Award Number: DAMD17-02-1-0473

TITLE: Benign Breast Disease: Toward Molecular Prediction of Breast Cancer Risk

PRINCIPAL INVESTIGATOR: Lynn C. Hartmann, M.D.

CONTRACTING ORGANIZATION: Mayo Clinic, Rochester
                                    Rochester, MN 55905

REPORT DATE: June 2007

TYPE OF REPORT: Annual

PREPARED FOR: U.S. Army Medical Research and Materiel Command
                          Fort Detrick, Maryland 21702-5012

DISTRIBUTION STATEMENT: Approved for Public Release;
                          Distribution Unlimited

The views, opinions and/or findings contained in this report are those of the author(s) and should not be construed as an official Department of the Army position, policy or decision unless so designated by other documentation.
# Benign Breast Disease: Toward Molecular Prediction of Breast Cancer Risk

**Optimal early detection and prevention strategies for breast cancer are predicated on our ability to identify individuals at significantly increased risk for this disease. The purpose of this Center is to bring molecular risk prediction for breast cancer into the clinical area. This will require progress on three fronts of scientific endeavor: (i) Establishment of a tissue repository of benign breast disease; (ii) Assessment of potential biomarkers of risk in this tissue set and (iii) Discovery of new, potentially relevant biomarkers of risk. We have made significant progress on these aims. Our current cohort comprises 9,376 women, 758 (8%) of whom have been diagnosed with breast cancer since the time of their benign biopsy. We established our tissue repository of benign breast tissue and have collected the subsequent breast cancer tissue. We assessed the significance of benign histology in predicting risk of future breast cancer, examining in detail the role of proliferative disease, atypia, papillomas, radial scars and involution. We explored the link between centrosome amplification, COX-2 expression and breast cancer outcomes and are currently exploring the significance of p16, ER and MIB-1. We have begun our work with Wayne State to characterize the histopathology in a cohort of African American women. Our focus in 2007-2008 will be on the Wayne State cohort and exploring additional molecular markers.**

**Subject Terms:** benign breast disease, biomarkers, histology, breast cancer

**Security Classification:**

- **a. REPORT:** U
- **b. ABSTRACT:** U
- **c. THIS PAGE:** U

**Limitation of Abstract:**

- **UU**

**Number of Pages:**

- **33**

**Telephone Number:**

- **USAMRMC**

<table>
<thead>
<tr>
<th>Security Classification of:</th>
<th>Limitation of Abstract</th>
<th>Number of Pages</th>
<th>Telephone Number</th>
</tr>
</thead>
<tbody>
<tr>
<td>a. REPORT</td>
<td>U</td>
<td></td>
<td></td>
</tr>
<tr>
<td>b. ABSTRACT</td>
<td>U</td>
<td></td>
<td></td>
</tr>
<tr>
<td>c. THIS PAGE</td>
<td>U</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
# Table of Contents

- Introduction ................................................................................................................. 4
- Body ............................................................................................................................... 4-10
- Key Research Accomplishments .................................................................................. 11
- Reportable Outcomes ..................................................................................................... 11-13
- Conclusions .................................................................................................................. 13
- Appendices ..................................................................................................................... 14-33
INTRODUCTION

Our fifth year Center of Excellence report details a total of 52 months of work involving human subjects. Delays during initial approval processes led to some delay in the start-up funding for the human subjects portion of the grant.

Three main aims of scientific activity exist within our Center of Excellence: 1) the establishment of a large tissue repository from a retrospective cohort of women with benign breast disease (BBD) (1967-1991); 2) the application of potential biomarkers of risk to this archival tissue set; and 3) the discovery of new, potentially relevant biomarkers of risk in fresh and frozen specimens of BBD. The Center includes a multi-institutional team of basic scientists, pathologists, epidemiologists, clinicians, statisticians, and advocates (Mayo Clinic; University of California San Francisco (USCF); Wayne State).

Task 1: Establish Retrospective Cohort of BBD and Nested Case-Control Study

A. Complete cohort follow-up
   We reported the details in our 2006 report. This task has been completed.

B. Validate reported breast cancers
   We reported the details in our 2006 report. This task has been completed.

C. Match appropriate controls to known breast cancer cases
   We described this process in our 2004 report. This task has been completed.

D. Construct test set for preliminary evaluation of markers
   We described the construction of our test set in our 2004 report. This subset consists of 124 cases and their two closest controls selected from the entire study period.

E. Construct validation set from remaining breast cancer cases, each matched with two controls.
   The remaining cases and controls will serve as the validation set.

Task 2: Biomarkers in Archived Tissues from Cases and Controls

A. Retrieve tissue slides/blocks of BBD specimens for all cases and controls
   We reported details in our 2006 report. This task has been completed.

B. Characterize benign histopathology
   1. General findings
      In 2006 we reported the benign histology for our entire cohort. This objective has been completed. We published the general histology findings in July 2005 in the New England Journal of Medicine.

   2. Atypia
      We reported on our atypia results in our 2006 report. These results were just published in the July 1, 2007 issue of Journal of Clinical Oncology (Appendix A). The following highlights the major findings:
Although atypical hyperplasia is a well-established risk factor for subsequent breast cancer, data regarding long-term absolute risk and factors for risk stratification are needed. Estimates of absolute risk with long-term follow-up are not well established. We found the following:

- With a mean follow-up of 13.7 years, 66 breast cancers (19.9%) occurred among 331 women with atypia. The relative risk of breast cancer with atypia was 3.88 (95% CI 3.00-4.94).
- Marked elevations in risk were seen with multifocal atypia (e.g., three or more foci with calcifications ((relative risk - 10.35, 95% CI, 6.13 – 16.4)).
- Multiple foci of atypia and the presence of histologic calcifications may indicate “very high risk” status, exceeding 50% risk at 20 years.
- Relative risk was higher for younger women, under 45 years of age (Relative risk 6.76, 95% CI 3.24-12.4)
- Risk was similar for atypical ductal and atypical lobular hyperplasia.
- Breast cancer risk remained elevated over 20 years, and the cumulative incidence approached 35% at 30 years.
- A positive family history does not further increase risk in women with atypia.

3. Papillomas
These data and the publication of these data were reported in 2006.

4. Involution
In our 2006 report we identified that the extent of lobular involution in breast tissue is an important risk indicator for the development of breast cancer. These results were recently published in the Journal of the National Cancer Institute in November, 2006 (see appendix B). We fond the following:

- Risk of breast cancer was associated with the extent of involution. Comparisons in relation to the Iowa SEER population revealed that the relative risk for women with no involution was 1.88 (95% CI = 1.59-2.21), partial involution 1.47 (CI = 1.33 – 1.61) and for complete involution the relative risk was 0.91 (CI = 0.75-1.10).
- Increased involution was positively associated with increased age and inversely associated with parity.
- The significant reduction in breast cancer risk noted with involution also existed in women at “high risk” based on atypia or young age.

We are pleased with the accompanying editorial (see appendix C) in which the authors Henson DW, Tarone RE, Nsouli H from George Washington University Cancer Institute asserted:

- “It truly is a remarkable event when traditional pathologic observations lead to new ideas about the prevention of cancer.”
- “It is the first study…..to substantiate a hypothesis….that delayed involution is a major risk factor for breast cancer…..”
- “One of the most striking findings in the study…..is the degree to which the strong association between extent of involution and breast cancer risk was independent of all known breast cancer risk factors that were investigated.”
- “Results of the Mayo study provide a new paradigm for breast cancer research and prevention.”


5. Radial Scars
The significance of radial scars to subsequent risk of breast cancer has been debated. Radial scars (RS) are benign breast lesions of uncertain etiology. The growth pattern in RS can resemble breast cancer and on mammogram a RS can be difficult to distinguish from breast cancer, prompting a biopsy. The literature is mixed about the risk of developing breast cancer following the diagnosis of a RS, leading to our interest in examining the significance of RS in the subsequent development of breast cancer. We found no increased breast cancer risk for women with radial scars when compared to the risk already present due to proliferative disease with or with atypia. Breast cancer risk was also not affected by the size or number of RS lesions.

This manuscript has been published online: Berg JC, Visscher DW, Vierkant RA, Pankratz VS, Maloney SD, Lewis JT, Frost MH, Ghosh K, Degnim AC, Brandt KR, Vachon CM, Reynolds CR, Hartmann LC. Breast cancer risk in women with radial scars in benign breast biopsies. *Breast Cancer Research and Treatment*. Published online May 22, 2007

C. Prepare tissue slides for biomarker analyses
Tissue slides have been prepared for the test set and two other subgroups of interest: women whose breast cancer occurred within 5 years of their diagnosis (n = 174) and women whose histopathology revealed atypia (n = 336).

D. Perform IHC of molecular markers
Our focus continues to be on the earliest possible changes that we might detect in these "premalignant" lesions. There is certainly no consensus on this point. Our decision was to begin with COX-2, ER alpha, MIB-1, gamma tublin and cyclin-D, and the test set and atypia subgroup have been stained for these markers.

1. COX-2 in atypia
We reported these findings in our 2006 report. A manuscript has been prepared and is being circulated among the coauthors. The following highlights the points that will be made:

- COX-2 is an enzyme responsible for the elaboration of multiple bioactive mediations important in carcinogenesis. It has been shown to be overexpressed in DCIS and invasive breast cancer. We sought to evaluate its expression in atypical hyperplasia. Our hypothesis is that the expression of COX-2 will be increased in subjects with atypical hyperplasia (AH), and will be associated with subsequent breast cancer development. The pharmaceutical availability of COX-2 inhibitors makes this a particularly interesting marker to explore due to the possibility of subsequent trials to examine the effectiveness of COX-2 inhibitors in women at high-risk for the development of breast cancer.
- The intensity of immunostaining was associated with the type of AH (p<0.001).
- Most atypical ductal hyperplasia (ADH) (77%) had no or weak (< 1) COX-2 staining
- Most atypical lobular hyperplasia (ALH) stain intensity was 2+ or 3+ positive (61.4%)
- Strong immunostaining was more likely with increasing age (p<0.01)
  - Of the women who were <45 years at the time of biopsy, 20% had 2+ or 3+ staining
  - Of the women >55 years, 52.3% had showed 3+ staining, 33.8% had lesions that were 2+
- COX-2 intensity was correlated with the increasing number of AH foci (p=0.02)
Among the 39 subjects who had three or more foci, 61.6% had strong (2+ or 3+) staining
- Of the subjects who had a single AH focus (131 total), 35.9% had strong staining
- The relative risks for subsequent breast cancer compared to a control population from Iowa SEER data, were 2.63 for ≤1 stain intensity, 3.56 for 2+ and 5.66 for 3+ (see Table below)

Table 1. Association of COX-2 staining intensity with risk of breast cancer after the diagnosis of atypical hyperplasia.

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>No. of women</th>
<th>No. of person-years</th>
<th>No. of Observed Events</th>
<th>No. of Expected Events</th>
<th>Relative Risk (95% CI)²</th>
</tr>
</thead>
<tbody>
<tr>
<td>Overall</td>
<td>235</td>
<td>3265</td>
<td>41</td>
<td>12.4</td>
<td>3.31 (2.38-4.49)</td>
</tr>
<tr>
<td>COX-2 Staining Intensity</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>0-1+</td>
<td>130</td>
<td>1869</td>
<td>18</td>
<td>6.9</td>
<td>2.63 (1.56-4.15)</td>
</tr>
<tr>
<td>2+</td>
<td>71</td>
<td>1004</td>
<td>14</td>
<td>3.9</td>
<td>3.56 (1.94-5.97)</td>
</tr>
<tr>
<td>3+</td>
<td>34</td>
<td>391</td>
<td>9</td>
<td>1.6</td>
<td>5.66 (2.59-10.75)</td>
</tr>
</tbody>
</table>

1: number of events expected on the basis of Iowa Surveillance, Epidemiology and End Results (SEER) data.
2: all analyses account for the effects of age and calendar period. CI denotes confidence interval.

