Comparison of Age Distribution Patterns for Different Histopathologic Types of Breast Carcinoma

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Abstract

Background: Historically, female breast carcinoma has been viewed as an etiologically homogeneous disease associated with rapidly increasing incidence rates until age 50 years, followed by a slower rate of increase among older women. More recent studies, however, have shown distinct age incidence patterns for female breast cancer when stratified by estrogen receptor (ER) expression and/or histopathologic subtypes, suggesting etiologic heterogeneity.

Materials and Methods: To determine if different age incidence patterns reflect etiologic heterogeneity (more than one breast cancer type within the general breast carcinoma), we applied “smoothed” age histograms at diagnosis (density plots) and a two-component statistical mixture model to all breast carcinoma cases (n = 270,124) in the Surveillance, Epidemiology, and End Results Program of the National Cancer Institute. These overall patterns were then reevaluated according to histopathologic type, race, and ER expression.

Results: A bimodal age distribution at diagnosis provided a better fit to the data than a single density for all breast carcinoma populations, except for medullary carcinoma. Medullary carcinomas showed a single age distribution at diagnosis irrespective of race and/or ER expression.

Conclusions: Distinct age-specific incidence patterns reflected bimodal breast cancer populations for breast carcinoma overall as well as for histopathologic subtypes, race, and ER expression. The one exception was medullary carcinoma. Of note, medullary carcinomas are rare tumors, which are associated with germ-line mutations in the BRCA1 gene. These descriptive and model-based results support emerging molecular data, suggesting two main types of breast carcinoma in the overall breast cancer population.

Introduction

Breast carcinomas among women are extremely diverse in clinical and histopathologic features, suggesting that these tumors also vary in etiology. Descriptive studies support this view. For example, data from the Danish Breast Cancer Cooperative Group and the Surveillance, Epidemiology, and End Results (SEER) Program posit that the classically recognized inflection point in age-specific breast cancer incidence rates overall around menopause (“‘Clemmesen’s Hook’”; ref. 1) may reflect the superimposition of two different rate curves (2–5), corresponding to estrogen receptor (ER)–negative and ER-positive tumors. Rates for ER-negative tumors increase rapidly until age 50 years, then flatten or decrease, whereas ER-positive tumors increase rapidly until age 50 years, then continue to increase at a slower pace.

In addition to the two age incidence patterns for ER expression, SEER data show three incidence rate curves according to histopathologic subtype (6–8). First, there is a rapid increase in incidence until age 50 years, then a slower increase for ductal, tubular, and lobular carcinomas, as for ER-positive tumors. Second, there is a rapid increase until age 50 years followed by leveling for medullary and inflammatory carcinomas, similar to curves for ER-negative tumors. Finally, there is a steady increase with aging for papillary and mucinous carcinomas, as is seen for epithelial tumors such as colorectal carcinoma (9, 10).

To explore further the age incidence patterns for different histopathologic types, we applied descriptive techniques (age-density plots) and a model-based analysis using two-component statistical mixture and single-density models.

Materials and Methods

We used the SEER Cancer Incidence Public-Use Database (November 2004 submission) to analyze invasive breast carcinoma among women, diagnosed during the years 1992 to 2002 (11). A total of 270,124 breast cancer cases were obtained from 13 SEER registries, including the Atlanta, Connecticut, Detroit, Hawaii, Iowa, New Mexico, San Francisco-Oakland, Seattle-Puget Sound, Utah, Los Angeles, San Jose-Monterey, rural Georgia, and Alaskan Native Tumor Registries. Data were stratified by histopathologic subtype, using codes from the International Classification of Diseases for Oncology, 3rd edition (ICD-O-3), and the WHO (12-14): duct carcinoma of no special type (duct NST; ICD-O-3 code 8500), tubular carcinoma (ICD-O-3 code 8211), lobular carcinoma (ICD-O-3 code 8520), medullary carcinoma (ICD-O-3 code 8510), inflammatory carcinoma (ICD-O-3 code 8530), papillary carcinoma (ICD-O-3 codes 8050, 8260, and 8503), and mucinous carcinoma (ICD-O-3 code 8480); other or unknown included all other ICD-O-3 codes.

