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TITLE: Exploring Early Detection Methods: Using the Intraductal Approach to Predict Breast Cancer

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**Exploring Early Detection Methods: Using the Intraductal Approach to Predict Breast Cancer**

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**Sponsoring Agency:**
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**Abstract**
Breast cancer is a leading cause of death among women in the U.S. Early diagnosis is believed to be key to minimizing mortality, thus, techniques to identify high-risk women are essential. This study is using an interdisciplinary approach to conduct a follow-up study on a group of 3413 women from the Santa Barbara, Ca area who had breast fluids drawn between 1970-1990 using one of the following three methods: nipple aspiration, ductography or ductal lavage. The follow-up study will determine if abnormal cytologic findings from the past are associated with a higher incidence of breast cancer development during the later years. Follow-up methods include direct contact using questionnaires, linkage with the California Cancer Registry (CCR), linkage with the California Department of Vital Statistics and the National Death Index. Data has been completed on 539 subjects, all mailings have been completed. Follow-up phone calls have begun and all follow up and data entry is to be completed by June 15, 2006. 380 cases of breast cancer have been identified via California Cancer Registry and a final merge will be done in June 2006. Analysis of data will begin in July 2006.

**Subject Terms**
breast fluids, breast cancer risk, nipple aspiration, ductal lavage
INTRODUCTION:
Nipple aspiration, ductal lavage and ductography are methods of obtaining breast fluids from women who are neither pregnant or lactating. Breast cells in these fluids can be classified as either normal or as showing various abnormalities including hyperplasia, atypical hyperplasia and cancer. In previous follow-up studies of women who participated in breast fluid and tissue studies, it was shown that women with proliferative cytology (hyperplasia or atypical hyperplasia) were significantly more likely to develop breast cancer than women with normal cytologic findings in breast fluids or than women from whom fluid could not be obtained. (Fabian et al., 2000; Wrensch et al., 2001) This study is following an additional cohort of women from Santa Barbara, CA that had fluids drawn between 1970-1990. Statistical methods of association will be used to determine if women with abnormal cytologic findings developed breast cancer at a higher rate than women with normal cytologic findings or women from whom fluid could not be obtained.

BODY:
The stated goals in the Statement of Work shall be addressed below:
Step 1 – Study Development: A research team is in place and consists of the study P.I., project coordinator and research assistant. The research team meets formally once a week to review study progress. The research team has completed all goals listed in Step 1 including the design and approval of a living and proxy questionnaire, creating a computer-based tracking system for subjects, IRB approval has been obtained from all necessary entities and Kimberly Baltzell has completed all work for her Ph.D. requirements.

Step 2 – A computer tracking database has been set up for all potential subjects. The new study coordinator is Terri Rice, MPH from the UCSF Department of Neurological Surgery. Pagan Morris is the study research assistant and is also part of the UCSF Department of Neurological Surgery. Both members of the research team have extensive experience conducting large epidemiologic follow-up studies. IRB approval was received from the DOD human subjects committee in August, 2005. A first mailing was sent to 2283 members of the living cohort in January, 2006. A first mailing of 100 was sent to proxies for deceased cohort members at the same time. The study dataset was linked with California Cancer Registry and approximately 380 cases of breast cancer were identified in the cohort. All members of the cohort needing address updates were sent to the DMV. Responses have been received from the DMV for all but 137 subjects.

Step 3 – A second mailing has been sent to 633 members of the cohort from whom we have no response. After a mediocre response to the proxy mailing, it was decided that all proxy contact will be done by phone to limit unnecessary costs associated with mailings to uncertain names and addresses. Follow up phone calling has begun to subjects who
have received 2 mailings but have not responded. All mailings have been completed at this time. Follow up phone calling to all subjects, both living and proxies, will be completed by June 15, 2006. The research team continues to meet formally once a week and via phone and email daily.

Step 4 – The study pathologist, Dr. Eileen King, has begun review of various cytologic diagnoses from the original study data abstraction. She is ranking cytology from least to most severe. We have not begun analyses on returned questionnaires at this time. All returned information will be data entered by June 15, 2006.

Step 5 – Not applicable at this point in time.

From the 3,204 members of the original cohort (total of alive and deceased), we have completed data from 539 members of the living cohort and 11 of the proxy cohort (we have not begun pursuit of this arm of the cohort via telephone yet). We have completed all second mailings and linkage with DMV for updated addresses. Phone calls have begun to all remaining subjects who have not responded. All phone contact attempts will be completed by June 15, 2006. Our data has been merged with California mortality tapes (years 1970-1999) and we are purchasing tapes from years 1999-2004 to do a final merge in July 2006. We are also requesting information from the California Vital Stats Department on Multiple Cause of Death (years 1970-2004) to identify additional breast cancer cases. We will request a final merge with California Cancer Registry in July 2006 to identify additional cases of breast cancer that have been diagnosed since our original merge approximately 1 year ago. We have requested IRB approval from Cottage Health Systems, a local cancer registry in Santa Barbara, CA, to link with their registry to identify breast cancer cases diagnosed between 1970-1988, prior to the California Cancer Registry’s data collection which began in 1988. All follow up contact attempts and linkages will be completed by July, 2006. Analysis of collected data will begin in July/August, 2006 with final study results intended to be completed in September, 2006.

KEY RESEARCH ACCOMPLISHMENTS:

Due to the human subjects review delay, we are still in the data collection phase and have no data analysis to present at this time.

REPORTABLE OUTCOMES:

-poster presentation: Oncology Nursing Society – Anahem, CA, April 2004

-poster presentation: DOD Era of Hope meeting – Philadelphia, PA, June 2005

- article – Breast Carcinogenesis – Can the Examination of Ductal Fluid Enhance Our Understanding? ONF, January 2005
- article – Strengths and Limitations of Breast Cancer Risk Assessment. ONF, May 2005
- Teaching opportunity - the grant recipient was co-faculty of record in fall quarter, 2005 for N265 – Cancer Prevention and Early Detection at UCSF Department of Physiological Nursing and has been asked to teach the course again in Fall, 2006
- Employment opportunity – Grant recipient has applied for an adjunct assistant professor position at UCSF Department of Physiological Nursing (4/06) based on experience supported by this grant

CONCLUSIONS:

This section is not applicable at this time.
References


Dear Ms. «LNAME»:

We are contacting you because you were seen by Dr. Otto Sartorius in Santa Barbara between 1970 and 1990. We are continuing to study personal and physiological characteristics of Dr. Sartorius’ patients with and without breast disease. As a prior patient of Dr. Sartorius, you are invited to participate in this research study. Dr. Sartorius was a pioneering physician who developed innovative procedures to help detect breast disease at early stages. Your participation in our study is very important to breast cancer research. By knowing about your current state of health, we can see if information contained in your records years ago predicts where you are today.

Participation involves filling out and returning the short questionnaire included in this packet. There are no medical procedures. All study information is coded so that your personal identity is not revealed. The computer file that contains your name and address is protected and maintained under strict confidentiality. If we do not receive your questionnaire or postcard in a few weeks, we may contact you by phone in the future.

We assure you that your privacy will be maintained. Your participation in this study is completely voluntary; you may refuse to answer any of the questions. Please contact Kimberly Baltzell, Co-Principal Investigator, or Pagan Morris, Research Assistant, at 1-866-282-5444 if you prefer to complete the questionnaire by phone or with any questions you may have regarding the study and/or study materials. You may return the attached postcard or call the above number if you would prefer not to participate. Please sign and return all pink documents and retain all white copies for your records.

Results from the follow-up study will greatly contribute toward establishing whether the techniques Dr. Sartorius pioneered in the 1970’s can predict who might develop breast disease in the future. Although you may have been contacted in the past several years regarding similar information, please complete the enclosed materials. We are continuing the study and would appreciate your most up-to-date information. Thank you very much for your time and consideration.

Sincerely,

Susan M. Love, M.D.                                      Margaret Wrensch, Ph.D.
President and Medical Director                            Professor
Susan Love MD Breast Cancer Research Foundation            UCSF Dept. of Epidemiology & Biostatistics

Marylin Dodd, R.N., Ph.D.                                  Kimberly Baltzell, R.N., Ph.D.
Associate Dean                                             Study Co-Principal Investigator
UCSF Dept. of Physiologic Nursing                           UCSF Dept. of Physiologic Nursing

Enclosures.
Patient Intro-letter

<< Sent Date >>
Dear Family of Ms. «FNAME» «LNAME»:

We are contacting you because Ms. «FNAME» «LNAME» was seen by Dr. Otto Sartorius in Santa Barbara between 1970 and 1990. UCSF and the Susan Love MD Breast Cancer Foundation are following-up Dr. Sartorius’s patients to study personal and physiological characteristics of women with and without breast disease. As the family member of a prior patient of Dr. Sartorius, you are invited to participate in this research study. Dr. Sartorius was a pioneering physician who developed innovative procedures to help detect breast disease at early stages. Participation in our study is very important to breast cancer research. By knowing about Ms. «LNAME»’s state of health, we can see if information contained in her records years ago predicted eventual health outcomes.

Participation involves filling out and returning the short questionnaire regarding Ms. «LNAME» included in this packet. There are no medical procedures. All study information is coded so that her personal identity is not revealed. The computer file that contains all names and addresses is protected and maintained under strict confidentiality. If we have not received the questionnaire or postcard in 2-3 weeks, we will attempt to contact you by phone.

You may not want to tell us about certain information. We assure you that your privacy and Ms. «LNAME»’s privacy will be maintained at all times. Please feel free to contact Kimberly Baltzell, Co-Principal Investigator, or Pagan Morris, Research Assistant, at 1-866-282-5444 if you prefer to complete the questionnaire by phone or with any concerns you may have regarding the study and/or study materials. Participation in this study is voluntary, you may return the attached postcard or call the above number if you would prefer not to participate. Please sign and return all pink documents and retain all white copies for your records.

Results from the follow-up study will greatly contribute toward establishing whether the techniques Dr. Sartorius pioneered in the 1970’s can predict who might develop breast disease in the future. You or Ms. «FNAME» «LNAME» may have been contacted in the last several years regarding this information. A new research team is continuing the study and the completion of the attached materials would be greatly appreciated. Thank you very much for your time and consideration.

Sincerely,

Susan M. Love, M.D.
President and Medical Director
Susan Love MD Breast Cancer Research Foundation

Margaret Wrensch, Ph.D.
Professor
UCSF Dept. of Epidemiology & Biostatistics

Marylin Dodd, R.N., Ph.D.
Associate Dean
UCSF Dept. of Physiologic Nursing

Kimberly Baltzell, R.N., Ph.D.
Study Co-Principal Investigator
UCSF Dept. of Physiologic Nursing

Enclosures.
Family intro-letter

Revised: 09/14/05 by KB
DR. OTTO SARTORIUS’ BREAST CLINIC FOLLOW UP STUDY

University of California, San Francisco
Department of Physiological Nursing &
Susan Love MD Breast Cancer Research Foundation

If you prefer to complete this questionnaire by phone or have any questions, please call:

1-866-282-5444

FOR OFFICE USE ONLY:

☐ Completed via phone
   DATE:_______  INTVWR:_______

☐ Received via mail
CONTACT INFORMATION

Please fill in the requested information below in the event that we need to contact you in the future.