2. ER
The estrogen receptor is essential to mediate the growth regulatory signals of estrogen in normal breast tissue and serves as a therapeutic target and predicative factor in breast cancer. The extent of ER staining in a well-characterized cohort of women with atypia, to our knowledge, has not yet been reported.

We are using the Automated Cellular Imaging System III (ACIS) to evaluate the intensity and percent ER staining in 231 women with atypia. This system is able to provide automated quantification of biomarkers. The areas of atypia were identified by the study pathologist and read into ACIS with reports produced for each identified area of interest. The preliminary analysis of the 10 most intensely stained areas shows:
- A mean of 56 percent stained cells (standard deviation 30.78, range 0.00 – 99.99)
- The mean intensity was 114 (standard deviation 28.97, range 0-206 (possible range 0 -256)).
- A linear multivariate mixed model examined percent staining and intensity differences based on atypia type (lobular, ductal), cancer status, and year of biopsy after controlling for repeated measures within a woman.
  - The only difference based on this initial analysis was stronger intensity of staining and greater percentage of staining of atypical ductal hyperplasia compared to atypical lobular hyperplasia (intensity: ADH mean of 117.68, standard error 2.5224 and ALH mean of 100.32, standard error 2.4537, p<0.0001; percentage: ADH mean of 64.1887, standard error 2.7860 and ALH mean of 44.2121, standard error 2.6720, p<0.0001).

We are currently completing the ER assessments for all areas of atypia and will finalize analyses subsequently.
3. Proliferation Status
Upon completion of the ER readings, we are set up to use the ACIS for measuring MIB-1 (Ki-67) stained areas. All the slides have been stained for Ki-67 and are ready for analysis.

4. p16
Dr. Tlsty from UCSF put forward p16 as the most significant marker to take forward into testing. We have stained the atypia slides for p16 and they have been read. Analysis will be forthcoming.

E. Perform centromere studies.
These data were presented in our 2006 report.

Task 3: Discovery - In Vitro Culturing and Gene Profiling Studies

A. Identify appropriate frozen proliferative BBD specimens at Mayo and Wayne State for profiling.
The purpose of these studies is to identify new, potentially relevant biomarkers in benign breast disease, markers that might correlate with subsequent breast cancer risk. When our grant was submitted, the technology was not available to do profiling studies in paraffin-embedded tissue (such as our BBD resource) and hence, we described doing profiling in frozen samples of BBD. A serious limitation of that approach, however, is that we do not have outcome information for our frozen repository samples, since these were accrued recently, and insufficient time has elapsed for the development of breast cancer. Fortunately, genomic profiling technology has proceeded significantly and there are now platforms available for us where microdissected, paraffin-embedded samples can be run. We are working currently to identify the quantity and quality of DNA and RNA that can be obtained from the paraffin-embedded samples.

B. Obtain fresh BBD tissue from appropriate patients at Mayo and Wayne State for culturing in vitro at UCSF. Revised 1/07 Obtain fresh BBD tissue from appropriate patients at Mayo for culturing in vitro at UCSF.
Forty-four samples were sent from Mayo to UCSF. Five of these samples were lost to contamination.

Multiple efforts to implement a prospective collection of fresh tissue in African-American women at Wayne State proved to be logistically impossible to launch. Thus, we have moved to develop a retrospective study in an African-American cohort at Wayne State, modeled after the Mayo (Caucasian) cohort. Through a collaboration with Dr. Hind Nassar, a junior pathologist at Wayne State, an IRB-approved protocol has been developed, to access paraffin-embedded samples of benign breast disease (BBD) from African-American women at Wayne State from 1992-2001. This will allow us to begin to look at the problem of BBD in African-American women. Moreover, because the population there is covered through the Detroit SEER database, we will have information about cancer outcomes. See task 5.

C. Culture BBD specimens and document their growth characteristics.
These data were reported in 2005.

D. Compare genomic expression levels of DCIS markers in BBD tissues.
We reported on this task in 2006. Task completed.

E. Profile BBD specimens.
We reported on this task in 2006. Task completed.
Task 4: Statistical Analyses

A. Establishment of relational database
This task is complete. The database is the foundation for tracking all tissue samples; entering clinical, pathologic, and molecular data; and analyzing results.

B. Enter epidemiologic and histopathologic data
This task is complete.

C. Enter culturing data (proportion of cells that break through proliferation barriers; slope of curve, etc.)
These data were entered as collected at UCSF.

D. Enter molecular data from culturing experiments (methylation of p16, p53 status, % proliferation versus apoptosis, etc).
These data were entered as collected at UCSF.

E. Enter gene profiling data.
These data were entered as collected at UCSF.

F. Calculate hazard function for breast cancer by age at BBD, family history, histology, and molecular marker data.
We have examined breast cancer risk by age at BBD, family history, histology [degree of proliferation, atypia yes/no, extent of involution, radial scar yes/no, presence of papilloma(s)], COX-2 expression, and centrosome status. We have summarized the findings in earlier sections of this report.

G. Assess accuracy of Gail model.
The Gail model is currently the main tool used in the clinical setting for risk assessment in patients with atypia. This is despite the fact that it has not been validated in this group of patients. We evaluated the Gail model in our group of women with atypia (N = 331). We used this model to predict 5 year and follow-up specific risks for each woman. The Gail model over-estimated the number of breast cancers that would occur in the first 5 years after biopsy (12.8 predicted, 8 occurred). However, we found that the Gail model underestimated the risk of breast cancer in women with atypia when using the current follow-up age of our participants (Gail model predicted 31.7 breast cancer while 58 occurred). Additionally, we found the concordance between Gail model individual-specific predicted outcomes and observed outcomes to show only modest improvement over chance alone (c-statistic 0.57, 95% CI: 0.52-0.63, p=0.011). This has significant implications for clinical practice. Healthcare providers should be cautious when using the Gail model in counseling patients with atypia regarding their risk of developing breast cancer. We are in the process of submitting a manuscript to report these findings.

-9-

Figure 1. Distributions of Gail model risk probabilities. Plot contains estimates for individualized risk at the end of the available follow-up. As risk predictions depend on age at BBD, and length of follow-up, the risk predictions were corrected for these factors prior to comparison.
Analyze expression data. This past year we have focused on COX-2, ER, p16 and MIB1(Ki-67). We have reported our current findings under Task 2.


A. Identify African-American women at Wayne State University who had a breast biopsy with benign results between 1992 and 2001.
One thousand one hundred forty-five women who had a benign breast biopsy during 1999-2000 were identified. To date 240 women have been identified for 1998.

B. Retrieve slides/blocks of BBD specimens.
Three hundred twenty slides have been obtained thus far.

C. Characterize benign histology of epithelium.
Our programmer developed a data entry tool with drop down option boxes for Dr. Nassar’s use. Dr. Nassar worked closely with Dr. Dan Visscher to ascertain consistency in definition and reading of cohort slides using the study pathology form. Dr. Nassar has read and entered the histology for 225 of these slides.

D. Cross list with Detroit SEER database to identify subsequent breast cancers.
Dependent on A-C.

E. Data clean-up, compare age, histology, involution status, and resulting risk with Mayo
Caucasian-American cohort and determine involution status by age of patient. To begin once A – F accomplished.

KEY RESEARCH ACCOMPLISHMENTS

- We identified the degree of risk associated with the common benign epithelial entities and the extent to which age at biopsy and family history influence the risk of breast cancer in women with proliferative or atypical lesions. The highest risk was among women who had proliferative disease with atypia, especially those of younger age.

- We identified a marked increased risk of breast cancer in women with three or more foci of atypia, especially for three or more foci with calcifications. Also, risk was higher in women diagnosed with atypical hyperplasia before age 45. Among women with atypia, risk was not affected by family history.

- We identified that a single papilloma without atypia imparts an increased risk of developing a subsequent carcinoma similar to other forms of proliferative breast disease without atypia. Atypical papilloma, particularly in the setting of multiple papillomas, imparts a breast cancer risk similar to or greater than conventional atypical ductal/lobular hyperplasias.

- We identified that the extent of lobular involution in breast tissue is an important risk indicator for the development of breast cancer. Increasing degrees of involution result in a significant reduction in breast cancer risk, even in women at “high risk” based on atypia or young age.

- We found that intense COX-2 expression is associated with a significantly greater likelihood of a subsequent breast cancer in women with atypia and represents one potential molecular target for chemoprevention strategies.

- We found no increased breast cancer risk for women with radial scars compared to the risk already present due to proliferative disease with or without atypia.

- We identified that centrosome amplification is seen more frequently in higher risk benign lesions (e.g. atypia) and is infrequently seen in non-proliferative lesions and in proliferative lesions without atypia.

- We have identified intense p16 expression as a biomarker that identifies women with a significantly greater likelihood for recurrence after lumpectomy only for DCIS. This biomarker is presently being applied to the BBD cohort.

- We found the Gail model to predict only slightly higher than chance alone the breast cancer risk of women with atypia. The model underestimated lifetime risk and current risk of our cohort of women with atypia.

REPORTABLE OUTCOMES

Manuscripts


Presentations
Poster Presentation at annual meeting of the United States and Canadian Academy of Pathology, Atlanta, GA, Feb. 11-17, 2006.
• Milanese TR, Hartman LC, Vierkant RA, Maloney SD, Frost MH, Pankratz VS, Visscher DW. The impact of lobular involution on breast cancer risk.

Podium Presentation at annual meeting of the United States and Canadian Academy of Pathology, Atlanta, GA, Feb. 11-17, 2006.
• Berg JC, Lewis JT, Maloney SD, Vierkant RA, Hartmann LC, Visscher DW. Analysis of cancer risk in women with radial scars of the breast.

Podium Presentation at annual meeting of American Association for Cancer Research, April 1-5, 2006 in Washington, D.C.

• Pankratz VS, Vierkant RA, Maloney SD, Frost MH, Visscher DW, Hartman LC. Assessment of the Gail model in a cohort of women with atypical hyperplasia.

Poster Presentation at Joint statistical Meetings, Minneapolis, MN, August 10, 2006.
• Pankratz VS, Vierkant SD, Maloney SD, Hartmann LC. Epidemiologic comparisons of disease incidence among populations: The person-years approach.

• Ghosh K, Hartmann LC, Maloney D, Vierkant RA, Milanese TM, Visscher DW, Pankratz VS, Vachon CM. Mammographic breast density is inversely associated with age-related involution.

• Lingle W, Negron V, Bruzek A, Murphy L, Riehle D, Vierkant RA, Pankratz VS, Hartmann LC, Visscher DW. Centrosome amplification is greatest in benign breast lesions associated with an increase in risk of cancer.

• Boughey JC, Hartmann LC, Degnim AC, Vierkant RA, Ghosh K, Vachon CM, Maloney SD, Reynolds C, Pankratz VS. Assessment of the accuracy of the Gail model in women with atypical hyperplasia.