Racial groups included White, Black, and Asian or Pacific Islanders. SEER did not record method of ER assay although immunohistochemical staining was probably used during the time period for this study (1992-2002). There was no centralized laboratory for hormone receptor expression for SEER; thus, each registry recorded hormone receptor expression as positive, negative, missing, borderline, or unknown. We combined missing, borderline, and unknown data into one group, designated as other or unknown.

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**Age-Specific Incidence Rates and Age Distribution.** Age-standardized incidence rates (2000 U.S. standard population) were calculated using SEER*Stat 6.1.4 and expressed per 100,000 woman-years. Age-specific incidence rates were charted for each histopathologic subtype for all women and by race (Fig. 1). Rates in Fig. 1 were plotted on a log-log scale, as originally described by Armitage and Doll (9, 10). We also approximated age incidence patterns with histograms using nonparametric probability density function curves in Figs. 2 and 3, as previously described (4, 15). In brief, the probability density function reflected a smoothed age distribution of cases at the time of primary breast carcinoma diagnosis. These density plots or curves are presented for each histopathologic subtype overall and by race (Fig. 2) as well as for each histopathologic subtype and ER expression (Fig. 3).

**Statistical Model.** We used a two-component mixture model to determine the probability density function for an early-onset versus a late-onset breast cancer population (or density; ref. 16). The probability density is defined as the derivative of a cumulative probability distribution function. The probability density function $f(x;y)$, where $x$ denotes the age at diagnosis, is then given by the mixture model $g(y;\theta) = f(y;\theta_0)(1 - p) + f(y;\theta_1)p$. In the mixture model, we interpreted $f(y;\theta_1)$ to be the probability density reflecting the early age at diagnosis population, whereas $f(y;\theta_1)$ was the probability density corresponding to the late age of onset population. The mixing probability $p$ represents the proportion of women within the breast carcinoma population with early age distribution. All the parameters in the model are denoted by $\theta$, and include the parameters of the component densities, the mixing proportion, and the parameter of the power transformation $i.e., \theta = (\theta_0, \theta_1, p, \lambda)$.

An extension of the two-component mixture model let $p$ depend on covariates such as race or ER expression through a logistic regression function. We employed two different parameterizations for the component densities $f(y;\mu, \sigma)$: normal probability densities $f = \phi(y;\mu, \sigma)$ with mean $\mu$, and SD $\sigma$, and semi-nonparametric densities, which multiply the normal density with a polynomial component, allowing for skewness and heavier tails than the normal density (17). We chose a polynomial of degree one, yielding $f(y;\mu, \sigma)(\alpha + \beta y)^r$, properly standardized.

To assess if the age distribution patterns were truly heterogeneous, we compared the fit of a mixture model with two components to that of a single density using the Akaike information criterion (AIC), which penalizes the log-likelihood by the number of parameters in the model (18, 19). Different parameterizations of the mixture model were compared using likelihood ratio test statistics, as these models are nested. Parameters in all models were estimated via a maximum likelihood procedure implemented in SAS 9.0 (SAS Institute, Cary, NC).

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**Results**

**Descriptive Statistics (Table 1A and B).** The SEER 13 Registry Database collected information for 270,124 invasive female breast carcinoma cases, diagnosed during the years 1992 to 2002. Similar to our previous study (8), infiltrating duct NST and lobular breast carcinomas were the most common histopathologic subtypes, accounting for 68.5% and 8.1%, respectively. Median age at diagnosis ranged from 51 years for medullary breast cancer to 81 years for mucinous breast carcinoma. The relative risk (RR, expressed as a rate ratio) of breast cancer for Black compared with White women was greatest for medullary [RR, 2.0; 95% confidence interval (95% CI), 1.8-2.2], papillary (RR, 1.8; 95% CI, 1.6-2.1), and inflammatory (RR, 1.4; 95% CI, 1.3-1.6) histologic subtypes. RR for Asian Pacific Islanders compared with White women was greatest for papillary breast carcinomas (RR, 1.2; 95% CI, 0.9-1.6). ER expression differed by histopathologic type. The RR of
ER-negative compared with ER-positive breast cancer ranged from 0.1 for mucinous type to 3.8 for medullary breast carcinoma.