Current address: ____________________________________________________________
    City:__________________________ State _____  Zip code ______
Home phone number: (___ ___) ___ ___ - ___ ___ ___
Work phone number: (___ ___) ___ ___ - ___ ___ ___
Best time to contact you: ____________________________________________

If completed by someone other than addressee, please list your name and relationship:
First name _________________________  Last name _________________________
Your relation to addressee: ______________________________________________

In case you move, or we are unable to reach you at the information above, please provide us with the name of a close friend or relative who would know how to contact you.

First name _________________________  Last name _________________________
Address of person on line above: ____________________________________________
    City:__________________________ State _____  Zip code __________
    Phone number: (___ ___) ___ ___ - ___ ___ ___
Relationship to you: ______________________________________________________

FOR OFFICE USE ONLY:
Remove front and back sheet from questionnaire and store in locked file.
SARTORIUS FOLLOW UP STUDY 2003-2005
University of California, San Francisco
Department of Physiological Nursing
&
Susan Love MD Breast Cancer Research Foundation

Please complete this questionnaire by **circling or filling in** the appropriate answers.

**DEMOGRAPHIC QUESTIONS**

1. Today’s date:  _____/_____/_____
   month /   day   /   year

2. What is your date of birth?  _____/_____/_____
   month /   day   /   year

3. Which number best describes the highest grade or degree that you achieved?
   1 = Less than high school
   2 = High school
   3 = Junior college, Associate’s Degree
   4 = Some college, no degree
   5 = Bachelor’s Degree
   6 = Master’s Degree
   7 = Doctorate Degree
   8 = Other, please specify:_____________________________

4. Please indicate the number that best describes the total pre-tax annual income of all
   members of your household in 2002.
   0 = Less than $24,999   5 = $150,000 to $199,999
   1 = $25,000 to $49,000   6 = $200,000 to $299,000
   2 = $50,000 to $74,000   7 = $300,000 to $399,000
   3 = $75,000 to $99,999   8 = $400,000 to $499,000
   4 = $100,000 to $149,000   9 = $500,000 or more

5. Which number best describes your race or ethnic background?
   1 = Caucasian/White
   2 = Black, African-American
   3 = Chinese-American
   4 = Japanese-American
   5 = Filipina-American
   6 = Mexican
   7 = Other Hispanic or Latina
   8 = Other, please specify:________________________________
BREAST CONDITIONS AND STATUS

6. Over the past 5 years, have you practiced breast self-examination?
   0 = no, never or rarely
   1 = yes, less than once every 6 months
   2 = yes, about once every 2-6 months
   3 = yes, about once every month
   4 = yes, more than once a month
   5 = other, please specify: ____________________________

7a. Have you ever had breast cancer?
   0 = no ➔ **SKIP TO QUESTION 9**
   1 = yes, right breast ➔ Year first found __________
   2 = yes, left breast ➔ Year first found __________
   9 = uncertain; please explain ______________________________________

7b. How was the first breast cancer found if more than one? (Circle all that apply)
   1 = self exam
   2 = clinical breast exam
   3 = mammogram
   4 = ultrasound
   5 = biopsy
   6 = other, please specify: _______________________________
   9 = uncertain; please explain __________________________________

8. Have you ever had a breast lumpectomy (as treatment for breast cancer)?
   0 = no, never
   1 = yes, right breast ➔ Year procedure was done ______
   2 = yes, left breast ➔ Year procedure was done ______
   9 = uncertain; please explain ______________________________________

9. Have you ever had a mastectomy (removal of entire breast) for either breast cancer treatment or prevention of breast cancer?
   0 = no, never
   1 = yes, right breast ➔ Year procedure was done ______
   2 = yes, left breast ➔ Year procedure was done ______
   9 = uncertain; please explain ______________________________________

10. Have you ever had a mammogram?
    0 = no, never
    1 = yes, less than once every 3 years
    2 = yes, about once every 2 years
    3 = yes, once a year
    4 = yes, more than once a year
    5 = yes, other, please specify: _______________________________
11. Have you ever had a breast tissue biopsy (part of or entire lump removed which may have resulted as benign or malignant)?
   0 = no, never → **SKIP TO QUESTION 12**
   1 = yes, LEFT only *please COMPLETE Column A in the table below*
   2 = yes, RIGHT only *please COMPLETE Column B in the table below*
   3 = yes, BOTH breasts *please COMPLETE Columns A & B in the table below*
   9 = uncertain; please explain __________________________________________

Information about your biopsy results. Please circle whether the “finding” was benign, malignant, or unknown. If the finding was **benign**, please circle hyperplasia, atypia, or unknown.

<table>
<thead>
<tr>
<th>COLUMN A – LEFT BREAST</th>
<th>COLUMN B – RIGHT BREAST</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Year of biopsy</strong></td>
<td><strong>Finding</strong> (please circle one)</td>
</tr>
</tbody>
</table>
| **Biopsy #1**          | 1 = benign📍  
                        | 1 = hyperplasia  
                        | 2 = atypia  
                        | 3 = unknown  
                        | 2 = malignant (cancer or in situ)  
                        | 9 = uncertain |
| **Biopsy #2**          | 1 = benign📍  
                        | 1 = hyperplasia  
                        | 2 = atypia  
                        | 3 = unknown  
                        | 2 = malignant (cancer or in situ)  
                        | 9 = uncertain |
| **Biopsy #3**          | 1 = benign📍  
                        | 1 = hyperplasia  
                        | 2 = atypia  
                        | 3 = unknown  
                        | 2 = malignant (cancer or in situ)  
                        | 9 = uncertain |
| **Biopsy #4**          | 1 = benign📍  
                        | 1 = hyperplasia  
                        | 2 = atypia  
                        | 3 = unknown  
                        | 2 = malignant (cancer or in situ)  
                        | 9 = uncertain |
| **Biopsy #5**          | 1 = benign📍  
                        | 1 = hyperplasia  
                        | 2 = atypia  
                        | 3 = unknown  
                        | 2 = malignant (cancer or in situ)  
                        | 9 = uncertain |

If you had more than 5 biopsies, please write the information in question 30.
12. Have you ever had a fine needle aspiration of your breast(s)?
   0 = no, never → **SKIP TO QUESTION 13**
   1 = yes, LEFT only *please COMPLETE Column A in the table below*
   2 = yes, RIGHT only *please COMPLETE Column B in the table below*
   3 = yes, BOTH breasts *please COMPLETE Columns A & B in the table below*
   9 = uncertain; please explain _________________________________________

Information about your aspiration results. Please circle whether the “finding” was benign, malignant, or unknown. If the finding was **benign**, please circle hyperplasia, atypia, or unknown.

<table>
<thead>
<tr>
<th>COLUMN A – LEFT BREAST</th>
<th>COLUMN B – RIGHT BREAST</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Year of aspiration</strong></td>
<td><strong>Finding</strong></td>
</tr>
<tr>
<td></td>
<td>(please check one)</td>
</tr>
<tr>
<td>Aspiration #1</td>
<td>1 = benign ↓</td>
</tr>
<tr>
<td></td>
<td>1 = hyperplasia</td>
</tr>
<tr>
<td></td>
<td>2 = atypia</td>
</tr>
<tr>
<td></td>
<td>3 = unknown</td>
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<tr>
<td></td>
<td>2 = malignant (cancer or in situ)</td>
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<tr>
<td></td>
<td>9 = uncertain</td>
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<tr>
<td>Aspiration #2</td>
<td>1 = benign ↓</td>
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<td></td>
<td>1 = hyperplasia</td>
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<tr>
<td></td>
<td>2 = atypia</td>
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<td>3 = unknown</td>
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<tr>
<td></td>
<td>2 = malignant (cancer or in situ)</td>
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<td></td>
<td>9 = uncertain</td>
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<tr>
<td>Aspiration #3</td>
<td>1 = benign ↓</td>
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<td></td>
<td>1 = hyperplasia</td>
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<td></td>
<td>2 = atypia</td>
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<td>3 = unknown</td>
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<td></td>
<td>2 = malignant (cancer or in situ)</td>
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<td>9 = uncertain</td>
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<tr>
<td>Aspiration #4</td>
<td>1 = benign ↓</td>
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<td></td>
<td>1 = hyperplasia</td>
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<td></td>
<td>2 = atypia</td>
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<td>3 = unknown</td>
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<td>2 = malignant (cancer or in situ)</td>
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<td>9 = uncertain</td>
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<tr>
<td>Aspiration #5</td>
<td>1 = benign ↓</td>
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<td></td>
<td>1 = hyperplasia</td>
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<td>2 = malignant (cancer or in situ)</td>
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<td></td>
<td>9 = uncertain</td>
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</table>

*If you had more than 5 aspirations, please write the information in question 30.*
13. During the last 5 years, would you say that your health in general was:
   1 = excellent   3 = good
   2 = very good   4 = poor

14. Have you ever had any cancer other than breast cancer?
   0 = no
   1 = yes

   (primary site is where the cancer first occurred, e.g. skin, colon, ovary)
   Primary site 1: ____________________________ Age at diagnosis ___
   Primary site 2: ____________________________ Age at diagnosis ___
   Primary site 3: ____________________________ Age at diagnosis ___
   9 = uncertain; please explain __________________________________________

15. Have you ever taken medication to prevent pregnancy? (do NOT include barrier methods such as condoms or diaphragms)
   0 = no, never
   1 = yes

   *please COMPLETE the following table for the medication(s)*
   9 = uncertain; please explain _________________________________________

   Please answer for medication(s) you have taken to prevent pregnancy. For medications you have NEVER TAKEN, please leave blank.