CONCLUSIONS
We have made significant progress on all three aims. Specifically, we have completed the cohort follow-up by questionnaires. Our pathologist has completed readings on the benign and cancer tissue for the entire cohort. We have evaluated the significance of the benign histologic categories (NP vs. PDWA vs. AH) and examined the risks associated with specific pathologic findings including atypia, papillomas, radial scars and involution. We calculated hazard functions for breast cancer by age at BBD and family history. We applied the Gail model to our study population of women with atypia and found the concordance between predicted individual risk and actual risk to be only slightly above chance. The Gail model significantly underestimated actual risk based on length of follow-up in our cohort of women with atypia. We have stained the test and atypia subgroups for several immunohistochemical markers. We have identified COX-2 as an important marker of an increased risk of breast cancer in women with atypia and are currently exploring ER, MIB-1 and p16. Additionally, we are working closely with Wayne State to characterize the histopathology and breast cancer outcomes in a cohort of African American women with benign breast disease.
Appendix A: Atypia Paper
Stratification of Breast Cancer Risk in Women With Atypia: A Mayo Cohort Study


A B S T R A C T

Purpose
Atypical hyperplasia is a well-recognized risk factor for breast cancer, conveying an approximately four-fold increased risk. Data regarding long-term absolute risk and factors for risk stratification are needed.

Patients and Methods
Women with atypical hyperplasia in the Mayo Benign Breast Disease Cohort were identified through pathway review. Subsequent breast cancers were identified via medical records and a questionnaire. Relative risks (RRs) were estimated using standardized incidence ratios, comparing the observed number of breast cancers with those expected based on Iowa Surveillance, Epidemiology, and End Results (SEER) data. Age, histologic factors, and family history were evaluated as risk modifiers. Plots of cumulative breast cancer incidence provided estimates of risk over time.

Results
With mean follow-up of 13.7 years, 66 breast cancers (19.9%) occurred among 331 women with atypia. RR of breast cancer with atypia was 3.88 (95% CI, 3.00 to 4.94). Marked elevations in risk were seen with multifocal atypia (eg, three or more foci with calcifications [RR, 10.35; 95% CI, 6.13 to 16.4]), RR was higher for younger women (< 40, RR, 6.76; 95% CI, 3.24 to 12.4). Risk was similar for atypical ductal and atypical lobular hyperplasia, and family history added no significant risk. Breast cancer risk remained elevated over 20 years, and the cumulative incidence approached 35% at 30 years.

Conclusion
Among women with atypical hyperplasia, multiple foci of atypia and the presence of histologic calcifications may indicate “very high risk” status (> 50% risk at 20 years). A positive family history does not further increase risk in women with atypia.

J Clin Oncol 25:2671-2677. © 2007 by American Society of Clinical Oncology

INTRODUCTION

Atypical hyperplasia is a well-established risk factor for subsequent breast cancer. Multiple studies corroborate an approximately four-fold increased risk of breast cancer in women undergoing surgical biopsy with a finding of atypia.1-7 Despite good concordance on the estimated relative risk (RR) with atypia, estimates of absolute risk with long-term follow-up are not well established. Reliable breast cancer risk estimates for women with atypia are crucial for risk-benefit analysis and decision making regarding risk-reduction strategies.

The Gail model in current use predicts a dramatically increased risk for those women who have both atypia and a family history (over that of atypia alone).8 Prior published literature has stated that the risk of breast cancer abates considerably after 10 years after a diagnosis of atypia,9 whereas more recent evidence indicates otherwise.10 It is also unclear whether breast cancer risk is higher in cases of atypical ductal hyperplasia (ADH) versus atypical lobular hyperplasia (ALH).

Here, we present a comprehensive description of breast cancer risk in women with atypical hyperplasia, based on 331 women with atypia in the Mayo Benign Breast Disease Cohort. Our investigation addresses the effect of family history on atypia risk, the effect of time since biopsy, the influence of ductal versus lobular histology, effects of age at atypia diagnosis, and presence of calcifications on breast cancer risk. In addition, we provide absolute risk estimates over time, and we also present a new histologic feature of atypia—multifocality—that stratifies breast cancer risk among women with atypia.
Study Population

Entry criteria for the study cohort have been previously described. Briefly, this comprises an institutional review board–approved study of women ages 18 to 85 years who had a benign breast biopsy via surgical excision during 1967 to 1991. The initial cohort included 9,887 women. With additional follow-up, data are now available for 9,976 women, 331 (3.5%) of whom had atypical hyperplasia.

Follow-Up

Follow-up for breast cancer events (including both invasive cancer and ductal carcinoma in situ [DCIS]) and risk-factor information were obtained for all women with atypia through the Mayo medical record and a study questionnaire. Family history was classified as negative, strong, or weak. The criteria for a strong family history were at least one first-degree relative with breast cancer before the age of 50 years or two or more relatives with breast cancer, with at least one being a first-degree relative. Any lesser degree of family history was considered weak.

Histology

All available archival hematoxylin and eosin–stained sections were evaluated by our breast pathologist (D.W.V.), without knowledge of the original histologic diagnoses or patient outcomes. The number of slides reviewed per case was variable because of the retrospective nature of the study, with a mean of 3.2 (standard deviation, 3.7). Calculations were recorded for each case when seen histologically. A diagnosis of ADH or ALH was based on the criteria of Page et al. ADH was characterized by filling and distension of involved ducts by an atypically complex proliferation of monotonous cells forming “punched out” (cuboidal form–like) secondary lumens or microcystic formations. Although well-developed examples of ADH share some morphologic features with low-grade DCIS, the latter is characterized by tumeformative growth (requiring complete involvement of >2 contiguous lumens) as well as greater nuclear enlargement and hyperchromatism. For each example of atypical hyperplasia, the number of separate foci of the atypia was defined. Multifocal atypia required its identification in more than one terminal duct lobular unit (TDLU) as defined by clear separation from another by nonneoplastic interlobular mammary stroma. All cases of multifocal atypia were agreed on by two study pathologists (D.W.V. and C.A.R.).

The primary study pathologist (D.W.V.) identified 332 cases of atypia from the entire benign breast disease cohort of 9,376. To address concerns of reproducibility in the diagnosis of atypia, we performed a nested study of concordance, blinding another pathologist (H.B.) to the study diagnoses in a random subset of several hundred samples from the original cohort, including nonproliferative lesions, proliferative disease without atypia, and atypical hyperplasia. Of 189 atypia samples reviewed for concordance, 165 (87.3%) atypias were similarly classified by subsequent independent review. Of the remaining 24 cases with differing interpretation, 18 were then judged to have atypia by joint review (D.W.V. and H.B.), and five of six remaining cases had atypia by review of a third “tiebreaker” breast pathologist (C.A.R.). The one case in question was excluded from further analysis, leaving a total of 331 subjects for study.

Statistical Analysis

Follow-up was defined as the number of days from benign biopsy to date of breast cancer diagnosis, death, or last contact. We estimated RRs with standardized incidence ratios (SIRs) and 95% CIs, dividing observed numbers of incident breast cancers by expected counts. We calculated expected counts by apportioning each individual’s follow-up time into 5-year age and calendar-period categories, and applying these person-years to population-based incidence rates, thereby accounting for differences in these variables. We used the Iowa Surveillance, Epidemiology, and End Results (SEER) registry as the reference population, because of its proximity to the Mayo Clinic catchment area and racial similarities compared with our cohort. We extrapolated incidence-rate data for cohort follow-up occurring outside the SEER time frames (1973–2002), such that person-years before 1973 were applied to 1973 to 1975 incidence rates, and person-years subsequent to 2002 were applied to 2001 to 2002 incidence rates. Assuming a two-sided test of hypothesis and a type I error rate of 0.05, we would have 80% power to detect SIRs as low as 0.61 if the expected event count is 2.5, as low as 2.97 if the count is 4.2, as low as 0.28 if the count is 0.5, and as low as 1.8 if the count is 17. Note that these expected counts reflect the approximate expected numbers of events in our cohort for women with three or more foci of atypia, two foci, one focus, and all subsets combined, respectively.

Recognizing that other biologic mechanisms may modify the association of atypia and breast cancer risk, we formally assessed the potential differential effects of these mechanisms using Poisson regression analyses. This approach allowed us to estimate SIRs with the flexibility that generalized linear models provide, such as covariate adjustment and tests for trend or heterogeneity across subgroups. For all analyses, the log-transformed expected event rate for each individual was modeled as the offset term.

We displayed observed and expected event rates using cumulative incidence curves and corresponding 95% confidence limits, accounting for the effects of death as a competing risk. Expected events were calculated for each 1-year follow-up interval in a manner similar to that used for determining SIRs. A modified Kaplan-Meier approach was used to cumulate expected incidence over these intervals. The expected curve was then smoothed using linear interpolation.

We compared the RR of ipsilateral versus contralateral breast cancer overall and across different medical characteristics using ratios of corresponding incidence rates. When calculating incidence for ipsilateral cancer, individuals with contralateral cancer were excluded at their diagnosis date, and vice versa. Women with missing laterality, or having bilateral biopsies or cancer, were excluded for both events. The RRs are equivalent to ratios of observed events, as the approach yields identical person-years for each event type. We thus used properties of the binomial distribution to obtain exact 95% CIs for these RRs. All statistical tests were postulated a priori and were two sided, and all analyses were conducted using the SAS software system (SAS Institute, Cary, NC).

Characteristics of Patients and Pathologic Specimens

A total of 331 women with atypia were identified in our cohort between 1967 and 1991. In Table 1, we present the patients’ vital status, breast cancer status, family history, age at biopsy, year of biopsy, indication for biopsy, and histologic features. Women were likely to be older than 55 at diagnosis of atypia (55.9%), and 42.9% had a family history of breast cancer (23.5% with a strong family history). Histologic findings included calcifications in most cases of atypia (68.6%); most cases (60.1%) had only one focus of atypia. The relative percentages of women with one, two, and three or more foci of atypia remained stable over the time period of the cohort. The proportions of women with ADH and ALH were similar.

Subsequent Breast Cancer Risk and Modifying Factors

The 331 women with atypia were followed for a total of 4,543 person-years (mean 13.7 years), with 66 (19.9%) observed breast cancer events to date. The histologic types are known in 61 of these, with 53 (86.9%) of 61 invasive cancers and eight (13.1%) of 61 DCIS. The majority of invasive cancers were ductal type (47 of 53, 89%), and the remaining six invasive lobular cancers were divided between the ALH and ADH subgroups. Table 2 shows the estimated RRs for breast cancer associated with various characteristics. The overall group with atypia demonstrates a four-fold RR of breast cancer (RR, 3.88; 95% CI, 3.00 to 4.94) compared with the general population.

Family History

There were no significant differences in RR seen among the subgroups with a strong family history (RR, 3.59; 95% CI, 1.96 to 2021) and no family history (RR, 2.50; 95% CI, 1.16 to 5.37). There was a trend toward a lower RR in those with strong versus weak family history (RR, 3.59; 95% CI, 1.96 to 2021 versus RR, 2.97; 95% CI, 1.30 to 6.75).

