage-restricted hormone-responsive signaling network in the normal breast (25). While higher expression of FOXA1 in the primary tumor is associated with better outcome, its overexpression in metastatic and/or anti-estrogen-resistant tumors is associated with rewiring of ERα signaling and poor outcome (26-29). In addition, it is suggested that FOXA1 gene is preferentially methylated in tumors of AA women (30). Because of its relative importance in breast cancer, we assessed our TMA for FOXA1 expression. Representative staining pattern of FOXA1 is shown in Fig. 2A and numerical values are presented in Table 1. While FOXA1 levels in KTB-normal tissues of AA women were modestly higher than in EA women, NATs of AA women had lower FOXA1 compared with KTB-normal tissues (Fig. 2B). Thus, tumors through their field effect may decrease FOXA1 in the surrounding breast tissues of AA women.

MSRB3 has recently been shown to be one of the downstream transcriptional targets of ZEB1 and it cooperates with ZEB1 during transformation of stem-like cells (19). To correlate ZEB1 expression with its activity, we measured the levels of MSRB3 using the same antibody used in the above study. We could measure positivity but not H-score because of low-level expression. The expression pattern was similar to that of ZEB1, as cells surrounding the ducts showed expression (Supplementary Fig. S1B). However, KTB-normal tissues of AA and EA women expressed similar levels of MSRB3 (Table 1; Supplementary Fig. S1C), which could be due to regulation by other transcription factors or to the low expression levels, making data interpretation difficult. Furthermore, except for a modest change in expression in NATs compared with KTB-normal tissues, no other differences were noted (Supplementary Fig. S1C and S1D).

In the mouse mammary gland, PROCR<sup>−</sup>/EpCAM<sup>−</sup> cells are purported to function as multipotent stem cells (18). In our previous study focused on evaluating ethnicity-dependent differences in the normal breast, we observed specific enrichment of PROCR<sup>−</sup>/EpCAM<sup>−</sup> cells in cultured normal breast epithelial cells from biopsies of healthy AA women compared with EA women (12). These cells are enriched for the expression of stemness-related transcription factor ZEB1 and have enhanced Wnt pathway activity compared with PROCR<sup>+</sup>/EpCAM<sup>+</sup> cells (12). ZEB1 has recently been demonstrated to limit oncogenic p53-driven DNA damage response in stem cells and thus increase the stem cells' intrinsic susceptibility to malignant transformation (19). ZEB1 cells coexpress the methionine sulfoxidase reducer (MSRB3), which protects against DNA damage (19). These observations raised the possibility that PROCR<sup>−</sup>/ZEB1<sup>+</sup> cells are naturally present at higher levels in the normal breasts of AA women and that failure to consider natural variation in gene expression pattern, influenced at least partially by genetic ancestry, could have an impact on identifying cancer-induced field effect on the adjacent normal breast. Measuring PROCR itself in the breast tissue is complicated because there are 4 haplotypes of PROCR due to SNPs and only 1 among them is a cell surface protein (20). Because ZEB1 expression is enriched in PROCR<sup>−</sup>/EpCAM<sup>−</sup> cells, we used ZEB1 as a surrogate marker for PROCR<sup>−</sup>/EpCAM<sup>−</sup> cells in unmanipulated breast tissues.

Normal breast tissue from biopsies of healthy AA women and those from AA and EA women are shown in Fig. 1A and statistical analyses are presented in Fig. 1B-D and in Table 1. Descriptive statistics of ethnicity, age, menstrual status, pregnancy and breastfeeding history, hormone replacement therapy, and family history of breast cancer for the KTB-normal cohort is shown in Supplementary Table S1. Highly discriminative 41-ancestry marker profiles of KTB-normal showed >75% African ancestry markers in samples from AA women and >80% EA ancestry markers in Caucasian women (Supplementary Fig. S1A; ref. 21). Characteristics of breast cancer in the tumor cohort are shown in Supplementary Table S2. ZEB1-expressing cells were localized outside the ductal structures of the normal breast and in the stromal part of the tumors (enlarged version on right side of Fig. 1A). KTB-normal breast tissue of AA women contained significantly higher levels of ZEB1<sup>+</sup> cells compared with KTB-normal breast of EA women (Fig. 1B). NATs of AA women showed a modest increase in ZEB1<sup>+</sup> cells compared with those of KTB-normal (Fig. 1C and D). The scenario was completely different in EA women; both NATs and tumors contained significantly higher levels of ZEB1<sup>+</sup> cells compared with KTB-normal tissue (Fig. 1C and D). NAT to tumor differences were noted only in EA women where an increase in ZEB1<sup>+</sup> cells was noted predominantly in ERα<sup>−</sup> tumors (Supplementary Table S3). Thus, ZEB1<sup>+</sup> cells are intrinsically higher in the normal breasts of AA women, whereas remarkably elevated ZEB1<sup>+</sup> cells in the breasts of EA women were observed only in the context of breast cancer. Increases in ZEB1<sup>+</sup> cells in KTB-normal tissue of AA women compared with EA women is less likely related to BMI differences. This was demonstrated by subdividing women above and below BMI of 30, irrespective of genetic ancestry: ZEB1 H-score but not positivity showed a marginal relationship (P = 0.04) to BMI above and below 30 (Supplementary Table S4).
ERα status in tumors influences differences between NATs and tumors