<table>
<thead>
<tr>
<th>Medications to Prevent Pregnancy:</th>
<th>Total years taken</th>
<th>Total months taken</th>
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</thead>
<tbody>
<tr>
<td>BIRTH CONTROL PILLS</td>
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<tr>
<td>1 = not now → Year first taken</td>
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<td>2 = now → Year first taken</td>
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<tr>
<td>Check if not taken continuously</td>
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<tr>
<td>DEPO PROVERA (injections)</td>
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<td>1 = not now → Year first taken</td>
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<td>2 = now → Year first taken</td>
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<td>Check if not taken continuously</td>
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<tr>
<td>NORPLANT (implants)</td>
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<td>1 = not now → Year first taken</td>
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<td>2 = now → Year first taken</td>
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<td>Check if not taken continuously</td>
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<td>PATCH</td>
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<td>1 = not now → Year first taken</td>
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<td>2 = now → Year first taken</td>
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<td>Check if not taken continuously</td>
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<td>OTHER, specify:</td>
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<td>1 = not now → Year first taken</td>
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<td>2 = now → Year first taken</td>
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<td>Check if not taken continuously</td>
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</tbody>
</table>
16. Have you ever taken medications (fertility drugs) to increase your chances of having a child?
   0 = no, never   1 = yes *(COMPLETE table below)*   9 = uncertain

<table>
<thead>
<tr>
<th>Name of medication</th>
<th>Year first taken</th>
<th>Total years taken</th>
<th>Total months taken</th>
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<tbody>
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</table>

17. Have you ever taken female hormones for menopause?
   0 = no, never   1 = yes *(COMPLETE table below)*   8 = not applicable, premenopausal

<table>
<thead>
<tr>
<th>Name of medication</th>
<th>Year first taken</th>
<th>Total years taken</th>
<th>Total months taken</th>
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</table>

18. Have you ever taken Tamoxifen or Raloxifene?
   0 = no, never   1 = yes *(COMPLETE table below)*   9 = uncertain

<table>
<thead>
<tr>
<th>Medication</th>
<th>Year first taken</th>
<th>Total years taken</th>
<th>Total months taken</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tamoxifen</td>
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<tr>
<td>Raloxifene</td>
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</tbody>
</table>

19. Have you ever taken other medications to prevent breast cancer?
   0 = no, never   1 = yes *(COMPLETE table below)*   9 = uncertain

<table>
<thead>
<tr>
<th>Name of medication</th>
<th>Year first taken</th>
<th>Total years taken</th>
<th>Total months taken</th>
</tr>
</thead>
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</table>
FAMILY HISTORY

Please answer the following questions only for your blood relatives (living or deceased)

20. Did your mother ever have breast cancer?
   0 = no
   1 = yes → Age at diagnosis ______
   9 = don’t know

21a. Do you or did you have any sisters?
   0 = no → SKIP TO QUESTION 22a
   1 = yes → How many? ______

21b. How many of your sisters ever had breast cancer? ______ or don’t know

21c. For each sister(s) who ever had breast cancer, how old was she when it was first diagnosed?
   #1 ______ #2 ______ #3 ______ #4 ______ #5 ______ or don’t know

22a. Do you or did you have any daughters?
   0 = no → SKIP TO QUESTION 23
   1 = yes → How many? ______

22b. How many of your daughters ever had breast cancer? ______ or don’t know

22c. For each daughter(s) who ever had breast cancer, how old was she when it was first diagnosed?
   #1 ______ #2 ______ #3 ______ #4 ______ #5 ______ or don’t know

23. Did your mother’s mother ever have breast cancer?
   0 = no
   1 = yes → Age at diagnosis ______
   9 = don’t know

24a. How many sisters does (did) your mother have? ______ or don’t know
   (If none, SKIP TO QUESTION 25)

24b. How many of your mother’s sisters ever had breast cancer? ______ or don’t know

24c. For each of your mother’s sister(s) who ever had breast cancer, how old was she when it was first diagnosed?
   #1 ______ #2 ______ #3 ______ #4 ______ #5 ______ or don’t know
25. Did your father's mother ever have breast cancer?  
   0 = no  
   1 = yes; Age at diagnosis ______  
   9 = don't know

26a. How many sisters does (did) your father have? ______ or don't know  
   (If none, SKIP TO QUESTION 27)

26b. How many of your father's sisters ever had breast cancer? ______ or don't know

26c. For each of your father's sister(s) who ever had breast cancer, how old was she when it was first diagnosed?  
   #1 ______  #2 ______  #3 ______  #4 ______  #5 ______  or don't know

MENSTRUAL AND PREGNANCY HISTORY

27. At what age did you start menstruating? ______

28. Are you still having periods?  
   1 = yes  
   2 = yes, but pregnant, postpartum, or breastfeeding now  
   3 = yes, but infrequently, probably perimenopausal  
   4 = yes, but due to hormone replacement therapy  
   5 = no, went through natural menopause  ➔ Age of last period ______  
   6 = no, went through natural menopause and later had hysterectomy  ➔ Age of last period ______  
   7 = no, had a hysterectomy  
   with womb and one ovary removed  ➔ Age of last period ______  
   8 = no, had hysterectomy  
   with womb and both ovaries removed  ➔ Age of last period ______  
   9 = no, had hysterectomy with only womb removed  ➔ Age of last period ______  
   10 = no, had a hysterectomy, type unknown  ➔ Age of last period ______  
   11 = no, due to chemo therapy or radiation treatment  ➔ Age of last period ______  
   12 = other, please specify: ____________________________
29. Have you ever been pregnant or are you pregnant now?
   0 = no → **SKIP TO QUESTION 30**
   1 = yes  *please COMPLETE the table below*
   2 = currently pregnant  *please COMPLETE the table below for previous pregnancies*

**Pregnancy information:**

<table>
<thead>
<tr>
<th>Pregnancy Number</th>
<th>Outcome (please circle one)</th>
<th>Year Pregnancy Ended</th>
<th>Number of Months the Pregnancy Lasted</th>
<th>Total Number of Months You Nursed (if live born infant)</th>
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</thead>
<tbody>
<tr>
<td>1st</td>
<td>1 = live birth</td>
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<td></td>
<td>2 = miscarriage</td>
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<td>3 = still birth</td>
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<td>4 = other, specify</td>
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<td>9 = uncertain</td>
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<td>1 = live birth</td>
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<td>2 = miscarriage</td>
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<td>3 = still birth</td>
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<td>4th</td>
<td>1 = live birth</td>
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<td>9 = uncertain</td>
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<td>Year</td>
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30. Is there anything else that you would like to tell us?

___________________________________________________________________________
___________________________________________________________________________
___________________________________________________________________________
___________________________________________________________________________
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___________________________________________________________________________
___________________________________________________________________________
___________________________________________________________________________

End of Questionnaire – Thank you very much for your help in this study
DR. OTTO SARTORIUS’ BREAST CLINIC FOLLOW UP STUDY

University of California, San Francisco
Department of Physiological Nursing
&
Susan Love MD Breast Cancer Research Foundation

If you prefer to complete this questionnaire by phone or have any questions, please call:

1-866-282-5444

FOR OFFICE USE ONLY:

☐ Completed via phone
   DATE: ________   INTVWR: ________

☐ Received via mail
CONTACT INFORMATION

Please fill in the requested information below in the event that we need to contact you in the future.

Current address: _______________________________________________
    City:____________________________ State _____ Zip code ______

Home phone number: (__ __ __) __ __ __ - __ __ __ __

Work phone number: (__ __ __) __ __ __ - __ __ __ __

Best time to contact you: _______________________________

If completed by someone other than addressee, please list your name and relationship:
First name _________________________ Last name _______________________
Your relation to addressee: ______________________________

In case you move, or we are unable to reach you at the information above, please provide us with the name of a close friend or relative who would know how to contact you.

First name _________________________ Last name _______________________
Address of person on line above: _______________________________________
    City:____________________________ State _____ Zip code __________
    Phone number: (__ __ __) __ __ __ - __ __ __ __

Relationship to you: ______________________________

FOR OFFICE USE ONLY:
Remove front and back sheet from questionnaire and store in locked file.
Please complete this questionnaire concerning your mother/wife/sister/daughter/friend by circling or filling in the appropriate answers regarding her breast cancer experience.

1. Today’s date: _____/_____/_____

   month / day / year

BREAST CONDITIONS AND STATUS

2. Did she ever have breast cancer?
   0 = no
   1 = yes, right breast only ➔ Year first found ___________ or don’t know
   2 = yes, left breast only ➔ Year first found ___________ or don’t know
   3 = yes, both breasts ➔ Year first found ___________ or don’t know
   9 = uncertain; please explain _________________________________________

2a. How was the first breast cancer found, if more than one? (Circle all that apply)
   1 = self exam
   2 = clinical breast exam
   3 = mammogram
   4 = ultrasound
   5 = biopsy
   6 = other, please specify: _______________________________
   9 = uncertain; please explain _________________________________________

3. Did she ever have a mastectomy (removal of a breast)?
   0 = no, never
   1 = yes, right breast only ➔ Year procedure was done ______
   2 = yes, left breast only ➔ Year procedure was done ______
   3 = yes, both breasts ➔ Year procedure was done ______
   9 = uncertain; please explain ________________________________

4. Did she ever have a mammogram (x-ray of a breast)?
   0 = no, never
   1 = yes ➔ Year of first mammogram ____ / Year of most recent mammogram ____
   9 = uncertain; please explain _________________________________________

5. Did she ever have a breast biopsy?
   0 = no, never ➔ skip to question 11
   1 = yes *please complete the following table*
   2 = yes, left only *please complete the following table*
   3 = yes, both breasts *please complete the following table*
   9 = uncertain; please explain _________________________________________
5a. Information about her biopsy results. Please circle (if information is available) whether the finding was either benign, malignant or unknown. If the finding was benign, please circle whether it was hyperplasia, atypia or don’t know.

<table>
<thead>
<tr>
<th>Left breast</th>
<th>Year of biopsy</th>
<th>Finding (please circle benign, malignant or uncertain)</th>
<th>Right breast</th>
<th>Year of biopsy</th>
<th>Finding (please circle one)</th>
</tr>
</thead>
</table>
| Biopsy #1   |                | 1 = benign if yes, circle 1, 2 or 3  
1 = hyperplasia  
2 = atypia  
3 = don’t know  
2 = malignant (cancer)  
9 = uncertain | Biopsy #1 |                | 1 = benign if yes, circle 1, 2 or 3  
1 = hyperplasia  
2 = atypia  
3 = don’t know  
2 = malignant (cancer)  
9 = uncertain |
| Biopsy #2   |                | 1 = benign  
1 = hyperplasia  
2 = atypia  
3 = don’t know  
2 = malignant (cancer)  
9 = uncertain | Biopsy #2 |                | 1 = benign  
1 = hyperplasia  
2 = atypia  
3 = don’t know  
2 = malignant (cancer)  
9 = uncertain |
| Biopsy #3   |                | 1 = benign  
1 = hyperplasia  
2 = atypia  
3 = don’t know  
2 = malignant (cancer)  
9 = uncertain | Biopsy #3 |                | 1 = benign  
1 = hyperplasia  
2 = atypia  
3 = don’t know  
2 = malignant (cancer)  
9 = uncertain |
| Biopsy #4   |                | 1 = benign  
1 = hyperplasia  
2 = atypia  
3 = don’t know  
2 = malignant (cancer)  
9 = uncertain | Biopsy #4 |                | 1 = benign  
1 = hyperplasia  
2 = atypia  
3 = don’t know  
2 = malignant (cancer)  
9 = uncertain |
| Biopsy #5   |                | 1 = benign  
1 = hyperplasia  
2 = atypia  
3 = don’t know  
2 = malignant (cancer)  
9 = uncertain | Biopsy #5 |                | 1 = benign  
1 = hyperplasia  
2 = atypia  
3 = don’t know  
2 = malignant (cancer)  
9 = uncertain |

6. What was the cause of death?  

End of Questionnaire – Thank you
University of California, San Francisco
Research Study Information Sheet
Dr. Otto Sartorius’ Breast Clinic Follow-up Study

A. Purpose and Background

You have been asked to participate in this research study because you had breast fluid specimens evaluated by Dr. Sartorius between 1970 and 1990. Dr. Sartorius collected nipple aspiration specimens as part of his standard clinical assessment. There was no original study outlined or intended at the time of the breast fluid collection. Researchers now believe that following up on this information may be important and are requesting your consent at this time. This study is being conducted by Marylin Dodd, R.N., PhD and Kimberly Baltzell, R.N., PhD (1-866-282-5444 ) in the Department of Physiological Nursing, UCSF, Margaret Wrensch, PhD in the Department of Neurological Surgery at UCSF and the Susan Love MD Breast Cancer Foundation.