Information downloaded from jco.ascopubs.org and provided by MAYO CLINIC LIBRARY on July 19, 2007 from 129.176.151.6. Copyright © 2007 by the American Society of Clinical Oncology. All rights reserved.
Table 1. Clinical and Histologic Characteristics Among the 331 Women With Atypical Hyperplasia From the Mayo Benign Breast Disease Cohort Study

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>No.</th>
<th>%</th>
</tr>
</thead>
<tbody>
<tr>
<td>Vital status</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Deceased</td>
<td>101</td>
<td>30.6</td>
</tr>
<tr>
<td>Alive</td>
<td>230</td>
<td>69.5</td>
</tr>
<tr>
<td>Breast cancer status</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Noncase</td>
<td>265</td>
<td>80.1</td>
</tr>
<tr>
<td>Case</td>
<td>66</td>
<td>19.9</td>
</tr>
<tr>
<td>Age, years</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mean</td>
<td>50</td>
<td></td>
</tr>
<tr>
<td>SD</td>
<td>12</td>
<td></td>
</tr>
<tr>
<td>Family history of breast cancer</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Unknown</td>
<td>42</td>
<td></td>
</tr>
<tr>
<td>None</td>
<td>185</td>
<td>57.1</td>
</tr>
<tr>
<td>Weak</td>
<td>56</td>
<td>18.4</td>
</tr>
<tr>
<td>Strong</td>
<td>68</td>
<td>23.5</td>
</tr>
<tr>
<td>Age at BBD, years</td>
<td></td>
<td></td>
</tr>
<tr>
<td>&lt; 45 at BBD Dx</td>
<td>46</td>
<td>13.9</td>
</tr>
<tr>
<td>45-55 at BBD Dx</td>
<td>100</td>
<td>30.2</td>
</tr>
<tr>
<td>&gt; 55 at BBD Dx</td>
<td>185</td>
<td>55.9</td>
</tr>
<tr>
<td>Year of BBD</td>
<td></td>
<td></td>
</tr>
<tr>
<td>1967-1971</td>
<td>15</td>
<td>4.5</td>
</tr>
<tr>
<td>1972-1976</td>
<td>36</td>
<td>10.6</td>
</tr>
<tr>
<td>1977-1981</td>
<td>40</td>
<td>12.1</td>
</tr>
<tr>
<td>1982-1990</td>
<td>96</td>
<td>28.9</td>
</tr>
<tr>
<td>1991-1997</td>
<td>145</td>
<td>43.8</td>
</tr>
<tr>
<td>Indication for biopsy</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Unknown</td>
<td>6</td>
<td></td>
</tr>
<tr>
<td>Palpable mass</td>
<td>139</td>
<td>42.8</td>
</tr>
<tr>
<td>Mammographic abnormality</td>
<td>160</td>
<td>57.2</td>
</tr>
<tr>
<td>Calcifications</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Without calcifications</td>
<td>104</td>
<td>31.4</td>
</tr>
<tr>
<td>With calcifications</td>
<td>227</td>
<td>68.6</td>
</tr>
<tr>
<td>Histologic subtype</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Lobular</td>
<td>175</td>
<td>52.9</td>
</tr>
<tr>
<td>Ductal</td>
<td>142</td>
<td>42.9</td>
</tr>
<tr>
<td>Lobular and ductal</td>
<td>14</td>
<td>4.2</td>
</tr>
<tr>
<td>No. of foci of atypia</td>
<td></td>
<td></td>
</tr>
<tr>
<td>1</td>
<td>100</td>
<td>60.1</td>
</tr>
<tr>
<td>2</td>
<td>81</td>
<td>24.5</td>
</tr>
<tr>
<td>≥ 3</td>
<td>51</td>
<td>15.4</td>
</tr>
</tbody>
</table>

Abbreviations: SD, standard deviation; BBD, benign breast disease; Dx, diagnosis.

6.03), a weak family history (RR, 5.59; 95% CI, 3.20 to 9.90), or a negative family history (RR, 3.81; 95% CI, 2.60 to 5.37; Table 2; Fig 1A).

Age at Biopsy

Women with atypia diagnosed at younger age had a higher RR compared with age-matched expected rates (Table 2; Fig 1C). The RR was 6.76 at age less than 45, 5.10 at age 45 to 55, and 2.87 at age greater than 55 years (P for trend = .01). The increased risk seen in younger women was not due to a positive family history, because there was no difference in risk for women with and without a family history in each age subgroup (data not shown).

Number of Foci of Atypia

Increasing risk was seen with increasing foci of atypia: RR = 2.33 with a single focus, 5.26 for two foci, and 7.97 for three or more foci, with a highly significant test for trend (P < .001; Fig 1B). The increased risk seen with multiple foci of atypia was not due to predominance of young (higher risk) women in those subgroups; women younger than 45 years constituted only 4.94% and 7.84% of the subgroups with two and three foci of atypia, compared with 19.1% of the subgroup with one focus. Multivariate Poisson regression analysis also confirmed that young age and multifocality contributed independently to increased risk.

Calculations

Risk was dramatically increased in the small group of women (n = 38) with both calcifications and three or more foci of atypia (RR, 10.4; 95% CI, 6.13 to 16.4). However, women with calcifications and less than three foci of atypia (RR, 3.1) had risk similar to that of patients with fewer than three foci of atypia and no calcifications (RR, 3.31).

Histologic Type of Atypia

Histologic type of atypia did not affect breast cancer risk, because the RR of breast cancer was the same for ADH and ALH, although the few individuals with both histologic types may have higher risk (Fig 1D).

Indication for Biopsy

Breast cancer risk was similar whether a palpable or mammographic concern prompted the biopsy.

At-Risk Time Interval and Cumulative Incidence of Breast Cancer

The RR of breast cancer for the entire group with atypical hyperplasia was elevated persistently beyond 15 years, with a 20-year cumulative risk of 21% (95% CI, 14.0% to 28%) and a 25-year cumulative risk of 29% (95% CI, 20.6% to 37%; Fig 2). Stratification based on number of foci of atypia demonstrates a cumulative incidence of 18% for a single focus, 45% for two foci, and 48% for three or more foci of atypia at 25 years of follow-up (Fig 3).

Laterality of Breast Cancer Risk

Of the 66 women with atypia who subsequently developed breast cancer, side of cancer and side of atypia are known in 57 cases. Although cancer was more frequent in the ipsilateral breast, this difference was not statistically significant for the overall group with atypia (RR, 1.38 for ipsilateral v contralateral event; 95% CI, 0.79 to 2.21). However, the 32 women with atypia who developed breast cancer within 10 years of their benign biopsy were 2.2 times more likely (95% CI, 1.02 to 4.86; P = .05) to develop cancer in the same breast versus the opposite breast. Women with ADH had higher ipsilateral risk (RR, 1.50; 95% CI, 0.62 to 3.82), and women with three or more foci also had higher risk of ipsilateral breast cancer (RR, 2.20; 95% CI, 0.71 to 4.52), although these increases did not reach statistical significance due to small numbers of events and modest statistical power for these analyses. Women with ALH had similar cancer risk in both breasts (RR, 1.08; 95% CI, 0.45 to 2.14).

Having reliable breast cancer risk estimates for women with atypical hyperplasia is imperative in order to tailor their care appropriately. For women with atypia, the Gail model is the only model available for risk...
Table 2. Risk Factors for Breast Cancer Among the 331 Women With Atypia From the Mayo Benign Breast Disease Cohort Study

<table>
<thead>
<tr>
<th>Variable</th>
<th>No.</th>
<th>Person-Years</th>
<th>Observed Events</th>
<th>Expected Events</th>
<th>RR</th>
<th>95% CI</th>
</tr>
</thead>
<tbody>
<tr>
<td>Overall atypia group</td>
<td>331</td>
<td>4,543</td>
<td>66</td>
<td>17.0</td>
<td>3.88</td>
<td>3.00 to 4.84</td>
</tr>
<tr>
<td>Age at benign biopsy, years</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>&lt; 45</td>
<td>46</td>
<td>678</td>
<td>10</td>
<td>1.5</td>
<td>6.76</td>
<td>3.24 to 12.40</td>
</tr>
<tr>
<td>45-55</td>
<td>102</td>
<td>1,540</td>
<td>26</td>
<td>5.1</td>
<td>5.10</td>
<td>3.33 to 7.48</td>
</tr>
<tr>
<td>&gt; 55</td>
<td>185</td>
<td>2,328</td>
<td>10.4</td>
<td>2.87</td>
<td>1.94</td>
<td>1.0 to 4.10</td>
</tr>
<tr>
<td>No. of foci of atypia</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1</td>
<td>199</td>
<td>2,792</td>
<td>24</td>
<td>10.3</td>
<td>2.33</td>
<td>1.49 to 3.46</td>
</tr>
<tr>
<td>2</td>
<td>81</td>
<td>1,088</td>
<td>22</td>
<td>4.2</td>
<td>5.28</td>
<td>3.29 to 7.98</td>
</tr>
<tr>
<td>≥3</td>
<td>51</td>
<td>666</td>
<td>20</td>
<td>2.6</td>
<td>7.97</td>
<td>4.07 to 12.30</td>
</tr>
<tr>
<td>Calculations</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Without</td>
<td>104</td>
<td>1,520</td>
<td>18</td>
<td>5.6</td>
<td>3.21</td>
<td>1.90 to 5.08</td>
</tr>
<tr>
<td>With</td>
<td>227</td>
<td>3,013</td>
<td>48</td>
<td>11.4</td>
<td>4.21</td>
<td>3.10 to 5.58</td>
</tr>
<tr>
<td>&lt; 3 foci</td>
<td>189</td>
<td>2,530</td>
<td>30</td>
<td>9.7</td>
<td>3.10</td>
<td>2.09 to 4.43</td>
</tr>
<tr>
<td>≥ 3 foci</td>
<td>38</td>
<td>478</td>
<td>18</td>
<td>1.7</td>
<td>10.4</td>
<td>6.13 to 16.40</td>
</tr>
<tr>
<td>Histologic subtype</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Lobular</td>
<td>175</td>
<td>2,535</td>
<td>34</td>
<td>9.3</td>
<td>3.67</td>
<td>2.54 to 5.13</td>
</tr>
<tr>
<td>Ductal</td>
<td>142</td>
<td>1,815</td>
<td>27</td>
<td>7.0</td>
<td>3.83</td>
<td>2.83 to 5.58</td>
</tr>
<tr>
<td>Lobular and ductal</td>
<td>14</td>
<td>194</td>
<td>5</td>
<td>0.7</td>
<td>7.10</td>
<td>2.31 to 16.5</td>
</tr>
<tr>
<td>Family history of breast cancer</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>None</td>
<td>165</td>
<td>2,220</td>
<td>32</td>
<td>8.4</td>
<td>3.81</td>
<td>2.60 to 5.37</td>
</tr>
<tr>
<td>Weak</td>
<td>56</td>
<td>763</td>
<td>16</td>
<td>2.9</td>
<td>5.59</td>
<td>3.20 to 9.09</td>
</tr>
<tr>
<td>Strong</td>
<td>68</td>
<td>1,029</td>
<td>14</td>
<td>3.9</td>
<td>3.59</td>
<td>1.96 to 6.03</td>
</tr>
<tr>
<td>Indication for biopsy</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Palpable mass</td>
<td>139</td>
<td>2,069</td>
<td>33</td>
<td>7.2</td>
<td>4.55</td>
<td>3.13 to 6.39</td>
</tr>
<tr>
<td>Mammographic abnormality</td>
<td>188</td>
<td>2,409</td>
<td>32</td>
<td>9.5</td>
<td>3.36</td>
<td>2.30 to 4.74</td>
</tr>
</tbody>
</table>

Note. RR and CI represent standardized incidence ratio and 95% confidence limits, comparing observed number of events to those expected based on Iowa Surveillance, Epidemiology, and End Results data. All results account for the effects of age and calendar period. Abbreviation: RR, relative risks.