Age-Specific Incidence Rates Curves. Age-specific incidence rates for all breast cases combined \((n = 270,124)\) and for each histopathologic type were stratified by White, Black, and Asian or Pacific Islander races. Similar to total rates for all breast cases combined (Fig. 1A), total rates for duct NST, tubular, and lobular breast carcinomas increased rapidly until age 50 years and then continued to increase at a slower pace (Fig. 1B-D). Rates among White women were generally similar to total rates, which is not surprising given that the White race accounted for the majority of breast cancer cases in SEER.

**Figure 2.** Age at diagnosis-density plots (or smoothed histograms) of female breast cancer diagnosed in SEER 13 Registry Database (1992-2002), stratified by race. Each plot includes a reference line at age 50 years, a reference line at age 70 years, and one histopathologic subtype with four superimposed charts: (i) total breast carcinoma cases for each histopathologic subtype; (ii) breast carcinoma cases defined by White race; (iii) breast carcinoma cases defined by Black race; and (iv) breast carcinoma cases defined by Asian or Pacific Islander race.

ER-negative compared with ER-positive breast cancer ranged from 0.1 for mucinous type to 3.8 for medullary breast carcinoma.

**Figure 3.** Age at diagnosis-density plots (or smoothed histograms) of female breast cancer diagnosed in SEER 13 Registry Database (1992-2002), stratified by ER-positive and ER-negative expression. Each plot includes a reference line at age 50 years, a reference line at age 70 years, and one histopathologic subtype with three superimposed charts: (i) total breast carcinoma cases for each histopathologic subtype; (ii) breast carcinoma cases defined ER-positive expression; and (iii) breast carcinoma cases defined by ER-negative expression.
In contrast to rates among White women, rates among Black women rose more slowly after age 50 years, whereas rates among Asian or Pacific Islander women plateaued or decreased after age 50 years for ductal, tubular, and lobular carcinomas.

Rates for medullary and inflammatory breast carcinomas increased until age 50 years, and then flattened or decreased, irrespective of race (Fig. 1E-F). Rates for papillary and mucinous breast carcinomas increased steadily with age, irrespective of race (Fig. 1G-H).

**Age Distribution.** The age distribution at diagnosis for all breast cases combined (n = 270,124) showed a bimodal pattern with incidence peaks (or modes) near ages 50 and 70 years (Figs. 2A and 3A). In addition, total cases for every histopathologic type except for medullary carcinoma were bimodally distributed.

### Table 1. Descriptive statistics for standard tumor characteristics in SEER 13 Registry Database among female breast cancer cases collected during 1992 to 2002

#### Table 1A:

<table>
<thead>
<tr>
<th></th>
<th>All breast cancer cases</th>
<th>Duct NST</th>
<th>Tubular</th>
<th>Lobular</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sample size</td>
<td>270,124</td>
<td>185,169</td>
<td>4,407</td>
<td>21,846</td>
</tr>
<tr>
<td>% Total cases</td>
<td>100.0</td>
<td>68.5</td>
<td>1.6</td>
<td>8.1</td>
</tr>
<tr>
<td>Median age (y)</td>
<td>62.0</td>
<td>61.0</td>
<td>62.0</td>
<td>66.0</td>
</tr>
<tr>
<td>Median tumor size (centimeters)</td>
<td>1.6</td>
<td>1.6</td>
<td>0.8</td>
<td>2.0</td>
</tr>
<tr>
<td>Rate (SE)</td>
<td>132.4 (0.26)</td>
<td>91.0 (0.21)</td>
<td>2.19 (0.03)</td>
<td>10.7 (0.07)</td>
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</tbody>
</table>