This study will determine breast cancer occurrence in women who participated in breast fluid studies with Dr. Otto Sartorius between 1970 and 1990. The purpose is to decide if women who had abnormal cells in breast fluid specimens were more likely to develop breast cancer than women with normal cells in breast fluid specimens or in women from whom fluid could not be obtained.

B. Procedures

If you agree to be in this study, you will do the following:
1) Fill out the enclosed questionnaire and return it to the investigator in the post-paid envelope. The questionnaire will take approximately 30-45 minutes to complete.
2) You can also participate in this study by calling 1-866-282-5444 to arrange for a telephone interview.

*You are free to decline to answer any questions*

C. Risk and/or Discomforts

The risk from this study is that you may feel some discomfort at recalling your medical history.

Confidentiality: Participation in research may involve a loss of privacy, but information about you will be handled as confidentially as possible. Your name will not be used in any published reports about this study. Study information will be coded and kept in locked files at all times. Only study personnel will have access to the files.
Representatives of the U.S. Army Medical Research and Material Command are eligible to review research records as part of their responsibility to protect human subjects in research. The UCSF Committee on Human Research may also review the research records.

D. Benefits

There is no direct benefit to you from participating in this study. The anticipated benefit from this study is confirmation that nipple aspirate fluid (cellular studies) may be a useful tool in addition to other screening methods to identify women who may be at high risk of breast cancer.

E. Alternatives

An alternative is not to participate in the study.

F. Costs and Reimbursements

There will be no costs and no reimbursements to you for taking part in this study.

G. Potential Conflict of Interest and Funding

The researchers conducting this study do not have any known financial interests that may affect the performance or interpretation of this research. Funding for this study has been provided by the Department of Defense, Breast Cancer Research Dissertation Award #BC021862.

H. Questions

If you have any questions or comments about participating in this study, you should first talk with Dr. Marylin Dodd or Dr. Kimberly Baltzell at 1-866-282-5444. If for some reason you do not wish to do this, you may contact the Committee on Human Research, which is concerned with protection of volunteers in research projects. You may reach the Committee office between 8:00 a.m. and 5:00 p.m. Monday-Friday, by calling 415-476-1814, or by writing to the Committee on Human Research, Suite 11, Laurel Heights Campus, Box 0962, University of California, San Francisco, CA 94143.

I. Consent

Enclosed please find a copy of the Experimental Subject’s Bill of Rights to keep.

PARTICIPATION IN RESEARCH IS VOLUNTARY. You have the right to decline to participate or to withdraw at any point in this study without jeopardy to your present or future status as a patient at UCSF.

____________________________________  _________________________
Participant’s Signature      Date
Exploring Early Detection Methods: Using the Intraductal Approach to Predict Breast Cancer Risk

Dr. Otto Sartorius’ Breast Clinic Follow-up Study

1. Purpose, Participation and Procedures

You are invited to participate in this research study because your relative, Ms. <<fname>> <<lname>> had breast fluid specimens evaluated by Dr. Sartorius between 1970 and 1990. Marylin Dodd, RN, PhD and Kimberly Baltzell, RN, PhD (1-866-282-5444) in the Department of Physiological Nursing, UCSF, Margaret Wrensch, PhD in the Department of Neurological Surgery at UCSF and the Susan Love MD Breast Cancer Foundation are conducting the study.

This study will determine breast cancer occurrence in women who participated in breast fluid studies with Dr. Otto Sartorius between 1970 and 1990. The purpose is to decide if women who had abnormal cells in breast fluid specimens were more likely to develop breast cancer than women with normal cells in breast fluid specimens or than women from whom fluid could not be obtained.

If you agree to participate in the study, you will do the following:
1) Fill out the enclosed questionnaire and return it to the investigator in the post-paid envelope. The questionnaire will take approximately 10-15 minutes to complete.
2) You can also participate in the study by calling 1-866-282-5444 for a telephone interview.

*You are free to decline to answer any questions*

2. Description of Risks

The risk from this study is that you may feel some discomfort at recalling Ms. <<lname>>’s medical history. Participation in research may involve a loss of privacy, but information about Ms. <<lname>> will be handled as confidentially as possible.

3. Confidentiality

Ms. <<lname>> will not be used in any published reports about this study. Study information will be coded and kept in locked files at all times. Only study personnel will have access to Ms. <<lname>>’s files. Representatives of the U.S. Army Medical Research and Material Command are eligible to review research records as part of their responsibility to protect
human subjects in research. The UCSF Committee on Human Research may also review the research records.

4. Benefits

There is no direct benefit to you from participating in this study. The anticipated benefit from this study is confirmation that nipple aspirate fluid (cellular studies) may be a useful tool in addition to other screening methods to identify women who may be at high risk of breast cancer.

5. Alternatives

An alternative is not to participate in the study.

6. Potential Conflict of Interest and Funding

The researchers conducting this study do not have any known financial interests that may affect the performance or interpretation of this research. Funding for this study has been provided by the Department of Defense, Breast Cancer Research Dissertation Award #BC021862.

7. Questions

If you have any questions or comments about participating in this study, you should first talk with Dr. Marylin Dodd or Kimberly Baltzell, RN, PhD at 1-866-282-5444. If for some reason you do not wish to do this, you may contact the Committee on Human Research, which is concerned with protection of volunteers in research projects. You may reach the Committee office between 8:00 a.m. and 5:00 p.m. Monday-Friday, by calling 415-476-1814, or by writing to the Committee on Human Research, Suite 11, Laurel Heights Campus, Box 0962, University of California, San Francisco, CA 94143.

8. Consent

Enclosed please find a copy of the Experimental Subject’s Bill of Rights to keep.

PARTICIPATION IN RESEARCH IS VOLUNTARY. You have the right to decline to participate or to withdraw at any point in this study without jeopardy to your present or future status as a patient at UCSF.

______________________________________________  _________________________
Participant’s Signature      Date
Sartorius Clinic Follow-up Study

id#_____________________

[ ] I may be interested in hearing more about the study.
You can contact me at:

( )_____________________ _____________________________
phone number good time to call

[ ] I am unable to participate in this study but please contact my spouse/relative/friend, ____________________________,
who may be able to participate as my proxy.

( )_____________________ _____________________________
phone number good time to call

[ ] I do not wish to participate in the follow-up study.
Please do not contact me further.

Sign here: ___________________________________________
Breast Carcinogenesis—Can the Examination of Ductal Fluid Enhance Our Understanding?

Kimberly Baltzell, RN, PhD(c), Suzanne E. Eder, NP, RN, and Margaret Wrensch, MPH, PhD

**Key Points...**

- Many breast carcinogenic theories support the notion of a cellular continuum from normal epithelium through multiple proliferative stages to malignancy.
- Examining breast epithelial cells over time to determine when premalignant changes occur may lead to enhanced risk prediction.
- Obtaining breast epithelial cells via nipple aspiration, ductal lavage, or periareolar fine-needle aspiration may be a less invasive way to acquire information on breast cancer risk than currently achieved by breast biopsy.

A n understanding of normal cellular transformation to malignancy is not defined clearly in the study of breast cancer. Deciphering the breast cancer pathophysiologic pathway is necessary for the design of effective cancer prevention strategies (Miller, Bates, & Nabell, 2002). Recent studies showing a significant association between proliferative breast cells and increased risk of breast cancer development highlight the importance of clarifying precursors to disease development (Fabian & Kimler, 2001; Wrensch et al., 2001). Studzinski and Harrison (2002) wrote that precise breast cancer diagnosis, monitoring, and treatment require understanding the control of cell growth, which may lead to the ultimate goal—prevention. Studying the progression from normal cell growth patterns to malignancy has been difficult because of the populations on whom most research has been performed. These populations typically include patients with advanced or metastatic disease. These studies may be limited in their usefulness because events surrounding carcinogenesis already have taken place (Briand & Lykkefeldt, 2001). Researchers generally agree that carcinogenesis is a result of a combination of inherited susceptibility (germline mutations) and acquired genetic changes (somatic mutations), possibly involving more than 200 genes (Miller et al.; Studzinski & Harrison). This article will discuss the current theories of breast carcinogenesis, emphasizing the progression of normal cells through malignant transformation. Carcinogenesis theory lends support to the idea of using breast epithelial cells to analyze possible precursors to malignancy, leading to enhanced breast cancer risk-prediction models. Types of intraductal sampling techniques will be reviewed, as well as the correlation between tissue cytology and intraductal cytology.

Kimberly Baltzell, RN, PhD(c), is a doctoral candidate in the Department of Physiologic Nursing at the University of California, San Francisco (UCSF). Suzanne E. Eder, NP, RN, is a nurse practitioner at the UCSF Breast Care Center, and Margaret Wrensch, MPH, PhD, is a professor in the Department of Neurological Surgery and Epidemiology/Biostatistics at UCSF. (Submitted December 2003. Accepted for Publication September 3, 2004.) (Mention of specific products and opinions related to those products do not indicate or imply endorsement by the Oncology Nursing Forum or the Oncology Nursing Society.)

Digital Object Identifier: 10.1188/05.ONF.33-39
Carcinogenesis Theory Overview

Development of Breast Cells

Breast cells begin to complete their growth during puberty. Prior to that time, the mammary gland consists of a fat pad with a primary duct and several ductal branches (Miller et al., 2002). With the onset of menarche, rapid growth occurs, regulated by estradiol and progesterone. The cyclic nature of estrogen exposure continues to act on the breast tissue, yet ductal development stops after puberty is completed.

Although the formation of ducts is over, the duct end buds continue interchanging rounds of growth and cessation in response to hormonal changes produced by the menstrual cycle (Miller et al., 2002). This balance of proliferation and apoptosis (cell death) keeps the breast epithelium in check, and imbalance in this system is the basis of many carcinogenesis theories. The protective effect of full-term, early pregnancy is linked to its association with ductal differentiation, leaving breast cells less vulnerable to these cyclic events that may lead to cancer development.

Estrogen is thought to play a key role in the development of normal breast cells, as well as the development of breast cancer cells (Allred, Mohsin, & Fuqua, 2001). Estrogen is responsible for the elongation of breast ducts and thickening of the epithelium that occurs in puberty (Rosen, 2001). Differentiation of the lobuloalveolar units occurs during puberty, with insulin, progesterone, and growth hormone contributing to the process (Mccarty & Tucker, 1992; Rosen). These changes continue through menstrual cycles, pregnancy, lactation, and menopause.

Carcinogenesis Theories

Carcinogenesis is described as a multistage process whereby normal cell proliferation continues unchecked because of aberrant genetic or chromosomal alterations, leading to invasive and metastatic growth (Briand & Lykkefeldt, 2001).

Cancer of the breast generally is divided into two etiologic origin groups. The first group is cancer that is deemed to arise from strong hereditary sources, primarily a mutation of either the BRCA1 or BRCA2 gene (Miller et al., 2002). These germline mutations are believed to be responsible for about 5%–10% of all breast cancers and 65% of all inherited breast cancers. The second group of breast cancers, the remaining 90%, is defined as sporadic and nonfamilial. The processes for both types of cancers, however, seem to be a combination of genetic susceptibility and environmental factors (Briand & Lykkefeldt, 2001). Epigenetic factors are defined as altered expression of genes, although base pairs remain unchanged (Tannock & Hill, 1998). Epigenetics may hold great promise for future interventions, given that epigenetic alterations are reversible and mutations are not.