Although we observed differences in ZEB1, GATA3, and ERα expression between NATs and tumors (Figs. 1D, 2D, 3D; Supplementary Fig. S2D), interpretation of these data is difficult because of differences in characteristics of breast cancer subtypes, particularly ERα+ and ERα− (32). To determine whether ERα+ and ERα− tumors have distinct effects on NATs, FOXA1, GATA3, and
ZEB1 expression data in NATs and tumors were subdivided on the basis of ER status of the tumor and reanalyzed. ZEB1-positivity and H-scores were higher in ER<sup>−</sup> than ER<sup>+</sup> tumors compared with NATs (Supplementary Table S3). Despite small sample size, these differences were noted only in EA women with ER<sup>−</sup> breast cancers (Supplementary Table S3). With respect to FOXA1, H-score but not positivity was marginally higher in ER<sup>−</sup> tumors compared with NATs of EA women (Supplementary Table S3). ER<sup>−</sup> tumors of EA but not AA women showed a significant decline in both positivity and H-score of FOXA1 compared with NATs (Supplementary Table S3). ER<sup>−</sup> tumors but not ER<sup>+</sup> tumors showed further increase in GATA3 positivity and H-scores in EA women, which further confirm the role of GATA3 in hormonal regulation of breast cancer (Supplementary Fig. S2D; Supplementary Table S3). When the analyses was done with paired NAT and tumors, the above noted differences between NATs and tumors in ZEB1, GATA3, and FOXA1 levels remained significant, although sample size was too small to subdivide samples based on genetic ancestry (Supplementary Table S5).

NATs of AA and EA women show differing levels of CD8, PD1, and PDL1<sup>+</sup> cells

Results thus far point to proinflammatory state of NATs of EA women based on the known link between ZEB1 and inflammatory cytokines (33). To address this further, we stained the above TMAs with CD8 for T cells, CD68 for macrophages, and PD1 for immune cells. We also examined epithelial/tumor cells for PDL1. To address this further, we stained the above and PDL1 immunostains (data not shown). Therefore, we analyzed stained results between NATs of AA and EA women and between NATs and tumors. Representative staining patterns in NATs and tumors are shown in Fig. 4.

CD8 immunostaining was localized to inflammatory cells (T lymphocytes) and not to tumor cells in the breast cancer core. No background reactivity was observed in any case. NATs of AA women showed statistically significantly higher CD8 positivity compared with EA women (Fig. 5; Supplementary Table S6). The tumors in EA women had more CD8 immunostaining compared with corresponding NATs but such differences were not seen in the AA women.

CD8 staining was localized to macrophages in the breast cancer cores (Fig. 4). CD68 had lower positivity compared with CD8 by both visual and the Aperio-positive pixel reads. CD68 positivity was higher in tumors compared with their NATs (P = 0.02) in EA women but no such differences were noted in AA women (Fig. 5).

PD1 immunostaining was localized to immune cells only and no background staining was observed (Fig. 4). There was no staining of tumor cells. NATs of AA women contained significantly higher PD1<sup>+</sup> cells, similar to CD8<sup>+</sup> cells, compared with NATs of EA women (Fig. 6; Supplementary Table S6). PD1 staining did not show any differences between NATs and tumors in both groups (Supplementary Table S3).