In this model, calculations of risk for women with atypia and a family history are dramatically higher, based on prior evidence from the Nashville study. Therefore, for a 50-year-old white woman with menarche at age 12, first birth at 24, and atypia on breast biopsy, the predicted lifetime risk of breast cancer is 17.5%. If that same woman also has a first-degree relative with...

Fig 1. Stratification of breast cancer risk for women with atypical hyperplasia. (A) Family history, 69 number of foci, 65 age at diagnosis, and (D) histologic type. Relative risks expressed as standardized incidence ratios and 95% confidence limits, comparing observed number of events to those expected based on Iowa Surveillance, Epidemiology, and End Results data. All results account for the effects of age and calendar period. Horizontal lines at 1.0 reflect the overall expected relative risk in the cohort. ALH, atypical lobular hyperplasia; ADH, atypical ductal hyperplasia.
breast cancer, her lifetime risk doubles to 34%. Our data indicate that the Gail model predicts inaccurately for such women because the increased risk of breast cancer associated with atypia is independent of the effect of family history.

Women in our cohort with atypia and a positive family history of breast cancer had no additional increased risk of breast cancer over that of atypia alone. This finding counters the commonly held view proposed by the Nashville study (ie, that atypia and a positive family history increase breast cancer risk additively). When data from other major studies of benign breast disease are considered along with the Mayo findings, the preponderance of evidence calls into question the result from the Nashville group. In that study, the subgroup of women with atypia and a family history was small (n = 39) with an RR of 8.9 (95% CI, 4.8 to 17), compared with 3.5 (95% CI, 2.3 to 5.5) in 193 women with atypia and no family history.2 In contrast, evaluation of a much larger population in the Breast Cancer Detection and Demonstration Project showed similar frequencies of breast cancer in women with atypia and family history (16 of 261, 6.1%) compared with those with atypia alone (51 of 1,044, 4.9%).3 Recent data from the Nurses’ Health Study confirm our finding that a family history of breast cancer in a first-degree relative does not further increase risk among women with atypical hyperplasia.13 To explain these findings, we postulate that atypical hyperplasia is a phenotype reflecting increased risk; this phenotype derives from both inherited risk and lifetime exposures. Thus, the histologic presence of atypia already reflects the increased breast cancer risk inherent in a positive family history.

We have identified a new histologic variable that appears to stratify risk in women with atypia: multifocality. The RR of breast cancer increases in a dose-response fashion for women with one, two, and three or more foci of atypia, with a statistically significant test for trend. With a single focus, the cumulative incidence of breast cancer reached 18% at 25 years. For women with two or more foci of atypia, the cumulative risk of breast cancer was greater than 40% at 25 years. Moreover, in the highest risk subgroup of women with three or more foci and histologic calcifications, the cumulative incidence exceeded 50% over 25 years. This level of risk approaches that reported for carriers of BRCA mutations.16 In line with our observation, differential risk based on extent of disease has been established for lobular neoplasia (ie, ALH or lobular carcinoma),17 and the number of foci of atypia found in core needle biopsy specimens correlates with the likelihood of finding cancer at surgical excision.18

Some may question whether multifocal atypias may actually represent subtle in situ carcinoma, particularly those of the ADH type. In cases of multifocal ADH, it should be emphasized that individual foci arose in separate and distinct terminal duct lobular units, none of which measured more than 2 mm. Hence, these examples failed to exhibit the confluent degree of cellular proliferation requisite for a diagnosis of DCIS. We submit that more widespread distribution of atypical foci within breast tissue signals a larger burden of at-risk tissue that has progressed along the continuum toward breast cancer. The data presented in this article provide evidence that the extent of pre-malignant breast change is related to subsequent cancer risk. Since this is the first report of the clinical relevance of this histologic finding, we recognize the need for validation and plan to evaluate this factor in a more recent cohort from our institution. Furthermore, we hope that other research groups with large numbers of patients with atypical hyperplasia will also examine the relevance of multifocal atypia in their study sets.

Age at the diagnosis of atypia also emerged as a significant modifier of subsequent breast cancer risk, with a higher RR in younger women. The Nurses Health Study16 and the Breast Cancer Detection and Demonstration Project1 have also shown higher risk in younger women with atypia. In our cohort, this increased risk in younger women is not explained by more frequent multifocal disease or a positive family history. Perhaps atypical hyperplasia present at a young age is the result of previous oncogenic events; alternatively, breast tissue with atypia may be unusually susceptible to proposed oncogenic estrogen metabolites associated with the premenopausal hormonal environment.19

When counseling women with atypical hyperplasia, the length of time at risk is a key element in planning risk-reduction strategies. Dupont and Page10 reported that the greatest risk of breast cancer after
a diagnosis of atypia lies in the first 10 years, with subsequent RR reduced by half \( (P = .06) \). By contrast, the Nurses Health Study found that risk does not decrease over time, with RR slightly higher more than 10 years after biopsy (RR, 3.6) compared with the first 10 years (RR, 3.2). Our data confirm that the RR for breast cancer after a biopsy demonstrating atypia remains significantly elevated for at least 15 years.

Data on long-term absolute risk are more useful than RR estimates when counseling patients. Our study provides estimates of absolute risk for women with atypia and indicates a higher cumulative incidence of breast cancer with long-term follow-up than has been reported by other studies. Figures from the study of Dupont and Page show a cumulative breast cancer incidence of 13% at 20 years and 23% at 25 years in women with atypia. The cumulative incidences identified in our cohort were higher: 21% at 20 years and 29% at 25 years. One factor contributing to this difference is our inclusion of DCIS as a recordable breast cancer event, whereas the Nashville study counted only cases of invasive breast cancer. Because DCIS currently receives local treatment (and in some cases, systemic treatment) similar to that for early-stage invasive breast cancer, it is reasonable to include cases of DCIS when estimating risk.

Our data on the laterality of subsequent breast cancer do not allow conclusions regarding atypical hyperplasia acting as a precursor lesion, yet there is a suggestion of predilection for the ipsilateral breast that requires ongoing study. Breast cancers occurring in the first 10 years after atypia diagnosis were significantly more likely to occur in the ipsilateral breast. A recent study of gene expression profiling identified remarkably similar alterations in gene expression among ADH, DCIS, and invasive cancers found in the same specimen, supporting the role of atypical hyperplasia as a precursor lesion. Regarding differences in ipsilateral risk for ductal versus lobular atypia, we found that risk was equal for both breasts after a diagnosis of ALH, which is consistent with the distribution of invasive breast cancers after a diagnosis of lobular carcinoma in situ. In contrast, ADH was more likely associated with a later ipsilateral breast cancer, as has been shown for DCIS untreated after diagnostic biopsy.

In conclusion, our study provides a comprehensive analysis of breast cancer risk associated with atypical hyperplasia. These findings confirm a four-fold RR of subsequent breast cancer in women with atypical hyperplasia. We estimate that the long-term absolute risk of subsequent breast cancer (in situ or invasive) is higher than previously reported—at least 25% at 25 years, and as high as 50% to 60% in a high-risk subgroup defined by multifocality and calcifications. A positive family history does not confer significantly increased risk in women with atypia. Improved risk prediction and stratification is now possible to guide risk reduction counseling for women with atypical hyperplasia.

The author(s) indicated no potential conflicts of interest.

---

**Authors' Contributions**

Conception and design: Amy C. Degnim, Daniel W. Visscher, Marlene H. Frost, Thomas A. Sellers, Robert A. Vierkant, Kartihi Gosh, Thera Tlsty, L. Joseph Melton III, Lynn C. Hartmann

Financial support: Lynn C. Hartmann

Administrative support: Lynn C. Hartmann

Provision of study materials or patients: Wilma L. Lingle, L. Joseph Melton III, Lynn C. Hartmann


Breast Cancer Risk in Women With Atypia


18. Sreed, M., Gurseman, S.J., Higginbotham, S., et al.: Formation of the deprotonated NGlycosine and 7-guanine adducts by reaction of DNA with hexestrol-3',4'-quinone or enzyme-activated 3'-hydroxyhexestrol. Implications for a unifying mech-
anism of tumor initiation by natural and synthetic estrogens. Steroids 70:37-45, 2005


Acknowledgment

We thank Joel Worra for database development; Teresa Allens, Jo Johnson, Mary Amundsen, Mary Campion, and Romayne Thompson for data collection; and Ann Harris and the Survey Research Center for patient follow-up.
Appendix B: Involution Paper
Age-Related Lobular Involution and Risk of Breast Cancer


Background: As women age, the lobules in their breasts undergo involution or regression. We investigated whether lobular involution in women with benign breast disease was associated with subsequent breast cancer risk. Methods: We examined biopsy specimens of 8736 women in the Mayo Benign Breast Disease Cohort from whom biopsy samples were taken between January 1, 1967, and December 31, 1991. Median follow-up for breast cancer outcomes was 17 years. We classified lobular involution in the background breast tissue as none (0% involuted lobules), partial (1%-75%), or complete (>75%). Subsequent breast cancer events and data on other risk factors were obtained from medical records and follow-up questionnaires. To estimate relative risks (RRs), standardized incidence ratios were calculated by use of incidence rates from the Iowa Surveillance, Epidemiology, and End Results (SEER) Registry. All statistical tests were two-sided. Results: Distribution of extent of involution was none among 1627 (18.6%) women, partial among 5197 (59.5%), and complete among 1912 (21.9%). Increased involution was positively associated with increased age and inversely associated with parity (both P<.001). The relative risk for the entire cohort of 8736 women, compared with the Iowa SEER population, was 1.40 (95% CI = 1.30 to 1.51). Risk of breast cancer was associated with the extent of involution (for no involution, RR [i.e., observed versus expected] = 1.00, 95% confidence interval [CI] = 0.90 to 1.10; for partial involution, RR = 1.00, 95% CI = 1.00 to 1.10; and for complete involution, RR = 0.99, 95% CI = 0.96 to 1.02; test for heterogeneity P=.001). Lobular involution modified risk in all subsets (e.g., among women with atypia, for no involution, RR = 0.79, 95% CI = 0.56 to 1.13; for partial involution, RR = 0.92, 95% CI = 0.58 to 1.43; and for complete involution, RR = 0.79, 95% CI = 0.41 to 1.47; P < .001). Conclusions: In this large cohort of women with benign breast disease, lobular involution was associated with reduced risk of breast cancer. Abrupt involution may be a biologically important phenomenon in breast cancer biology. [J Natl Cancer Inst 2006;98:1600-7]
Histology

All slides were reviewed by a breast pathologist (DWV) without knowledge of patient age, cancer outcome, or original histologic diagnosis. Biopsy punchings were classified by the most extreme degree of hyperplasia as nonproliferative, proliferative disease without atypia, or atypical hyperplasia, as previously reported (9).

Each biopsy specimen was also categorized according to the extent of lobular involution in the background breast tissue. Involved terminal duct lobular units (TDLUs) contain only a few to several small acini that may be distorted by cystic change (Fig. 1). Involved lobules also have flattened inconspicuous acinar epithelium with atypia of specialized intralobular stroma. The degree of involution for each specimen was categorized as none (0% TDLUs involved), partial (1%–74% TDLUs involved), or complete (≥75% TDLUs involved). These cut points were set by the pathologists at the initiation of the study to best distinguish the extremes of no involution from near-complete involution.