What is known about carcinogenic pathways? Five individual steps necessary for malignant transformation have been proposed (Hahn & Weinberg, 2002). The steps are independence from mitogenic stimulation, evasion of apoptosis, immortalization, resistance to exogenous growth-inhibitory signals, and angiogenesis.

Mitogenic stimulation independence may occur as a result of the mutation of an oncogene (e.g., ras, HER2-neu), in essence turning on a cell’s ability to override its own growth control checks. Cancer cells do not depend on external signals to make a commitment to proliferate (Hahn & Weinberg, 2002). A breast cancer oncogene of interest is HER2-neu.

Mutations of these genes may occur by base substitutions, translocation, amplification, or viral insertions. Whatever the method of mutation or activation employed, the affected cell takes on an enhanced capacity for growth. HER2-neu is an oncogene that frequently is overexpressed in tumors (Miller et al., 2002). Tumors with an abundance of this oncogene often have poorer responses to chemotherapy; however, this is an exciting area of exploration for new treatment modalities.

The evasion of apoptosis might occur as a result of a mutated tumor suppressor gene (e.g., p53) inhibiting the back-up system in place for both cell overgrowth and damaged cell surveillance and repair. Mutated p53 is present in about 30%–40% of human cancers (Dickson & Lippmann, 2000). It is the most frequently studied tumor suppressor gene, which, under normal circumstances, functions as an apoptosis inducer or inhibitor of cell overgrowth. Mutated p53 interferes with normal p53, and researchers have speculated that restoring normal p53 may inhibit cancer growth (Yin, Tainsky, Bischoff, Strong, & Wahl, 1992).

Immortalization results from damage to telomeres (the chromosomal end caps), allowing cells to maintain their proliferative potential indefinitely. Even in the presence of proper nutrients and space, normal cells stop dividing as the telomeres shorten and no longer can stabilize chromosomes.

A malignant cell, in contrast, maintains its proliferative potential indefinitely. Molecular mechanisms that inhibit this cell senescence are unclear (Tannock & Hill, 1998).

Resistance to exogenous growth-inhibitory signals works in tandem with one of the other behaviors of cancer cells, independent mitogenic stimulation, allowing cells to proliferate unchecked. All interrupted pathways lead to the hallmarks of malignancy: an increase in cell proliferation and lack of cell death. Finally, the ability of a cell to create additional blood flow appears to be a trait of cancer cells. Circulatory access is believed to be necessary for a tumor to grow larger than two centimeters.

Hormones play a major role in the development of breast cancer. Henderson, Pike, Bernstein, and Ross (1996) wrote that the role of hormones involves their effects on breast cell proliferation and that this increased cell division is vital for the genesis of human cancer. They also cited the activation of oncogenes and mutation of tumor-suppressor genes as necessary for the development of a malignant phenotype. This progression is illustrated in Figure 1.

![Figure 1. Progression to Malignant Phenotype](image)

*Note. Based on information from Henderson et al., 1996.*
Knudson (1971) inspired many carcinogenesis models based on his theory of a multistep process involving an initial "hit" of one of the tumor-suppressor gene alleles, inactivating it, resulting in homozygosity of the chromosome. In addition, cell division is required for all processes leading to breast cancer development. This theory supports a cellular continuum of normal cell appearance through an abnormal proliferative phase, followed by the progression to a malignancy.

Other studies have debated the hypothesis that cancer arises from mutations. Prehn (1994) wrote that mutations may have limited biologic significance. Cancer is hypothesized to give rise to mutations, rather than mutations giving rise to cancer. This theory is based on the epigenetic events surrounding breast cancer development; however, progression from a normal cellular state through abnormalities into malignancy is supported.

Vineis (2003) proposed a Darwinian approach to carcinogenesis whereby epigenetic events influence a cell's decision to progress to malignancy. The two phases are genetic change followed by selective advantage. The resistance of cells to events such as apoptosis allows for survival of the fittest, allowing mutated cells to adapt more readily to specific environmental niches better than normal cells. Vineis used this hypothesis to explain the difference in international rates of breast cancer because genetic differences account for only a small portion of the variation. Changes in environment as well as the presence of "selective advantage" combine to create cancer rates for specific populations. Willet, Rockhill, Hankinson, Hunter, and Colditz (2000) attributed the increase in breast cancer incidence in women who migrated from low-risk countries (primarily Asian) to high-risk countries (primarily Northern European) to the length of time spent in the high-risk country and adoption of the destination country's lifestyle.

Briand and Lykkezenfeld (2001) reviewed a decade of work on a human breast epithelial cell line, HMT-3522, to formulate an epigenetic model for breast carcinogenesis. They cautioned that following breast cancer events in advanced cases does not illuminate events related to how carcinogenesis actually begins. They believed that cell culture is an appropriate medium for exploring the events that lead to malignant transformation. The study's hypothesis suggested that mutation is a necessary step in the carcinogenesis process; however, epigenetic events influence which cells progress to cancer.

The primary assumption made in the study of breast carcinogenesis is the notion of cells progressing on a continuum. Although which cells will progress to a malignant state from a proliferative state (hyperplasia or atypia) is unknown, recent studies showing an increased risk of breast cancer development in women with proliferative findings have suggested a relationship (Wrensch et al., 2001). The ability to invade surrounding breast tissue and metastasize is present in 20%–50% of breast precancers (O'Shaughnessy, 2000). If hyperplasia and atypical hyperplasia are the result of the first several steps in the process outlined by Hahn and Weinberg (2002), identifying these cellular changes prior to circulatory access and commitment to metastasis is critical. The theory of malignant transformation using cell culture supports the concept of malignant conversion (Martin, 1996). By recognizing the progression of abnormal cell development as a continuum, some borderlines have been created between benign states and malignancies. Page and Rogers (1992) disputed the idea of categorizing cells as either benign or malignant. All tumor cells are believed to have sprung from a single cell, and tumor progression is a phenomenon that concludes that benign tumors often evolve into malignancies (Martin). A malignant phenotype arises from the cell population with the most rapid and favored growth pattern. The earlier discussion supports the idea of benign cells revealing changes that may be indicative of a progression to cancer. Perhaps the analysis of breast epithelial cells will illuminate important precursors to breast cancer. Evidence of intraductal and atypical hyperplasia in epithelial cells may allow for prediction and prevention of breast cancer, whereas advanced progression to invasive cancer requires more aggressive vigilance and treatment (see Figure 2.).

**Evaluating Breast Cancer Risk**

The most commonly used models for evaluating breast cancer risk are the Gail model, the Claus model, and BRCAPRO (developed by statisticians at the Duke University Institute for Statistics and Decision Sciences). Each model was designed from a different population, and, because the models are not used uniformly in clinical practice, the accuracy of the results is a function of healthcare providers' knowledge.

The Gail model uses age, age at menarche, number of prior breast biopsies, age at first live birth, and number of first-degree relatives affected by breast cancer to assess risk. Absolute risk is calculated for five years from the time of assessment and lifetime risk up to age 90 (Gail et al., 1989). The model is most appropriate for evaluating risk in women with limited family history of breast cancer. The Gail model uses limited family history of breast cancer and tends to overestimate risk in young women (Kelly, 2000).

Another breast cancer risk assessment model was developed by Claus, Risch, and Thompson (1993). The model addressed several of the alleged shortcomings of the Gail model by incorporating more extensive family history into the analysis. In addition, the Claus model integrates age at diagnosis of breast cancer into its calculations. This information has become more important since the discovery of BRCA1 and BRCA2 mutations, allowing healthcare professionals to consider the possibility of recommending genetic testing. This model is most helpful in determining risk for women with a strong family history of breast cancer. The nonfamily history information included in the Gail model is not considered in the Claus calculations.

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**Goal: Identify women at highest risk so they can be targeted for a proactive risk management strategy.**

- **Normal duct**
- **Intraductal hyperplasia**
- **Atypical ductal hyperplasia**
- **Ductal carcinoma in situ**
- **Invasive ductal carcinoma**

<table>
<thead>
<tr>
<th>Predict and prevent</th>
<th>Detect and treat</th>
</tr>
</thead>
<tbody>
<tr>
<td>Normal duct</td>
<td>Invasive ductal carcinoma</td>
</tr>
<tr>
<td>Intraductal hyperplasia</td>
<td></td>
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<tr>
<td>Atypical ductal hyperplasia</td>
<td></td>
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<tr>
<td>Ductal carcinoma in situ</td>
<td></td>
</tr>
</tbody>
</table>

**Figure 2. Cellular Progression From Normal Duct Epithelium to Carcinoma**

*Note. Image courtesy of Cytyc Corporation and affiliates. Used with permission.*
Computer programs also have been designed to assess women's risk of a BRCA1 or BRCA2 mutation. The BRCAPro program is considered to be the most comprehensive estimate of genetic mutation risk and has been compared favorably against the assessment of experienced risk counselors (Euhus, Smith, et al., 2002).

Although each of the models is useful in specific populations, no tool completely captures the many factors believed to contribute to a woman's risk of developing breast cancer. Viewing cells directly from the breast duct epithelium would allow the addition of biologic information to models of risk assessment. Cells can be obtained through nipple aspiration, ductal lavage, and periareolar fine-needle aspiration (FNA).

**Obtaining Epithelial Cells to Evaluate the Carcinogenic Process**

The ability to study breast epithelial cells for precancerous changes is necessary to evaluate where in the carcinogenic process intervention is most effective. Studies that have found a strong association between the presence of hyperplasia and atypical hyperplasia and future breast cancer development give this exploration credibility (Fabian et al., 2000; Wrensch et al., 2001). Tissue biopsy is an unrealistic screening tool in large populations of women. Other less invasive methods of obtaining breast epithelial cells include nipple aspiration, ductal lavage, and periareolar FNA. Although no specific screening guidelines exist at present, all results obtained from these methods are interpreted in the context of a breast cancer risk assessment. Appropriate candidates for epithelial cell study include women with a family history of breast cancer, a known genetic mutation such as BRCA1 or BRCA2, or a prior history of breast cancer (to assess the contralateral breast). Additionally, these women should be asymptomatic with a normal breast examination and screening mammogram.

**Nipple Aspiration**

Obtaining breast epithelial cells through a simple suction technique is known as nipple aspiration. This technique was pioneered by George Papanicolaou, MD, based on cytopathologic evaluation of cervical specimens and their relationship to cervical cancer (Papanicolaou, Holmquist, Bader, & Falk, 1958). Studies have shown varying degrees of success in obtaining nipple aspirate fluid (NAF) using aspiration. Sauter et al. (1997) concluded that NAF can be obtained in essentially all eligible subjects. Other studies have reported that nipple aspiration is far inferior to other techniques such as ductal lavage in obtaining an adequate number of cells for evaluation (Dooley et al., 2001). Past studies have obtained NAF from as few as 25% to as many as 95% of study subjects (Rose, Lahti, Kettunen, & Wynder, 1986). Wrensch et al. (2001) noted that obtaining fluid depends on the quantity of fluid present, duct and nipple characteristics, subject age, and the skill of the technician collecting the fluid. Wrensch et al. (1990) found that four important factors were positively related to the ability to obtain breast fluid: age up to 35–50 years, earlier age at menarche, non-Asian compared to Asian ethnicity, and history of lactation. Of interest is the finding that women who do not yield fluid may be less likely to develop breast cancer than women who do yield fluid (Wrensch et al., 1992).