#### Table 1B:

<table>
<thead>
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<th></th>
<th>Medullary</th>
<th>Inflammatory</th>
<th>Papillary</th>
<th>Mucinous</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sample size</td>
<td>2,619</td>
<td>3,004</td>
<td>1,678</td>
<td>6,881</td>
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<tr>
<td>% Total cases</td>
<td>1.1</td>
<td>1.2</td>
<td>0.7</td>
<td>2.8</td>
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<tr>
<td>Median age (y)</td>
<td>51.0</td>
<td>56.0</td>
<td>70.0</td>
<td>71.0</td>
</tr>
<tr>
<td>Median tumor size (cm)</td>
<td>2.0</td>
<td>5.3</td>
<td>1.5</td>
<td>1.5</td>
</tr>
<tr>
<td>Rate (SE)</td>
<td>1.29 (0.03)</td>
<td>1.50 (0.03)</td>
<td>0.81 (0.02)</td>
<td>3.31 (0.04)</td>
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</table>

#### Table 1C:

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<th>Variable</th>
<th>N (%)</th>
<th>Rate</th>
<th>RR (95% CI)</th>
<th>N (%)</th>
<th>Rate</th>
<th>RR (95% CI)</th>
<th>N (%)</th>
<th>Rate</th>
<th>RR (95% CI)</th>
<th>N (%)</th>
<th>Rate</th>
<th>RR (95% CI)</th>
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</table>

#### Table 1D:

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<th>Age (y)</th>
<th>N (%)</th>
<th>Rate</th>
<th>RR (95% CI)</th>
<th>N (%)</th>
<th>Rate</th>
<th>RR (95% CI)</th>
<th>N (%)</th>
<th>Rate</th>
<th>RR (95% CI)</th>
<th>N (%)</th>
<th>Rate</th>
<th>RR (95% CI)</th>
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#### Table 1E:

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<th>ER receptors</th>
<th>N (%)</th>
<th>Rate</th>
<th>RR (95% CI)</th>
<th>N (%)</th>
<th>Rate</th>
<th>RR (95% CI)</th>
<th>N (%)</th>
<th>Rate</th>
<th>RR (95% CI)</th>
<th>N (%)</th>
<th>Rate</th>
<th>RR (95% CI)</th>
</tr>
</thead>
</table>

**NOTE:**

1. N, sample size; 2. Rate, age-adjusted (2000 U.S. standard) incidence rate per 100,000 woman-years; 3. RR, expressed as a rate ratio where a high-risk characteristic is compared to a low-risk characteristic with an assigned RR of 1.0.
AIC mixture & $-1107039.0$ & $-752190.4$ & $-720027.2$ & $-17053.0$ & $-86327.3$  
\hline
Sample size & 270,124 & 185,169 & 4,407 & 21,846 &  
\hline
Rate (SE) & 132.4 (0.26) & 91.0 (0.21) & 2.2 (0.03) & 10.7 (0.07) &  
\hline
\end{tabular}

Key: Rate, age-adjusted (2000 US standard) incidence rate per 100,000 woman-years; SE, standard error; \( p \) for the early-onset group given race; \( p \) for the early-onset group given ER.

When total cases for each histopathologic subtype were stratified by race and ER expression, the density plots showed varying mixtures of bimodal breast carcinoma types with early-onset and late-onset modes or peak frequencies (Figs. 2B-H and 3B-H). These findings were also confirmed when we fit the mixture models stratified by race and ER. However, whereas the fraction of cases in the early-onset and late-onset groups changed by race and ER among each histopathologic subtype, the modal ages remained unchanged, near ages 50 and 70 years.