In general, viewing 5 to 10 lobules was sufficient to assess the extent of involution. One slide from a breast specimen typically contained a dozen or more lobules. There are two exceptions to this statement: 1) when involution was extensive and there are only a few lobular remnants on the slide (which is sufficient to state that complete lobular involution has occurred) and 2) when the entire sample consists of an epithelial hyperplastic lesion, as was the case for 640 (6.8%) of the 9376 women in our original cohort.

Risk Factor Information and Follow-up

To obtain information about family history, reproductive history, and use of hormone replacement therapy, a study-specific questionnaire was sent to patients: 5352 (61.7%) of the 8736 women or their next of kin returned the questionnaire. Follow-up for breast cancer events was obtained through comprehensive (inpatient and outpatient) Mayo medical records and the questionnaire. Family history of breast cancer was categorized as strong, weak, or negative. A strong family history was defined as the patient having 1) at least one first-degree relative with breast cancer diagnosed before age 50 years or 2) two relatives with breast cancer at any age, with at least one being a first-degree relative. Patients with family history of breast cancer who did not meet the above criteria were categorized as having a weak family history (5).

Statistical Analysis

Data were summarized descriptively by use of frequencies and percentages. We initially compared the unadjusted distribution of breast cancer risk factors across levels of involvement with chi-square tests of statistical significance. Subsequent comparisons were made after accounting for the effects of age by use of multivariate, nominal logistic regression analysis (9). We summarized results from these analyses by use of adjusted percentiles, carried out by calculating log odds estimates for each 10-year age category (<40, 40–49, 50–59, 60–69, 70–79, or ≥80 years), back-transforming to percent estimates, and then averaging the corresponding percentiles across all sets of age. This approach was similar to a least-squares means estimate in an analysis of variance setting. Among 245 women with synchronous bilateral biopsy examinations, we assessed the level of agreement across the two readings by use of weighted kappa statistics and their corresponding 95% confidence intervals (CI).

Participants and Methods

Study Population

Inclusion and exclusion criteria for the study cohort have been previously described (8). Briefly, the study population consisted of women aged 18–85 years who had benign breast disease (i.e., a breast biopsy examination with benign lesion) diagnosed via surgical excision at Mayo Clinic between January 1, 1967, and December 31, 1991. The Mayo Benign Breast Disease Cohort included 9087 women with 15 years of follow-up at initial report (8). Since that report, we obtained cancer follow-up data for an additional 289 women, for whom that information was lacking previously, bringing the cohort to a total of 9376 women with a median of 17 years of follow-up. For 640 breast biopsy samples, the biopsy specimen consisted entirely of the index lesion; there was no background breast tissue in which to determine the degree of lobular involution. Thus, the final cohort for this analysis included 8736 women.

All protocol procedures and patient contact materials were reviewed and approved by the Institutional Review Board of the Mayo Clinic. Return of the patient contact materials was considered implied consent.

Journal of the National Cancer Institute, Vol. 98, No. 22, November 15, 2006

ARTICLES 1601
The length of follow-up for each woman in the study was calculated as the number of days from her biopsy examination to the date of her breast cancer diagnosis, death, or last contact. We estimated relative risks (RRs) on the basis of standardized incidence ratios by dividing the observed numbers of incident breast cancers by expected numbers of population-based incident breast cancers. Expected values were calculated by apportioning each woman's person-years of follow-up into 5-year age and calendar-period categories and multiplying these by the corresponding breast cancer incidence rates from the Iowa Surveillance, Epidemiology, and End Results (SEER) Registry. This reference population was chosen because of its demographic similarities to the Mayo Clinic population (80% of cohort members reside in the upper Midwest). Potential heterogeneity in relative risks across levels of involution was assessed by use of Poisson regression analysis, with the log-transformed expected event rate for each individual modeled as the offset term.

In addition to assessing overall breast cancer risk, we also compared rates of ipsilateral to contralateral breast cancer in relation to the side of the biopsy lesion, both overall and by levels of involution. When calculating incidence for ipsilateral cancer, individuals with contralateral cancer were censored at their date of diagnosis, and vice versa. Women with missing latency information, bilateral biopsy examination results, or bilateral breast cancer were censored for both events in these analyses. This approach yielded identical numbers of person-years for each type of event. As a result, the length of follow-up was not a factor in the analysis, and the rate comparisons reduced to simple comparisons of the number of events. Thus, we were able to assess whether the relative rate of ipsilateral cancer (compared with contralateral cancer) differed across levels of involution using simple chi-square tests of statistical significance. All statistical tests were two-sided, and all analyses were carried out with the SAS software system (SAS Institute, Inc., Cary, NC).

RESULTS

Extent of Lobular Involution

We characterized the extent of lobular involution in the benign breast biopsies of a cohort of 8756 women with tissue sampled between January 1, 1967, and December 31, 1991, at the Mayo Clinic. The distribution of the patients by level of lobular involution was as follows: no involution among 1627 (18.6%) women, partial involution among 5197 (59.5%) women, and complete involution among 1912 (21.9%) women.

Factors Associated With Involution

As shown in Table 1, the degree of lobular involution increased progressively with age at diagnosis of benign breast disease (P<0.001). Complete involution of lobular units was observed in only 19.3% (154) of the 566 women who were younger than 30 years at their benign biopsy; in 55 (5.1%) of the 1037 women aged 30-39 years; in 142 (5.8%) of the 2446 women aged 40-49 years; in 455 (21.6%) of the 2109 women aged 50-59 years; in 724 (45.3%) of the 1600 women aged 60-69 years; and in 519 (53.1%) of the 978 women aged 70 years or older. The gradual nature of the involution process is apparent in that it is already present at least to a partial degree in more than half of the women younger than 40 years and is still ongoing in women older than 70 years.

Table 1. Association of lobular involution with age at diagnosis of benign breast disease, family history, type of hormone replacement therapy,

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>None</th>
<th>Partial</th>
<th>Complete</th>
<th>P-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Overall</td>
<td>1627 (18.6)</td>
<td>5197 (59.5)</td>
<td>1912 (21.9)</td>
<td>&lt;0.01</td>
</tr>
<tr>
<td>Age at BBD, y</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>15-29</td>
<td>308 (54.4)</td>
<td>239 (42.2)</td>
<td>19 (2.6)</td>
<td>0.47</td>
</tr>
<tr>
<td>30-39</td>
<td>417 (40.2)</td>
<td>507 (47.7)</td>
<td>35 (3.1)</td>
<td>0.32</td>
</tr>
<tr>
<td>40-49</td>
<td>643 (28.3)</td>
<td>1603 (67.9)</td>
<td>142 (5.8)</td>
<td>0.36</td>
</tr>
<tr>
<td>50-59</td>
<td>218 (10.5)</td>
<td>1438 (68.1)</td>
<td>455 (21.6)</td>
<td>0.27</td>
</tr>
<tr>
<td>60-69</td>
<td>291 (3.1)</td>
<td>847 (52.3)</td>
<td>724 (45.3)</td>
<td>0.42</td>
</tr>
<tr>
<td>&gt;70</td>
<td>12 (1.3)</td>
<td>447 (45.7)</td>
<td>519 (53.1)</td>
<td>0.001</td>
</tr>
<tr>
<td>Family history of breast cancer</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>None or weak</td>
<td>796 (21.1)</td>
<td>2717 (56.4)</td>
<td>911 (22.4)</td>
<td>&lt;0.01</td>
</tr>
<tr>
<td>Strong</td>
<td>223 (24.5)</td>
<td>556 (53.9)</td>
<td>159 (17.5)</td>
<td>0.26</td>
</tr>
<tr>
<td>Parity</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Nulliparous</td>
<td>113 (17.6)</td>
<td>421 (55.3)</td>
<td>177 (27.1)</td>
<td>0.90</td>
</tr>
<tr>
<td>Parous</td>
<td>893 (22.6)</td>
<td>2780 (56.2)</td>
<td>856 (21.2)</td>
<td>0.52</td>
</tr>
<tr>
<td>No. of children</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>0</td>
<td>113 (17.6)</td>
<td>431 (59.3)</td>
<td>177 (27.1)</td>
<td>0.87</td>
</tr>
<tr>
<td>1</td>
<td>72 (7.2)</td>
<td>299 (69.7)</td>
<td>131 (23.0)</td>
<td>0.29</td>
</tr>
<tr>
<td>2</td>
<td>224 (22.2)</td>
<td>891 (85.2)</td>
<td>238 (21.5)</td>
<td>0.91</td>
</tr>
<tr>
<td>3</td>
<td>257 (23.9)</td>
<td>742 (58.5)</td>
<td>205 (17.6)</td>
<td>0.62</td>
</tr>
<tr>
<td>&gt;4</td>
<td>235 (16.6)</td>
<td>871 (55.3)</td>
<td>242 (16.3)</td>
<td>0.37</td>
</tr>
<tr>
<td>Children breast fed</td>
<td></td>
<td></td>
<td></td>
<td>&lt;0.01</td>
</tr>
<tr>
<td>No</td>
<td>464 (21.9)</td>
<td>1555 (57.4)</td>
<td>426 (21.2)</td>
<td>0.42</td>
</tr>
<tr>
<td>Yes</td>
<td>431 (22.3)</td>
<td>1302 (55.9)</td>
<td>364 (40.8)</td>
<td>0.56</td>
</tr>
<tr>
<td>Hormone replacement therapy</td>
<td></td>
<td></td>
<td></td>
<td>0.16</td>
</tr>
<tr>
<td>Ever</td>
<td>481 (22.3)</td>
<td>1300 (57.1)</td>
<td>494 (20.9)</td>
<td>0.20</td>
</tr>
<tr>
<td>Never</td>
<td>516 (21.8)</td>
<td>1837 (55.4)</td>
<td>456 (23.9)</td>
<td>0.52</td>
</tr>
</tbody>
</table>

*For age at diagnosis of benign breast disease (BBD), percentage values were unadjusted. For all other variables, percentage values were adjusted for age.