**Ductal Lavage**

Clinically, ductal lavage is used as a risk assessment tool and in the assessment of suspicious nipple discharge. Ductal lavage has its most important clinical application as a risk assessment tool and is best used in a breast cancer prevention program that addresses the broader issues of breast cancer prevention. Ductal lavage is described as a procedure that uses a microcatheter to cannulate identified ductal orifices for the collection of breast epithelial cells for analysis (Dooley et al., 2001) (see Figure 3). The procedure is performed with only topical anesthesia to facilitate cannula insertion. Dooley et al. found that of 507 women tested, a majority (78%) of subjects' samples were adequate for analysis. The study used comparison groups, examining specimen adequacy of ductal lavage versus nipple aspiration. Of the subjects who underwent ductal lavage, a median of 13,500 cells were collected per duct, with 24% of the subjects showing cellular abnormalities ranging from mild atypia to malignancy. The procedure was well tolerated, with most subjects rating the pain on par with mammography. In addition, ductal lavage was 3.5 times more likely to result in a cytologic diagnosis than nipple aspiration (p < 0.001). The abundance of cells available from ductal lavage makes it a promising tool to enhance risk assessment. Informed consent is obtained prior to the procedure. When educating a woman about ductal lavage, healthcare providers should discuss the procedure, possible adverse effects, possible results, and their implications.

Ductal lavage has five potential cytoclinical interpretations: benign, inadequate cellular material for diagnosis, mild atypical cells, marked atypical cells, or malignant. In discussions about the implications of ductal lavage, healthcare providers must explain that ductal lavage is not a screening tool for breast cancer. Ductal lavage is not a substitute for screening tests such as mammography. The false-negative rate of ductal lavage has not been defined. Women should be counseled about the possible results of ductal lavage and their implications. When the result is benign, the woman must be

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**Figure 3. Steps in Nipple Aspiration and Ductal Lavage**

*Note: Image courtesy of Susan Love, MD. Used with permission.*
counseled that several ducts have not been sampled. A benign result gives information on only the ducts sampled. Follow-up would include ductal lavage performed on a yearly basis for continued risk assessment. The frequency of follow-up ductal lavage remains, however, a study question. It currently is based on the frequency of traditional screening methods used in breast cancer, such as mammography.

Limitations of this method include the possibility of infection, injury to the breast, and technical problems that affect cell collection (e.g., dehydration, cold) (Esserman, Adduci, Chew, & Ljung, 2003). In addition to these limitations, ductal lavage is not yet considered the standard of care in breast cancer prevention. Most insurance companies will not authorize or provide reimbursement for ductal lavage. The current fee for ductal lavage is about $900 per duct. During a ductal lavage, as many as four ducts may be accessed. Patients may receive ductal lavage by participating in study protocols, in which case they are not burdened with providing payment.

**Fine-Needle Aspiration**

FNAC often is recommended for clinical diagnosis of suspicious breast lumps (Hughes, Mansel, & Webster, 2000). This procedure provides highly accurate information (99% accuracy rate) when performed by skilled practitioners and read by experienced cytopathologists (Barrows, Anderson, Lamb, & Dixon, 1986). In addition to providing diagnostic information about breast lumps, periareolar FNAC is being explored as a potential methodology for assessing cellular characteristics leading to increased breast cancer risk (Fabian et al., 2000). Fabian et al. suggested that limitations of other methods discussed earlier point to the feasibility of using periareolar FNAC to obtain specimens for risk assessment. In their study, which updated results from a cohort of 480 high-risk women (defined as having one of the following major risk factors: family history of breast cancer, prior lymph node-negative breast cancer, or a prior biopsy indicating atypical lobular or ductal hyperplasia or carcinoma in situ), cyologic evidence of atypical hyperplasia was predictive of breast cancer development. The authors cautioned that this procedure is best employed with women who are premenopausal or those who are postmenopausal and receiving hormone replacement therapy (HRT) because of the limitations of periareolar FNAC in obtaining adequate specimens in fatty or involuted breast tissue. HRT delays the development of fatty breast tissue, maintaining a breast structure similar to premenopausal breast tissue. Other studies have used periareolar FNAC to enhance individual risk assessment (Euhus, Cler, et al., 2002). Using loss of heterozygosity in breast epithelium as the marker of interest, Euhus, Cler et al. were able to demonstrate that periareolar FNAC may be a feasible method for molecular analysis to define subsets of high-risk women. Masood (1999) emphasized the importance of standardizing both the practice and interpretation of periareolar FNAC to justify its use in breast cancer studies, paying particular attention to well-established cytomorphic criteria (see Table 1).

**Correlation Between Tissue Cytology and Intraductal Cytology**

If any of the methods of extracting breast epithelial cells are to be useful in assessing risk, a strong correlation must be present between findings in tissue biopsy (the current gold standard for analyzing breast cell changes) and less invasive means of obtaining those cells. Because 90% of breast cancers are believed to be of ductal-lobular origin, analyzing cells from the ducts to determine whether any precancerous changes have taken place is logical. King, Chew, Petrakis, and Ernster (1983) assigned strict criteria for evaluating cytomorphic changes in breast epithelial cells. The most important finding of their study was the significant association between atypical hyperplasia found in epithelial cells in nipple fluid and atypical hyperplasia found in biopsy tissue. The authors also concluded that the relationship between atypical hyperplasia in the two sources was most significant for women with more marked changes. Using epithelial cells from breast fluid was less reliable for women with benign breast disease. In addition, the study was one of the first to compare cytology between nipple fluid and biopsy using morphologic terms applied to tissue biopsy. One study, which evaluated cells from nipple aspiration only, found cytologic and histologic correlation only when ductal carcinoma in situ and extensive nipple involvement were found in the tissue biopsied (Krishnamurthy et al., 2003). This may be a limitation overcame by using one of the other methods outlined earlier, such as ductal lavage or FNAC.

**Sensitivity and Specificity Issues**

To provide meaningful information, methods of obtaining epithelial cells must have acceptable levels of sensitivity and specificity. Sensitivity is defined as the ability of the test to truly determine the presence of a real breast cancer precursor, and specificity is the ability of the test to correctly identify cells that would not lead inevitably to breast cancer (Last, 2001). Sensitivity is the rate of true positives; specificity is the rate of true negatives.

Ductal lavage yields abundant epithelial cells for evaluation (Dooley et al., 2001). Cytologic studies are performed easily on these specimens; however, what to do with the information remains unclear. Recent studies have questioned the sensitivity and specificity of this method, suggesting that it remains a breast cancer detection method best used in clinical trials (Domchek, 2002). Dooley et al. found ductal lavage to be 3.2 times more sensitive in detecting abnormalities in breast cells than nipple aspiration (79 versus 32 breasts) in a study of 507 women. Sensitivity is less of a concerning issue than specificity in ductal lavage. Until breast carcinogenesis theory is elucidated further, what actions to take in response to abnormal findings remains unclear.

Nipple aspiration is less invasive than ductal lavage; however, the number of cells available for study from aspiration is limited. Dooley et al. (2001) compared cellular yield between ductal lavage and nipple aspiration and found a significant difference (13,500 cells versus 120 cells, respectively). Additional studies have found that cytologic evaluation of nipple aspiration is not useful given its low predictive value (Krishnamurthy et al., 2003; Shao & Nguyen, 2001). The authors speculated that if breast cancer was present, the ducts probably were obstructed and cancer cells would not be aspirated. Because the precise precursors to carcinogenesis have not been defined clearly, searching for more accurate tumor markers is recommended as a priority.

FNAC is associated with a high rate of accuracy under optimal circumstances (Barrows et al., 1986). A study of 1,158 FNAs concluded that the procedure is sensitive and specific.
<table>
<thead>
<tr>
<th>Method</th>
<th>Description</th>
<th>Pros</th>
<th>Cons</th>
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| Nipple aspiration   | Use of simple suction technique employing a handheld device; droplets of nipple fluid are collected via capillary tube for analysis. | • Completely noninvasive  
• Inexpensive  
• Can be done by any trained healthcare professional  
• Can be collected outside the clinical setting | • Ability to collect fluid depends on ability of healthcare professional if woman has secreting ducts.  
• Fewer cells are available for cytologic diagnosis compared to ductal lavage. |
| Ductal lavage       | Use of microcatheter to cannulate ductal orifices; saline wash removes cells in collection container for analysis. | • Performed with a topical anesthetic only  
• Yields large number of cells for analysis | • More invasive than nipple aspiration  
• Low risk of infection or injury to the breast  
• Not all ducts are sampled. |
| Periareolar fine-needle aspiration | Use of a small needle to remove cells from the breast tissue for analysis | • Do not need intact ductal system to obtain cells for analysis | • Invasive procedure  
• Accuracy of readings  
• Depends on experience of healthcare professional performing procedure |

When used to evaluate clinically suspicious breast masses (Ariga et al., 2002). In groups of women divided by age (<40 years and younger versus 41 years and older), sensitivity was 99% and 98% and specificity 99% and 97%, respectively. Having established a cytologic and histologic correlation in FNA, its usefulness as a risk assessment tool is being studied (Fabian et al., 2000).

Sensitivity and specificity traditionally have been used as markers to evaluate the accuracy of a diagnostic tool. These evaluation standards are not applied easily to the use of breast epithelial cells as markers of breast cancer risk versus as markers of actual breast cancer. An important distinction must be made between using breast epithelial cells for the purpose of diagnosis versus the use of the cells as a measure of risk assessment. At the present time, these cells are best used as an enhancement to risk assessment, not as an independent diagnostic tool. Therefore, measures of sensitivity and specificity must be defined in relation to the risk assessment goals of breast epithelial cell evaluation.

### Using Ductal Fluid to Explore Carcinogenesis

The paths to carcinogenesis appear to be varied and numerous. Only by viewing the process as a work in progress will researchers develop interventions that may allow for true cure or prevention. As the majority of breast cancer cases are not the result of known germline mutations, an understanding of the genetic and epigenetic events that lead to malignancy is necessary to further the creation of new treatment modalities. This understanding may be advanced by viewing cells to sort out true precursors from benign changes. Access to breast epithelial cells via the nipple orifices or through periareolar FNA is pivotal for studying women who have developed breast cancer as well as those who have not developed it. Perhaps the study of changes in breast epithelial cells over time will allow researchers to begin to specify when premalignant changes take place and the events related to those changes. The methods outlined in this article for obtaining breast epithelial cells may determine when proliferative cells progress to something more ominous or regress back to normal. The carcinogenic continuum may be illuminated by viewing cytologic or molecular changes over time that are correlated with cancer development.