Given that all of the parameters in the mixture model, with the exception of the mixing proportion, were unaffected by the covariates, we incorporated race and ER expression into the model via logistic mixing proportions. For all histopathologic subtypes, except tubular carcinomas, Asian or Pacific Islanders were more likely than either Whites or Blacks to have early-onset tumors with peak incidence at age 50 years. For example, among duct NST (Table 2), the probability of a single Asian or Pacific Islander case belonging to the early-onset group was 0.80 (SE < 0.01), compared with 0.47 (SE < 0.01) for a single White case and 0.27 (SE < 0.01) for a single Black case. Similarly, ER-negative tumors were more likely than ER-positive tumors to show early-onset modes at age 50 years.

\begin{table}[h]
\centering
\begin{tabular}{lcccccc}
\hline
Variable & \multicolumn{2}{c}{All breast cancer cases} & \multicolumn{2}{c}{Duct NST} & \multicolumn{2}{c}{Tubular} & \multicolumn{2}{c}{Lobular} \\
\hline
Sample size & 270,124 & 185,169 & 4,407 & 21,846 &  
\hline
Rate (SE) & 132.4 (0.26) & 91.0 (0.21) & 2.2 (0.03) & 10.7 (0.07) &  
\hline
\end{tabular}
\caption{Estimates for early-onset and late-onset modes and group probabilities given a specified histopathologic subtype of breast carcinoma, race, or ER expression.}
\end{table}
Parenthetically, we have observed similar bimodal incidence patterns according to tumor size, axillary lymph nodes, tumor grade, and progesterone receptor expression (20). That is, large tumors were more likely than small tumors to show early-onset modes at age 50 years. Tumors with positive axillary lymph nodes, high grade, and negative progesterone receptor expression were also more likely to show early-onset modes at age 50 years than were breast cancers with negative nodes, low grade, and positive progesterone receptor expression. In sum, high-risk tumors (large size, positive lymph nodes, high grade, negative ER and progesterone receptor expression) have predominantly early-onset age distributions at diagnosis, whereas low-risk tumors (small size, negative lymph nodes, low grade, positive ER and progesterone receptor expression) have mostly late-onset age distributions at diagnosis.

Discussion

Emerging molecular analyses of female breast carcinoma suggest the existence of two main pathways for mammary carcinogenesis based on epithelial cellular origin (luminal or basal) and/or ER expression (positive or negative; refs. 5, 21). There is (a) derivation of tumors from stem cells committed to luminal differentiation and ER expression and (b) development of neoplasms from stem cells programmed to display basal differentiation and lacking ER expression (22,24). If confirmed, these observations may form the basis for revised conceptual frameworks and, therefore, should be validated in the general population.

Indeed, our analysis of population-based SEER data provided evidence for two main breast cancer populations according to age at onset, mixed within breast cancer overall. One breast cancer type was mostly early-onset with peak incidence near age 50 years. The second breast cancer type was largely late-onset with its mode occurring at age ~70 years. Although covariates had varying proportions of early-onset and late-onset age distributions, modal ages remained near 50 and 70 years, respectively.

The age structure of the breast cancer population (Figs. 2 and 3) seemed to define the shape of the age-specific incidence rate curve (Fig. 1). For example, medullary and inflammatory carcinomas with predominant early-onset age distributions (Figs. 2E-F and 3E-F) produced age-specific incidence rate curves that flattened or decreased after age 50 years (Fig. 1E-F). Papillary and mucinous carcinomas with late-onset age distributions (Figs. 2G-H and 3G-H) yielded age-specific rate curves that rose steadily before and after age 50 years (Fig. 1G-H). Finally, duct NST, tubular, and lobular carcinomas with bimodal breast cancer populations (Figs. 2B-D and 3B-D) produced intermediate age-specific rate curves (i.e., age-specific rates increased rapidly until age 50 years, and then continued to increase at a slower pace; Fig. 1B-D).

In their seminal study, Sorlie et al. (25) used a 534 “intrinsic” gene set to subdivide ER-positive and ER-negative tumors into five subtypes: luminal A, luminal B, basal-like, HER2-overexpressing, and “normal-like” subgroups. Luminal A and luminal B are ER-positive subtypes, whereas basal, HER2, and the normal-like subgroups are ER-negative subtypes. A straightforward interpretation of their results regards each molecular signature as a distinct breast cancer entity (24, 25). To test this concept, we constructed and analyzed age-density plots using breast cancer cases (n = 122) from Sorlie et al.’s data set (Fig. 4). After matching molecular portraits with ages at diagnosis from their supplementary data (25), luminal A and B cases had a predominant late-age distribution with a mode at age 74 years (SE = 1.7), as did ER-negative tumors in SEER. Therefore, and despite their small sample size, these results show only two modes with small variances, occurring at ages similar to those in the SEER population. Thus, where SEER’s two age-specific incidence rate patterns for ER and three rate patterns for histopathologic subtypes reflected bimodal mixtures of early-onset and late-onset breast cancer populations, Sorlie et al.’s molecular phenotypes also displayed bimodal breast cancer populations.