PDL1 immunostaining was seen localized in the tumor cell cytoplasm and cell membrane (Fig. 4). In a few EA cases, only lymphocytes were stained. PDL1 staining of NATs of EA women was significantly lower than EA cases (Fig. 6; Supplementary Table S6). Although PDL1 staining did not differ between NATs and tumors of AA women, its levels were marginally lower in ER<sup>−</sup> tumors but not ER<sup>+</sup> tumors compared with NATs in case of EA women (Fig. 6; Supplementary Table S3). It is interesting that PD1 and PDL1 staining scores in NATs of AA is the reverse of the patterns seen in EA women. In summation, the immune environment in NATs is different from that in KTB-normal tissue with further differences between NATs and tumors, showing variations based on genetic ancestry.

**Discussion**

Recent studies have shown cancer-induced field defects influencing gene expression patterns in histologically normal tissues surrounding cancer (3, 34, 35). These observations raise a concern, as well as provide an opportunity for further investigation. The concern is the use of tumor adjacent normal as a "normal" control, whereas the opportunity pertains to the development of cancer-induced field defects in the adjacent normal as early markers of cancer. However, recent discovery of inter-individual differences in gene expression patterns due to SNPs in gene regulatory regions and genetic ancestry–dependent enrichment of SNPs with breast cancer protective or elevated risk characteristics necessitate the use of ancestry-matched control samples from healthy individuals to develop molecular features of tumor adjacent normal as cancer-initiation or progression markers (11, 13, 36, 37). Ethnicity contributing to inter-individual differences in normal biology is just beginning to be explored, as evident from a recent study that demonstrated distinct gut microbiota in different ethnic groups with shared geography (38).

Furthermore, genetic ancestry has been shown to influence mutation patterns in cancer (7). Resources available at the Komen Tissue Bank at Indiana University (Indianapolis, IN), namely ancestry-mapped breast tissues from >5,000 healthy women, should enable us to take these factors into consideration as we...
develop molecular features of NATs as cancer detection markers. Utilizing a small fraction of those tissues, we provide evidence for ancestry-dependent differences in the number of ZEB1$^+$ and GATA3$^+$ cells in the normal breast, as well as cancer-induced field effects on ZEB1, GATA3, and FOXA$^+$ cells in the tumor-adjacent normal tissue. Recently discovered functions of ZEB1 have raised considerable interest in this molecule within the oncology field. The

Figure 2.
FOXA1 expression pattern in KTB-normal, NATs, and in breast tumors. A, Representative IHC of KTB-normal, NATs, and tumors of AA and EA women. Enlarged view of a KTB-normal is shown on right (top). B, Differences in FOXA1 expression (positivity and H-score) between KTB-normal of AA and EA women. C, Differences between KTB-normal and NATs in AA and EA women. D, Differences between NATs and tumors in AA and EA women.
regulatory regions of this gene remain in a bivalent state, enabling the regulatory regions to respond readily to the tumor microenvironment and increase breast cancer plasticity and tumorigenicity (39). Another study showed elevated ZEB1 expression in normal breast stem cells, and it functionally protects stem cells from p53-mediated cell death in response to oncogene activation-induced DNA damage and promotes tumorigenicity with limited genomic instability (19). It was

Figure 3.
ERα expression pattern in KTB-normal, NATs, and in breast tumors. A, Representative IHC of KTB-normal, NATs, and tumors of AA and EA women. Enlarged view of a KTB-normal is shown on right (top). B, Differences in ERα expression (positivity and H-score) between KTB-normal of AA and EA women. C, Differences between KTB-normal and NATs in AA and EA women. D, Differences between NATs and tumors in AA and EA women.
also reported that ZEB1 is expressed in both tumor and stromal cells of the breast (40). ZEB1 directly increases the expression of proinflammatory cytokines such as IL6 and IL8, and it promotes vascular mimics of breast cancer cells by remodeling extracellular matrix (33, 41). We had previously demonstrated that cytokines such as TNF induce the expression of ZEB1 (42). These observations along with our unique observations of genetic ancestry–dependent differences in ZEB1+ cells in the normal breast, elevated number of ZEB1+ cells in NATs compared with healthy breast tissues of women of EA ancestry, and its localization outside the ductal structures raise several questions about the function of ZEB1+ cells in the normal and tumor adjacent normal breast. We have shown previously that cytokeratin-positive, PROCR+/EpCAM+ cells of the normal breast, which are enriched in the normal breast of AA women compared with cytokeratin-positive, PROCR+/EpCAM+ cells of the normal breast, which are enriched in the normal breast of AA women compared with EA women, express 50-fold higher ZEB1 compared with cytokeratin-positive, PROCR+/EpCAM+ cells of the breast (12, 43). Thus, we suspect that ZEB1+ cells in the normal breast correspond to PROCR+/EpCAM+ cells and that cancer-induced field effect leads to expansion/proliferation of such cells in the breast of EA women. Signaling pathways leading to proliferation of ZEB1+ cells in NATs of EA women are unknown, but the Wnt pathway is the prime suspect as it is activated in cells surrounding cancer due to altered DNA methylation (3). In this respect, Wnt and ZEB1 constitute a reciprocal feed-forward signaling loop where ZEB1 enhances TCF4/β-Catenin–mediated transcription and Wnt signaling converts ZEB1 from a transcription repressor to an activator (44).