Reevaluating the use of current breast cancer risk assessment models by incorporating a more biologic component may enable healthcare professionals to more accurately assess risk. Nipple aspiration and ductal lavage are important adjuvants to risk assessment that could be performed easily in an outpatient setting. RNs and advanced practice nurses who work in the area of breast cancer risk assessment could perform these procedures safely and competently and inform patients regarding results in the context of individual risk assessment. Currently, nurse practitioners are trained by surgeons to perform ductal lavage and nipple aspiration. Institution-specific protocols are developed jointly by nurse practitioners and surgeons and guide practice. The skill set required is similar to that of placing an IV catheter.

Nipple aspiration, ductal lavage, and periareolar FNA are tools that hold great promise for exploring the breast carcinogenesis process. Through the observation of cellular and molecular abnormalities, opportunities for intervening in carcinogenesis will be revealed.

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### References


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**Oncology Nursing Forum – Vol 32, No 1, 2005**
CONTINUING EDUCATION

Strengths and Limitations of Breast Cancer Risk Assessment

Kimberly Baltzell, RN, PhD(c), and Margaret R. Wrensch, PhD

Purpose/Objectives: To evaluate current definitions of breast cancer risk and breast cancer risk assessment models, including the Gail, Claus, and BRCAPRO models, and discuss potential markers to enhance and standardize individual risk assessment.

Data Sources: Published articles, conference proceedings, and textbooks.

Data Synthesis: Defining high risk for breast cancer development is explored, and options for high-risk women are discussed. The risk factors frequently used for risk evaluation, including age, age at menarche, age at first live birth, past history of breast biopsy, family history of breast cancer, and the presence of atypical hyperplasia, are reviewed.

Conclusions: Current models of breast cancer risk assessment are limited. Exploiting the progression from healthy tissue to malignancy through techniques such as fine needle aspiration, ductal lavage, and nipple aspiration may lead to more precise individualized risk prediction.

Implications for Nursing: More accurate information regarding personal breast cancer risk is necessary. Oncology nurses may facilitate the use of appropriate tools that provide the most individualized risk assessment.

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cear of developing breast cancer is well founded among women in the United States. Breast cancer is the leading cause of death among women aged 35–50 years and the second-leading cause of death in women older than 50 years (Jemal et al., 2005). Approximately 40,000 women will die from this disease in the United States in 2005. Refining the science of breast cancer risk assessment has become more important with the availability of genetic testing for mutations associated with an increased risk of breast cancer development and the manufacture of medications to reduce breast cancer risk (Hollingsworth, Nall, & Dill, 2002).

A standardized algorithm for breast cancer risk assessment is not available at this time in the clinical setting. Women are categorized as either having possible genetic or hereditary risk or as having risk factors unrelated to a family history of breast cancer. Genetic testing is limited as a risk assessment tool because only a small percentage of women carry known genetic mutations that result in an increased risk of breast cancer development. Mathematical models calculate probabilities of developing breast cancer over specified periods of time; however, the factors included in the models contribute a relatively small degree of risk for the eventual development of breast cancer. Hollingsworth et al. (2002) suggested that

Key Points . .

- Assessing individual breast cancer risk has not been articulated in the United States despite an abundance of research devoted to risk factors.
- Currently employed risk assessment tools include the Gail model, the Claus model, and BRCAPRO.
- Exploring biologic markers such as atypical hyperplasia using minimally invasive methods (e.g., fine needle aspiration, ductal lavage, nipple aspiration) may enhance risk prediction.

Goal for CE Enrollees:

To enhance nurses’ knowledge about breast cancer risk factors, risk assessment models, and potential areas for refinement.

Objectives for CE Enrollees:

1. Summarize the impact of known risk factors on the development of breast cancer.
2. Discuss the strengths and limitations of currently used breast cancer risk assessment models.
3. Describe the potential role of pathologic information in more precisely determining breast cancer risk.

Kimberly Baltzell, RN, PhD(c), is a doctoral candidate in the Department of Physiological Nursing and Margaret R. Wrensch, PhD, is a professor in the Department of Epidemiology and Biostatistics and the Department of Neurological Surgery, both at the University of California, San Francisco. (Submitted December 2003. Accepted for publication July 18, 2004.)

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tissue- or serum-based strategies should be the next step in refining risk assessment, given that 70% of women who develop breast cancer have no identifiable risk factors.

Addressing inadequacies in breast cancer risk assessment may help to illuminate warning signs to women and healthcare providers as to who is at greatest risk for breast cancer development. This article will discuss risk assessment currently undertaken using the Gail and Claus models. In addition, the BRCAPRO program for assessing the probability of having known breast cancer genetic mutations will be discussed. Significant risk factors used in the clinical setting to determine risk will be outlined, as well as prevention options available to women deemed high risk. Abnormal epithelial breast cell cytology will be discussed as a potentially important risk factor to enhance current prediction models.

The Concept of High Risk

Defining High Risk

When is a woman at high risk for developing breast cancer? The generally agreed-upon risk factors currently used in various combinations in risk assessment models include being older than 65 years, experiencing early menarche (before 12 years of age), being nulliparous or having a first child after age 30, having a history of breast biopsy, and having a family history of breast cancer (Singletary, 2003). Radiation exposure at a young age (i.e., < 12 years) or as a treatment for Hodgkin disease also is associated with a higher risk of breast cancer development; however, it is not used as a risk factor in current risk assessment models (Clemens, Loijens, & Goss, 2000). The presence of atypical hyperplasia in breast tissue or fluid samples as a risk marker has shown significance in several studies (Fabian et al., 2000; Wrench et al., 2001). Various techniques to obtain this finding through histology and cytology have been discussed in greater detail in another article (Baltzell, Eder, & Wrench, 2005). Other factors contributing smaller degrees of risk for breast cancer development include drinking more than two alcoholic beverages per day, having a high body mass index in women older than 55 years, using hormone replacement therapy, and experiencing menopause after 55 years of age. Singletary succinctly listed the risk factors for breast cancer development (see Table 1). As more of these risk factors are present, the chance of developing breast cancer increases. The presence of a mutated BRCA1 or BRCA2 gene is currently the generally agreed-upon definition of high risk for breast cancer development. Multiple first-degree relatives with breast cancer and no mutated BRCA1 or BRCA2 gene in a woman's family history may suggest high-risk status, perhaps related to unknown genetic mutations.

If high risk was defined as a woman who has risk factors carrying a relative risk of greater than 2 (relative risk is the ratio of breast cancer risk among women with identified risk factors to the risk of breast cancer among women without those identified risk factors), then risk factors such as age, past personal history of breast cancer, lobular carcinoma in situ (LCIS), ductal carcinoma in situ (DCIS), biopsy findings of hyperplasia with atypia, atypia with a positive family history of breast cancer, first-degree relative with premenopausal breast cancer, more than two first-degree relatives with breast cancer, and known BRCA1 or BRCA2 mutations would provide information correlated with high risk. However, the majority of women seen in the clinical setting will not have information about their cellular or genetic risk factors (i.e., LCIS, DCIS, hyperplasia with atypia, BRCA1 and BRCA2 mutations). Obtaining information about these cellular or genetic risk factors may lead to a more concise and accurate definition of "high risk."

Accurate risk assessment is becoming increasingly important as potential prevention options, particularly prophylactic surgery and chemoprevention (Singletary, 2003), become available; however, these options are accompanied by their own set of risks. A decision to proceed with prophylactic surgery or chemoprevention should be made with as precise an assessment as possible. Because each of the currently available assessment tools uses different variables to assess risk, a precise definition is elusive. According to Verp, Cummings, and Olapade (2001), most cancers develop as a result of a combination of genetic and environmental factors. Despite years of research dedicated to articulating the risk factors leading to breast cancer development, no model completely calculates a woman's risk with great accuracy, with the exception of genetic testing indicating the presence of a BRCA1 or BRCA2 mutation (Winer, Morrow, Osborne, & Harris, 2001). Even genetic testing models are limited, given that they are based on very few of the possible mutations that increase breast cancer risk and are only definitive in families in which these mutations have been demonstrated (Berry et al., 2002).

Hamolsky and Facione (1999) described the importance of assisting women in making realistic appraisals of their personal risks. They reported that breast cancer risk estimations are misleading for many women because each woman has her own unique circumstances. According to Kelly (2000), although most women have beliefs regarding the cause of breast cancer, not all of those beliefs fit with current scientific findings. Women consistently overestimate their risk of developing breast cancer, which can lead to screening avoidance and psychological morbidity (Armstrong, Eisen, & Weber, 2000; Black, Nease, & Tofteson, 1995). Not every woman who has all of the currently recognized risk factors will develop breast cancer; therefore, more accurate risk assessment tools must be developed. Given that prophylactic surgery or chemopreventive drugs are the currently available breast cancer prevention choices, a woman must feel confident that her risk assessment is as complete as possible.

Breast Cancer Prevention Options

In the clinical setting, a limited number of breast cancer prevention options are available for women determined to be at extremely high risk for developing breast cancer (i.e., BRCA1 or BRCA2 mutations, a strong family history of breast cancer in first-degree relatives). These options include prophylactic surgery, chemopreventive drugs, and lifestyle modifications. If an extensive family history of breast cancer is found, genetic counseling or testing, if appropriate, should be offered to ascertain whether a BRCA1 or BRCA2 mutation is present. Although high penetrance genes are thought to account for only 10%–20% of breast cancers, the risk of developing breast cancer in the presence of these genes is high (Hamolsky & Facione, 1999).

Prophylactic mastectomy is associated with a risk reduction of more than 90% in women with strong family histories of breast cancer (Hartmann et al., 1999). The risk reduction associated with this procedure was similar for women with a strong family history and a subset of women with positive
Table 1. Risk Factors for Breast Cancer