We also found a strong, inverse association (P<0.001) between lobular involution and parity (Table 1). Specifically, the likelihood of complete involution was 27.1% (95% CI = 24.1% to 30.1%) in multiparous women, 28.0% (95% CI = 24.7% to 31.4%) in women who had one child, 21.5% (95% CI = 19.2% to 23.8%) in women who had two children, 20.4% (95% CI = 17.8% to 23.0%) in women who had three children, and 18.0% (95% CI = 16.1% to 20.0%) in women who had four children or more. Separating women into categories of ever versus never breastfeeding did not reveal any relationship with extent of lobular involution (P = 0.428). Women who reported having used hormone replacement therapy were slightly less likely to have complete involution (20.3%) than those with no history of hormone replacement therapy use (22.9%) (P = 0.16). Breast tissue from women with a strong family history of breast cancer was less likely than that from women with no or a weak family history of breast cancer to demonstrate lobular involution, i.e., after adjustment for age, more women with a strong family history had no involution (24.5%) than those with no or a weak family history.
Table 2. Association of involution and other risk factors with breast cancer after the diagnosis of benign breast disease

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>No. of women</th>
<th>No. of person-years</th>
<th>No. of observed events</th>
<th>No. of expected events*</th>
<th>Relative risk (95% CI)**</th>
</tr>
</thead>
<tbody>
<tr>
<td>Degree of involution</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>None (0%)</td>
<td>1627</td>
<td>32,721</td>
<td>150</td>
<td>79.6</td>
<td>1.88 (1.19 to 2.21)</td>
</tr>
<tr>
<td>Partial (1%-35%)</td>
<td>5191</td>
<td>90,469</td>
<td>440</td>
<td>300.1</td>
<td>1.47 (1.23 to 1.61)</td>
</tr>
<tr>
<td>Complete (35%)</td>
<td>1912</td>
<td>28,376</td>
<td>106</td>
<td>116.5</td>
<td>0.91 (0.75 to 1.10)</td>
</tr>
<tr>
<td>Histologic type</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Nonproliferative</td>
<td>5736</td>
<td>101,201</td>
<td>355</td>
<td>321.5</td>
<td>1.00 (0.59 to 1.23)</td>
</tr>
<tr>
<td>Proliferative without atypia</td>
<td>2077</td>
<td>45,418</td>
<td>276</td>
<td>158.1</td>
<td>1.75 (1.35 to 1.86)</td>
</tr>
<tr>
<td>Proliferative with atypia</td>
<td>532</td>
<td>4456</td>
<td>65</td>
<td>16.6</td>
<td>3.96 (3.82 to 4.08)</td>
</tr>
<tr>
<td>Age at biopsy, y</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>&lt;45</td>
<td>2682</td>
<td>52,955</td>
<td>158</td>
<td>108.8</td>
<td>1.45 (1.24 to 1.70)</td>
</tr>
<tr>
<td>45-55</td>
<td>2559</td>
<td>49,246</td>
<td>254</td>
<td>169.0</td>
<td>1.50 (1.32 to 1.70)</td>
</tr>
<tr>
<td>&gt;55</td>
<td>3485</td>
<td>49,734</td>
<td>286</td>
<td>218.8</td>
<td>1.59 (1.53 to 1.64)</td>
</tr>
<tr>
<td>Family history of breast cancer*</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>None or weak</td>
<td>4624</td>
<td>81,514</td>
<td>329</td>
<td>269.4</td>
<td>1.22 (1.09 to 1.36)</td>
</tr>
<tr>
<td>Strong</td>
<td>928</td>
<td>18,385</td>
<td>145</td>
<td>59.3</td>
<td>1.03 (0.99 to 1.25)</td>
</tr>
<tr>
<td>Age at birth of first live child, y</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Nulliparous</td>
<td>711</td>
<td>13,021</td>
<td>71</td>
<td>51.0</td>
<td>1.23 (1.20 to 2.00)</td>
</tr>
<tr>
<td>&lt;20</td>
<td>4121</td>
<td>77,710</td>
<td>327</td>
<td>257.9</td>
<td>1.07 (1.03 to 1.14)</td>
</tr>
<tr>
<td>&gt;20</td>
<td>388</td>
<td>7691</td>
<td>30</td>
<td>22.4</td>
<td>1.30 (0.50 to 0.92)</td>
</tr>
</tbody>
</table>

*Number of events expected on the basis of Iowa Surveillance, Epidemiology, and End Results breast cancer incidence data.
**All analyses control for the effects of age and calendar period. CI = confidence interval.

Risk ratio at parity was available for 2129 of 3736 women.

(21.1%), and fewer women with a strong family history had complete involution (19.5%) than those with no or a weak family history (22.4%). Logistic regression analysis comparing distribution of involution across levels of family history, P < 0.001.

We also examined the extent of lobular involution by category of benign breast disease. Among women with nonproliferative disease, 27.2% had complete involution. However, among women with proliferative disease without or with atypia, only 11.5% had complete lobular involution (P < 0.001).

Lobular Involution and Breast Cancer Risk

This cohort of women with benign breast disease was, overall, at increased risk of breast cancer when compared with age-matched women in the general population. Specifically, the relative risk for the entire cohort of 3736 women, compared with the Iowa SEER population, was 1.40 (95% CI = 1.30 to 1.51). In our cohort, degree of involution was associated inversely with breast cancer risk (Table 2; e.g., for no involution, RR = 1.88, 95% CI = 1.59 to 2.21; for partial involution, RR = 1.47, 95% CI = 1.33 to 1.61; and for complete involution, RR = 0.91, 95% CI = 0.75 to 1.07; test for heterogeneity P < 0.001).

Figure 2 illustrates the observed associations between the extent of involution and breast cancer risk among strata of age, histology, family history, and parity. Extent of lobular involution modified age-related breast cancer risk (e.g., for a woman older than 55 years with no involution, RR = 3.21, 95% CI = 1.90 to 5.08, and for a similar woman with complete lobular involution, RR = 0.92, 95% CI = 0.74 to 1.14). The same pattern was observed in all age groups.

Similarly, progressive increases in lobular involution in background breast tissue was associated with reduced risk of breast cancer among women with benign proliferative disease, even those with atypia (Fig. 2). Among women with atypia, no involution was more strongly associated with a higher risk of breast cancer (RR = 7.79, 95% CI = 3.56 to 14.81) than complete involution (RR = 1.49, 95% CI = 0.41 to 3.82) or partial involution (RR = 4.06, 95% CI = 3.03 to 5.33) (test for heterogeneity, P = 0.003).

Among women with proliferative disease without atypia, no involution was associated with a higher risk of breast cancer (RR = 2.94, 95% CI = 2.26 to 3.78) than complete involution (RR = 1.11, 95% CI = 0.68 to 1.71). The same pattern held true for those with nonproliferative breast disease; i.e., those with no invasion had a higher risk than those with complete or partial involution.

Lobular involution modified the influence of family history on risk of breast cancer (Fig. 2). Among women with a strong family history of breast cancer, no invasion was associated with the highest risk of breast cancer (RR = 2.77, 95% CI = 1.94 to 3.84), followed by partial invasion (RR = 2.72, 95% CI = 1.92 to 2.20) and then by complete involution (RR = 1.61, 95% CI = 0.92 to 2.61). Among women with no or a weak family history of breast cancer and complete involution, the risk of breast cancer (RR = 0.59, 95% CI = 0.41 to 0.81) was approximately half of that for the general population, which was based on Iowa SEER data, and approximately 50-fold less than the risk of those with strong family history and no invasion (i.e., RR = 2.77, 95% CI = 1.94 to 3.84).

Lobular involution also modified the risk associated with parity or age at birth of the first live child. Among nulliparous women and women whose first live child was born when she was at least 30 years old, no lobular involution was associated with increased risks of breast cancer (RR = 0.16, 95% CI = 0.65 to 2.41, and RR = 1.74, 95% CI = 1.31 to 2.30, respectively). However, among those same two groups of women, when complete lobular involution had occurred, there was no increase in risk (RR = 1.02, 95% CI = 0.53 to 1.78, and RR = 0.48, 95% CI = 0.10 to 1.40, respectively). Among women whose first live child was born when she was younger than 30 years old, complete lobular involution was associated with a decreased risk of breast cancer (RR = 0.85, 95% CI = 0.44 to 0.91).

We also asked whether the extent of involution in which the biopsy examination was performed—nominally, before or after widespread adoption of mammography—affected the results. In the first 15 years of the cohort (from 1967 through 1981), 78% of biopsy examinations were done because of a palpable concern (i.e., a palpable...
lump detected during a clinical breast examination or by the patient), and 22% were done because of an abnormal mammo-gram. From 1982 through 1991, 40% of the biopsy examinations were done because of a palpable concern, and 50% were done because of an abnormal mammogram. The relative risks of breast cancer by involution status and by dates (Table 3) indicated that associations between extent of involution and risk were similar in the pre- and postmammography time periods.

Lobular Involvement: Localized or Field Effect

To address whether or not the degree of involution was relevant only to the area of the biopsy or was representative of the field of breast tissue, we examined 1) whether, for women with bilateral benign biopsy examination results, involution results were concordant and 2) whether the degree of involution at the benign biopsy site was associated with the risk of ipsilateral breast cancer or with both ipsi- and contralateral breast cancers. A subset of 245 women had bilateral biopsy examinations performed at the same time. In 203 (83%) of these women, the same category of involution (no, partial, or complete) was found in the biopsy tissue from both breasts. In 41 (17%), there was a difference of one category between the two breasts. Only one individual had complete involution in the biopsy tissue of one breast and no involution in the contralateral sample. These results indicate a high level of agreement in involution measured across multiple biopsy specimens within a woman (kappa coefficient = 0.72, 95% CI = 0.64 to 0.80; test for agreement beyond due expected by chance P < 0.01). We next investigated the extent of involution and the laterality of subsequent breast cancers. In our cohort overall, there is a slight predominance of ipsilateral breast cancers (55.5%) over contralateral breast cancers (44.5%), as reported previously (8); this result is thought to reflect the presence of some direct precursors among these lesions. To determine whether involution at the site of the benign breast disease was relevant to the contralateral breast, we examined the ratio of ipsilateral to contralateral events by degree of involution. With no involution, the ratio was 55.6% ipsilateral to 46.4% contralateral; for partial involution, the ratio was 55.9% to 44.1%; and for complete involution, the ratio was 53.5% to 46.5% (chi-square test for difference in percent ipsilateral across involution status; P = .05). Thus, the relationship between involution extent and breast cancer risk was observed in both the ipsi- and contralateral breast.

DISCUSSION

We characterized the degree of lobular involution in the background breast tissue in a large cohort of women with benign...
breast disease. Our data demonstrate a strong, inverse relationship between degree of involution and breast cancer risk. To our knowledge, this is the first study to systematically examine age-related involution in the context of breast cancer risk. Furthermore, greater degrees of involution reduced breast cancer risk even in high-risk subsets defined by age, atypical reproductive history, or family history. There was a strong direct association between involution and increasing age. There was an inverse association between involution and parity.

As in this study, others have found that older women tend to have fewer lobules or only lobule remnants (4,5). Cowan and Herbert (4) performed a detailed autopsy study of the breast tissue from 102 women, aged 50–104 years, who died without known breast disease. Although considerable individual variability was present, they described a progressive loss of lobules with increasing age. Earlier reports state that age-related involution has already begun in women under the age of 40 years (1,5). Our data confirm that this process is present, at least to a partial degree, in many younger women.

We hypothesize that the degree of involution detected at the benign biopsy site reflects that of the overall field of a woman’s breast tissue. We believe that this hypothesis is reasonable because of our results showing a similar likelihood of contralateral and ipsilateral breast cancers by involution status at the site of the benign breast disease and because of the high concordance in involution status in women who had bilateral biopsy examinations. However, our study design cannot answer this question definitively. To do so would require examination of the extent of involution throughout all of a woman’s breast tissue.

It is widely appreciated that, as women age, their risk of breast cancer increases. But the role of involution is not as clear. The degree of involution can be assessed by a variety of methods, including qualitative analysis of breast tissue samples, quantitative analysis of breast tissue volume, and radiographic examination of breast density. However, these methods are not always accurate and may not provide a comprehensive picture of the extent of involution.

We examined various factors besides age for their association with degree of involution. We found an inverse association between lobular involution and parity. Others have also reported that the more children a woman has, the more likely she is to have persistence of lobular structures (1,5), which we found was associated with increased risk of breast cancer. Yet, multiparity is generally considered to reduce the risk of breast cancer (13,44).

Several factors may explain this apparent contradiction. First, we do not have data on the age at each child’s birth for the women in our cohort. Some epidemiologic work has suggested that full-term pregnancies after 35 years of age are associated with an increased risk of breast cancer (7,15). Thus, data on a woman’s age at each pregnancy and on her age at breast biopsy examination would help to evaluate more definitively the relationships of parity, involution, and breast cancer risk. In addition, our study was limited by the relatively large size of the group of women categorized as having partial involution. More specific, quantitative measures of degree of involution should be explored to determine whether the association between parity status and degree of involution can be defined more precisely. Given the inverse association between complete involution and multiparity and given that both are associated with reduced risk of breast cancer, we hypothesize that the breast cancer risk modification associated with parity is independent of involution status.