The reason for an intrinsically higher number of ZEB1+ cells in AA women is unknown. TNBCs in AA compared with EA women display elevated Wnt pathway activation and it could be that Wnt pathway activity is intrinsically higher in AA women leading to elevated ZEB1 expression (8). It has also been demonstrated that vitamin D through Vitamin D Receptor (VDR) represses ZEB1 expression and serum vitamin D levels are significantly lower in AA than EA individuals (45, 46). Therefore, lower VDR activity and resulting increase in the activity of proinflammatory cytokines could be responsible for higher number of ZEB1+ cells in the normal breast of AA women, which needs further investigation.
In contrast to stemness-associated ZEB1, FOXA1, and GATA3, which are expressed predominantly in differentiated luminal cells, showed opposite pattern in AA women. While the normal breasts of AA women had higher number of FOXA1 \(^+\) cells compared with EA women, a decline in FOXA1 \(^+\) cells in NATs as a consequence of cancer field effect was observed only in AA women. How tumors cause downregulation of FOXA1 in NATs is unknown but could involve inflammatory cytokines, as cytokine-inducible transcription repressors such as TWIST1 repress FOXA1 expression (47, 48). In this regard, we observed genetic

**Figure 5.**
Statistical analyses of CD8 and CD68 positivity in NATs and tumors (T) of AA and EA women. All statistically significant differences are indicated with \(P\).
ancestry–dependent differences in the levels of immune cells in NATs; NATs of AA women contained an elevated number of CD8⁺ T cells and PD1⁺ immune cells compared with NATs of EA women. In addition, FOXA1 regulatory regions are highly susceptible for DNA methylation and transcriptional repression, particularly in the context of BRCA1 deficiency (49). Furthermore, ER⁻ tumours in AA women show elevated FOXA1 DNA methylation compared with ER⁻ tumours of EA women (30). Recent studies have also demonstrated racial differences in plasma levels of cytokines with CCL2, CCL11, IL4, and IL10 being higher in EA women, and IL1RA and IFNα2 being higher in AA women (50).

**Figure 6.** Statistical analyses of PD1 and PDL1 TPS scores in NATs and tumors (T) of AA and EA women. All statistically significant differences are indicated with $P$.

Differential expression of GATA3 in the normal breasts of AA and EA women is intriguing, as GATA3 is one of the major signaling molecules required for hormonal response and differentiation of normal breast epithelial cells (25). Our results suggest that hormonal- and differentiation-signaling networks show genetic ancestry–dependent differences and it is likely that ERα,GATA3–dependent transcriptional program is more active in the normal breast of EA compared with AA women. Whether such difference between EA and AA persist in ERα⁺ tumors is unknown and potentially worth investigating as it is relevant for response to antiestrogen therapy.
Collectively, data presented in this study suggest the need to consider the following aspects for cancer biomarker discovery: (i) NATs are molecularly abnormal and thus are not suitable as controls; (ii) these abnormalities can be detected only when true normal breast tissues are used as controls and differences in normal gene expression attributable to genetic ancestry are taken into consideration; (iii) ZEB1 and GATA3 show unique expression pattern in the normal breast, which is influenced by the genetic ancestry and could potentially be developed as biomarkers of breast cancer initiation of women of EA ancestry; and (iv) genetic ancestry has an influence on the immune environment of tumors as well as NATs.

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

Authors’ Contributions
Conception and design: H. Nakshatri, C. D’Souza-Schorey
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Acquisition of data (provided animals, acquired and managed patients, provided facilities, etc.): H. Nakshatri, M.L. Cox, M. Jacobsen, A.M.V. Storniolo
Analysis and interpretation of data (e.g., statistical analysis, biostatistics, computational analysis): H. Nakshatri, B. Kumar, H.N. Burney, M. Jacobsen, G.E. Sandusky, C. D’Souza-Schorey
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References


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