Our study was limited by the usual concerns related to analyses of registry data: nonstandardization of histopathologic diagnosis and/or ER testing and incomplete data collection. However, it is reassuring that the age distribution for different histopathologic subtypes and known ER expression is similar to that in other reported series (26). In previous studies, we have also shown that cases with unknown data are generally similar to all cases combined (20, 27-29). Moreover, in a sensitivity analysis, we examined age incidence patterns in different SEER registries, over different time periods, and among different birth cohorts. SEER bimodal age incidence patterns were robust to geographic location as well as to calendar period and/or birth cohort effects (data not shown). Tarone and Chu (3) also showed that SEER’s distinct ER-positive and ER-negative incidence patterns were unaffected by adjustments for calendar period and birth cohort effects or artifacts, implying a true age-related biological phenomenon. Moreover, bimodal female breast cancer populations were first described by von Pirquet in 1930 (30) and have been observed worldwide in Africa (31), Taiwan (32), Italy (33), New Zealand (34), Europe, and America (35), suggesting a universal breast cancer phenomenon.

Figure 4. Age at diagnosis-density plots (or smoothed histograms) were applied to data provided in Supplementary Table S2 (http://www.pnas.org) for all tissue samples in the extended Norway/Stanford extended data set, described by Sorlie et al. (25): (i) all tissue samples (n = 122) in the Norway/Stanford data set; (ii) luminal A and luminal B cases in the Norway/Stanford data set; and (iii) basal-like and HER2-overexpressing cases in the Norway/Stanford data set.
In conclusion, age incidence patterns showed bimodal early-onset and late-onset breast cancer types, mixed within SEER's overall breast cancer population. Bimodal age distribution—the occurrence of two incidence peaks or modes—is always of interest. It suggests etiologic heterogeneity due to at least two biological subtypes or causal pathways (36). Although bimodal age incidence patterns are acknowledged for malignancies such as Hodgkin's lymphoma (37, 38), bimodality is not very well established for breast cancer. Moreover, whereas the two modes for Hodgkin's disease are widely separated in early and late adulthood, bimodal breast cancer has a narrower range of ages at 50 and 70 years.

Of note, the modal ages for breast cancer of 50 and 70 years were consistent across molecular portraits, race, ER, as well as other tumor characteristics (20); however, the relative distributions for early-onset and late-onset breast cancer types varied by covariate. In general, high-risk tumors such as ER-negative expression were surrogates for bimodal breast cancer populations, shifted towards the early-onset mode near age 50 years. Low-risk tumors such as ER-positive expression were surrogates for bimodal breast cancer, weighted towards the late-onset mode at age 70 years.

Additional etiologic clues may be provided by medullary carcinoma, the one histopathologic subtype with a single mode at age 51 years. Medullary carcinomas are rare tumors that have been associated with early-onset hereditary breast carcinomas due to the BRCA1 gene (39, 40). Given that age at onset is the strongest predictor for a genetic versus an acquired breast cancer phenotype (41, 42), bimodality in the general breast cancer population might be a reflection of early-onset hereditary (or familial) versus late-onset nonhereditary (acquired or sporadic) breast cancer types. Because age 50 years is both the early-onset mode and a valid menopausal surrogate (43, 44), we further speculate that premenopausal exposures will have greater effect on early-onset than on late-onset breast cancers. Future population-based studies that include assessment of family history, breast cancer risk factors, environmental exposures, standardized histopathology review, and molecular characterization are needed to examine the conceptual framework of a two-class or bimodal breast cancer model.

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