There are several biologic mechanisms by which involution or lack thereof could alter a patient’s breast cancer risk. The decrease in epithelial cell number that accompanies involution may decrease breast cancer risk simply because there are fewer epithelial cells to undergo malignant transformation. Another possibility is that aberrant involution may be a marker or phenotype reflecting underlying constitutional susceptibility for breast cancer that is present in the epithelial or stromal compartment or in their relationship with each other. Yet another possibility is that failure to undergo timely or appropriate involution allows prolonged exposure of epithelial cells to intrinsic and/or extrinsic mutagenic stresses (16–20). In this model, the prime targets of such mutagenic processes, such as stem cells or early progenitors, may become quiescent during the process of involution. Experiments to characterize the epithelial and stromal mediators present in tissue with and without involution, in women with and without subsequent cancer, should help to clarify the mechanisms of risk reduction.

For our work to date, we divided extent of involution into three categories. We recognized that, although the morphologic patterns of age-related lobular involution have been defined (4,5), no histologic standard exists for evaluating the extent of breast involution. In particular, there is no well-characterized method for grading partial degrees of involution. For this reason, we attempted to classify degree of involution with the least amount of subjectivity. Thus, by deciding only whether breast tissue had no lobular involution versus almost complete involution and then by combining the remainder into one category of partial involution, we minimized the subjectivity inherent in judging percent involution.
Our study has several limitations. First, these findings do not necessarily pertain to all women because the cohort studied included women who had a breast biopsy because of some concern. Nevertheless, women with benign breast disease make up a large proportion who are understandably concerned about their breast cancer risk (estimated at 1 million US women each year) (21–23). Another limitation lies in our current very broad category of partial involution. This category encompasses a wide range of involution extent (1%–74% of lobules involved). We expect that more specific gradations would support more refined association studies. Finally, we did not have complete risk factor data for all the women in the cohort, largely because the women with biopsy examinations in the earlier years of the cohort are now elderly or deceased. Fortunately, for purposes of this report, we did not have to depend on the questionnaire for involution status or for cancer outcomes (which were available from our comprehensive Mayo medical record). We had completed questionnaires for 63.8% of the patients with breast cancer and 61.6% of the patients without breast cancer in the cohort.

There are other approaches to the study of involution and breast cancer risk. Henson and Tarone (7) suggested an autopsycase-control series to look at involution as a possible risk factor for breast cancer. Although this approach would provide access to extensive amounts of breast tissue, the availability of clinical risk factor information and of a sufficient number of subjects could be limiting. Other women who have breast tissue removed in the course of clinical care are those who have reduction mammoplasty or prophylactic mastectomy. These women are, respectively, those who have breast hypertrophy or a hereditary predisposition to breast cancer. Although involution (or lack thereof) in these women is of considerable interest, their tissue is not necessarily representative of the general population.

The mechanisms controlling age-related involution are of considerable interest. Molecular programs that control postlactational involution in rodents have been studied extensively (24). With postlactational involution, there is dramatic reversal of the developmental changes wrought by pregnancy. Specifically, there is widespread apoptosis of alveolar epithelial cells followed by removal of apoptotic debris and remodeling of the stroma and extracellular matrix (24). These events occur within a matter of days of abrupt weaning and restore the gland to its prepuberty state. In contrast, the molecular orchestration of age-related involution, to our knowledge, has not yet been characterized.

In the past, for women with benign breast biopsy results, the type and extent of epithelial proliferation present in their biopsy has been the principal way to stratify their risk. Results of our study indicate that assessing the status of lobular involution in the biopsy tissue may ultimately add to risk prediction capabilities. It is notable, as shown in Fig. 2C, that some of the most extreme risk estimates are observed in women whose involution status is unusual for their age—namely, young women with complete involutions (RR = 0.43, 95% CI = 0.13 to 1.55) and women older than 55 years with no involutions (RR = 5.2, 95% CI = 1.90 to 5.08). It is tempting to speculate that the process of complete involution may be protective and, conversely, that lack of involution identifies higher risk groups. However, confidence intervals were wide around the estimates for these less common categories.

In summary, we have evaluated the extent and effect of age-related lobular involution in a cohort of approximately 9000 women who had a benign breast biopsy examination. We observed a statistically significant reduction in risk of breast cancer among women whose breast tissue had undergone extensive lobular involution, which was apparently independent of other modifiable risk. Among women with benign breast disease, assessment of extent of involution may help to pre-tune current risk prediction approaches. Elucidation of the mechanism of lobular involution may reveal ways to promote the process as a means of risk reduction.

REFERENCES


1605 ARTICLES

NOTES

Supported by a Department of Defense Center of Excellence Grant (PREDAMD17-02-1-0573-1), a grant (RO1 CA63332) from the National Institutes of Health, and grants from the Susan G. Komen Breast Cancer Foundation (DCCTR 99-2152), Armstrong Foundation, Breast Cancer Research Foundation, and Regis Foundation for Breast Cancer Research. The authors take full responsibility for the study design, collection of the data, analysis and interpretation of the data, the decision to submit the manuscript, and the writing of the manuscript.

We are indebted to Joel Wern and De Pietro de Groot for database development; to Wilma Ligth, PhD, and the Biopspecimen Core for tissue processing; to Teresa Allen, Mary Amorello, Mary Campbell, Joanne Johnson, Melanie Kassner, Maggie Logman, and Lois Perhach for data collection; to Ann Harris and the Survey Research Center for patient follow-up; and to Vicki Shew for help in preparing the manuscript.

Funding to pay the Open Access publication charges for this article was provided by a grant from the Department of Defense.

Manuscript received July 18, 2006; revised August 31, 2006; accepted September 22, 2006.
Appendix C: Editorial Accompanying Involution Paper
Lobular Involution: the Physiological Prevention of Breast Cancer

Donald Earl Henson, Robert E. Tarone, Hala Nsouli

It truly is a remarkable event when traditional pathologic observations lead to new ideas about the prevention of cancer. In this issue of the Journal, Milanese et al. (1), through a histologic review of breast biopsy specimens, show that the extent of age-related lobular involution is strongly associated with a reduced risk of breast cancer. Breast cancer risk decreased with increasing extent of involution in both high-risk and low-risk subgroups defined by age, epithelial atypia, reproductive history, and family history of breast cancer.

Beginning in the premenopausal period, lobular involution is a physiologic process that occurs over many years whereby the parenchymal elements in the breast progressively atrophy and disappear (2, 3). The study reported by Milanese et al. represents a unique application of the Mayo Benign Breast Disease Cohort to investigate prospectively involution as a risk factor for breast cancer. It is the first study, to our knowledge, to substantiate a hypothesis that is based on pathologic and epidemiologic considerations that delayed involution is a major risk factor for breast cancer (4, 5).

As for an explanation of the effect of lobular involution on breast cancer risk, it has been suggested that a reduction in mammary gland tissue that results from involution should lead to a reduction in breast cancer because a progressively smaller amount of epithelial tissue is available for malignant transformation (1, 5). The result of involution, therefore, can be considered physiologically analogous to a partial prophylactic mastectomy, with a corresponding reduction in breast cancer risk.

Although a reduction in mammary tissue is a plausible explanation, the underlying issue is one of aging or, more precisely, the failure of breast tissue to age normally. The aging process in the breast is under control of various hormones and does not follow the pattern seen in other organs or tissues. Pathologists have long commented on the possibility that persistent atypical lobules might be precursors of invasive breast cancer (6, 7). It seems paradoxical that an organ that normally undergoes complete or near complete physiologic atrophy would be a site in which cancer rates steadily increase with age. The continuing increase in breast cancer risk with age is likely associated with the persistence of glandular epithelium beyond the time of normal involution, reflecting an abnormal delay of the aging process in the breast (4, 5).

Except for morphologic observations concerning age of onset and progression with age, practically nothing is known about the process of involution. Even less is known about factors that control involution or that delay or accelerate the process. In this context, it is unknown whether the rate of involution is genetically determined and whether known breast cancer risk or environmental factors alter the rate of involution.

Evidence indicates that some risk factors for breast cancer may interfere or affect the process of involution. In the Mayo study, women with benign proliferative breast disease were substantially less likely to have complete involution than were women with benign nonproliferative disease, and women with a strong family history of breast cancer had slightly less advanced involution than women without such history (1). Late age of menopause, which increases the risk of breast cancer, is likely to result in delayed involution because of persistence of estrogen activity (8). Women whose first full-term pregnancy occurs after age 35 years have an elevated risk for breast cancer compared with nulliparous women or with women whose first pregnancy was at a much younger age (9, 10). After the commencement of involution, late pregnancy with its concomitant increase in the proliferation of the ductal – alveolar epithelium is likely to interrupt the normal process of involution, which typically begins between 30 and 40 years of age.

Oophorectomy, which protects against breast cancer (11), leads to the same type of atrophy of breast parenchyma in young women as that seen in older women (12). The reduction in risk may be due to the acceleration of involution induced by oophorectomy.

One of the most striking findings in the study of Milanese et al., however, is the degree to which the strong association between extent of involution and breast cancer risk was independent of all known breast cancer risk factors that were investigated (1). This observation suggests that factors unrelated to known risk factors are responsible for the protective effect of involution. For this reason, a greater understanding of the biologic basis for involution will be required to elucidate the mechanisms of the protective effect of lobular involution on breast cancer risk.

The observations reported by the Mayo group may provide practical applications for risk prediction (1). It may be useful for pathologists to report the extent of involution in addition to any epithelial changes found in breast biopsy specimens that do not contain cancer. It will be important to determine the extent to which mammographic breast density serves as a surrogate for the extent of involution. By taking extent of involution into account, it should be possible to increase the predictive ability of breast cancer risk models.

Results of the Mayo study provide a new paradigm for understanding the biology of breast cancer risk and may lead to new insights into the prevention of this disease.
for breast cancer research and prevention. Age has always seemed the opponent because of the increasing risk of breast cancer with age, but age may now become an ally. The challenge will be to unravel the natural history of involution and the normal process of aging in the breast. Eventually, involution could become a useful surrogate endpoint for research in breast cancer prevention. A possible approach to prevention may be to develop strategies that achieve complete involution as early as possible after childbearing is completed.

Affiliations of authors: Office of Cancer Prevention and Control (DEH) and Department of Epidemiology and Biostatistics, School of Public Health and Health Services (HN), The George Washington University Cancer Institute, Washington, DC; International Epidemiology Institute, Rockville, MD (RET); Department of Medicine and Vanderbilt-Ingram Cancer Center, Vanderbilt University Medical Center, Nashville, TN (RET).

Correspondence to: Donald E. Henson, MD, Office of Cancer Prevention and Control, The George Washington University Cancer Institute, Ross Hall, Rm. 502, 2300 “ I ” St. NW, Washington, DC 20037 (e-mail: patdeh@gwumc.edu).

See “ Note ” following “ References. ”

DOI: 10.1093/jnci/djj454
© The Author 2006. Published by Oxford University Press. All rights reserved. For Permissions, please e-mail: journals.permissions@oxfordjournals.org.

REFERENCES


NOTE

The authors would like to acknowledge the constructive comments from Dr. William Anderson.