<table>
<thead>
<tr>
<th>Risk Factor</th>
<th>Category at Risk</th>
<th>Comparison Category</th>
<th>Relative Risk</th>
</tr>
</thead>
<tbody>
<tr>
<td>Alcohol intake</td>
<td>2 drinks per day</td>
<td>Nondrinker</td>
<td>1.2</td>
</tr>
<tr>
<td>Body Mass Index</td>
<td>80th percentile, age 55 or greater</td>
<td>20th percentile</td>
<td>1.2</td>
</tr>
<tr>
<td>Hormone replacement therapy with estrogen and progesterone</td>
<td>Current user for at least 5 years</td>
<td>Never used</td>
<td>1.3</td>
</tr>
<tr>
<td>Radiation exposure</td>
<td>Repeated fluoroscopy</td>
<td>No exposure</td>
<td>1.6</td>
</tr>
<tr>
<td></td>
<td>Radiation therapy for Hodgkin’s disease</td>
<td>No exposure</td>
<td>5.2</td>
</tr>
<tr>
<td>Early menarche</td>
<td>Younger than 12 years</td>
<td>Older than 15 years</td>
<td>1.3</td>
</tr>
<tr>
<td>Late menopause</td>
<td>Older than 55 years</td>
<td>Younger than 45 years</td>
<td>1.2–1.5</td>
</tr>
<tr>
<td>Age at first childbirth</td>
<td>Nulliparous or 1st child after 30</td>
<td>1st child before 20</td>
<td>1.7–1.9</td>
</tr>
<tr>
<td>Current age</td>
<td>65 or older</td>
<td>Less than 65</td>
<td>5.8</td>
</tr>
<tr>
<td>Past history of breast cancer</td>
<td>Invasive breast carcinoma</td>
<td>No history of invasive breast carcinoma</td>
<td>6.6</td>
</tr>
<tr>
<td>Other histologic findings</td>
<td>Lobular carcinoma in situ</td>
<td>No abnormality detected</td>
<td>16.4</td>
</tr>
<tr>
<td></td>
<td>Ductal carcinoma in situ</td>
<td>No abnormality detected</td>
<td>17.3</td>
</tr>
<tr>
<td>Breast biopsy</td>
<td>Hyperplasia without atypia¹</td>
<td>No hyperplasia</td>
<td>1.9</td>
</tr>
<tr>
<td></td>
<td>Hyperplasia with atypia</td>
<td>No hyperplasia</td>
<td>5.3</td>
</tr>
<tr>
<td></td>
<td>Hyperplasia with atypia and positive family history</td>
<td>No hyperplasia, negative family history</td>
<td>11.0</td>
</tr>
<tr>
<td>Cytology (fine-needle aspiration, nipple aspiration fluid)</td>
<td>Proliferation without atypia¹</td>
<td>No abnormality detected</td>
<td>2.5</td>
</tr>
<tr>
<td></td>
<td>Proliferation with atypia</td>
<td>No abnormality detected</td>
<td>4.9–5.0</td>
</tr>
<tr>
<td></td>
<td>Proliferation with atypia and positive family history</td>
<td>No abnormality detected</td>
<td>8.1</td>
</tr>
<tr>
<td>Family history</td>
<td>1st-degree relative 50 years or older with postmenopausal breast cancer</td>
<td>No 1st- or 2nd-degree relative with breast cancer</td>
<td>1.8</td>
</tr>
<tr>
<td></td>
<td>1st-degree relative with premenopausal breast cancer</td>
<td>No 1st- or 2nd-degree relative with breast cancer</td>
<td>3.3</td>
</tr>
<tr>
<td></td>
<td>2nd-degree relative with breast cancer</td>
<td>No 1st- or 2nd-degree relative with breast cancer</td>
<td>1.5</td>
</tr>
<tr>
<td></td>
<td>Two 1st-degree relatives with breast cancer</td>
<td>No 1st- or 2nd-degree relative with breast cancer</td>
<td>3.6</td>
</tr>
<tr>
<td>Germ line mutation</td>
<td>Heterozygous for <strong>BRCA1</strong>, age &lt; 40</td>
<td>Not heterozygous for <strong>BRCA1</strong>, age &lt; 40</td>
<td>200.0</td>
</tr>
<tr>
<td></td>
<td>Heterozygous for <strong>BRCA1</strong>, age 60–69</td>
<td>Not heterozygous for <strong>BRCA1</strong>, age 60–69</td>
<td>15.0</td>
</tr>
</tbody>
</table>

¹ There is controversy over whether pathologic hyperplasia detected in breast biopsy samples is directly equivalent to cytologic hyperplasia detected in samples obtained through FNA (fine-needle aspiration) or nipple aspiration.

² Begg (2002) has suggested that these relative risks are subject to ascertainment bias and may overestimate the true risk associated with germ line mutations in BRCA genes.


**BRCA1** and **BRCA2** mutations. Although genetic testing is not suggested routinely for screening, a detailed family history indicating many relatives with breast or ovarian cancers may warrant offering genetic counseling. If a woman is found to be positive for genetic alterations of genes **BRCA1** or **BRCA2**, prophylactic mastectomy may be recommended. Love, Newcomb, and Trentham-Dietz (2002) recognized the magnitude of suggesting such a prevention strategy by stating, "In the absence of clinically applicable comprehensive risk models for individual patients, indications for prophylactic mastectomy must be strong and specific" (p. 210).

The removal of a woman's ovaries, or prophylactic oophorectomy, has been effective in reducing breast cancer risk in women with a known **BRCA1** or **BRCA2** mutation. Removing the ovaries in premenopausal women diminishes the amount of estrogen circulating that can stimulate breast cancer cells. When this source of estrogen is eliminated in women with genetic mutations known to increase risk of breast cancer development, risk has been reduced by approximately 50% (Olopade & Artioli, 2004).

Chemoprevention is described as "the use of specific natural and synthetic chemical agents to reverse or suppress carcinogenesis and prevent the development of invasive cancer" (Hamolsky & Facchine, 1999, p. 427). At present, the agents used for chemoprevention are a group known as selective estrogen receptor modulators (SERMs). Tamoxifen is the most widely prescribed SERM, and raloxifene is currently being evaluated for its effectiveness in preventing breast cancer development. SERMs act as estrogen agonists in some tissues (e.g., bone, endometrial) and as estrogen antagonists in other tissues (e.g., breast) (Brinton, Lacey, & Devesa, 2002). In the National Surgical Adjuvant Breast and Bowel Project (NSABP), a 49% lower risk of breast cancer was found in a tamoxifen-treated group versus a placebo-treated group (Fisher et al., 1998). Differences were apparent in groups within various studies; in a trial at the Royal Marsden Hospital, Eeles and Powles (2000) found that
SERMs were less effective in women with BRCA1 and BRCA2 mutations. Fisher et al. reported that the greatest risk reduction was in women with atypical hyperplasia. Risks associated with taking SERMs include stroke, deep vein thrombosis, and uterine cancer. Brinton et al. noted that although the overall results of SERM trials are informative, the analyses are less useful to individuals and their clinicians trying to make informed decisions regarding the appropriateness of this prevention strategy. That is, clinical guidelines are not yet clear about the recommendation of SERMs for breast cancer prevention.

Lifestyle changes have been examined in an effort to determine which may modify breast cancer risk. Dietary fat has been studied extensively as a risk factor for breast cancer development. According to Kushi and Giovannucci (2002), recommendations to reduce fat intake to prevent cancer risk are unwarranted. Drake (2001) reported that female joggers were less likely to develop breast cancer than those who did not jog. In another study, lifelong physical activity was potentially useful in reducing breast cancer risk (Bernstein, Henderson, Hanisch, Sallivan-Halley, & Ross, 1994). Physical activity in young women is associated with delayed menarche and anovulatory cycles, perhaps reducing overall lifetime exposure to estrogen. Although studies have not found a highly significant association between lifestyle variables and breast cancer prevention, a reduced-fat diet and increased exercise may be beneficial in regard to other diseases (e.g., cardiovascular disease). Love et al. (2002) created a table of possible primary prevention strategies categorized by age group (see Table 2). These interventions revolve around the idea that breast tissue development and the role of hormonal changes leading to breast cancer susceptibility but do not necessarily include truly feasible or desirable modifications or programs for women. To recommend breast cancer prevention strategies, a comprehensive breast cancer risk assessment is necessary.

**Risk Factors**

Age, age at menarche, age at first live birth, family history of breast cancer, past history of breast biopsy, and the presence of atypical hyperplasia are risk factors that can be taken into account when assessing breast cancer risk. Table 3 summarizes the potential modifiability of these risk factors.

**Age**

Of all the commonly used risk factors to predict breast cancer, increasing age is believed to have the most significance (Winer et al., 2001). In more than 50% of women diagnosed with breast cancer, increasing age is the only identifiable risk factor (Madigan, Ziegler, Benichou, Byrne, & Hoover, 1995). Risk of breast cancer development increases steadily until age 70, at which point risk actually declines (Kelly, 2000). The commonly quoted 1 in 8 risk is derived from the addition of age-stratification risk numbers. Women aged 20–30 years have a 2% risk of breast cancer development (1 in 50), women aged 50–70 years have a 6% risk of breast cancer development (1 in 17), and women aged 70–80 years have a 3% risk (1 in 33) (Kelly). These are generalized risk numbers that cannot be used effectively for individual risk assessment. In nonhereditary breast cancers, the increased risk of breast cancer with advancing age may come more from “wear and tear” on genetic material, providing an opportunity for mutations to occur or from decreased immune surveillance. Recent statistics are listed in Table 4 and show the increased number of diagnoses as women age (Jemal et al., 2005).

**Age at Menarche**

Risk assessment often categorizes age at menarche as less than 12 years or more than 15 years, representing higher versus lower risk, respectively. If lifetime exposure to estrogen is associated with risk determination for breast cancer, then the number of actual cycles an individual has provides important estrogen exposure information. Age at menarche has received more attention in recent years because of observations of earlier onset of puberty in the United States (Lee, Guo, & Kulik, 2001). The combinations of higher fat and protein diets and effective disease control are believed to have had an impact on lowering the age of menarche (Henderson, Pike, Bernstein, & Ross, 1996). MacDonald et al. (1982) reported that establishment of ovulatory cycles and increased hormone levels found in women who experienced early menarche play a role in promoting breast cancer risk. Henderson et al. suggested that for women of equivalent age, those with more than 40 years of menstruation have twice the risk of those with fewer than 30 years of menstruation. Strategies for decreasing risk may
Table 3. Summary of Risk Factor Modification Feasibility

<table>
<thead>
<tr>
<th>Risk Factors</th>
<th>Risk Modifiable?</th>
<th>Risk Modifiable at Age of Concern</th>
<th>Advantages</th>
<th>Disadvantages</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age</td>
<td>No</td>
<td>No</td>
<td>Not applicable</td>
<td>Adolescence is the time of increased body image distortion and onset of eating disorders. The effect on other disease development is unknown.</td>
</tr>
<tr>
<td>Age at menarche</td>
<td>Possibly</td>
<td>No</td>
<td>Encouragement of increased exercise and lifelong healthy habits</td>
<td></td>
</tr>
<tr>
<td>Age at first live birth</td>
<td>Yes</td>
<td>No</td>
<td>Could confer a protective period postpregnancy at critical time for breast carcinogenesis</td>
<td>Economic instability associated with young maternal age may create other health issues that are more threatening than breast cancer development.</td>
</tr>
<tr>
<td>Past history of breast biopsy</td>
<td>Partially</td>
<td>No</td>
<td>Obtain information related to high-risk cellular abnormalities via less invasive methods (e.g., fine needle aspiration, nipple aspirate fluid, lavage), Less invasive methods are not commonly practiced; accurate pathology reading is crucial for risk information.</td>
<td></td>
</tr>
<tr>
<td>Family history of breast cancer</td>
<td>No</td>
<td>No</td>
<td>Not applicable</td>
<td></td>
</tr>
<tr>
<td>Atypical hyperplasia</td>
<td>Unknown</td>
<td>Possibly</td>
<td>Not applicable</td>
<td></td>
</tr>
</tbody>
</table>

* Age of concern is defined as the age at which risk for breast cancer development increases significantly. For purposes of this table, age 40 begins the “age of concern” based on the probability increase from 1 in 228 (age birth to 39) to 1 in 24 (age 40–59).

* Patrikis et al. (1986) found an increase in cytologic detection of epithelial hyperplasia in breast fluids after increased consumption of soy protein in a small study of women aged 30–59. This indicates the possibility of exogenous influences in altering the progression of atypical hyperplasia.

include looking at adolescence as an effective intervention age. Encouraging increased amounts of exercise and healthy eating habits may influence menarche onset by a small margin; however, each year of menarche delay may provide a significant decrease in later breast cancer risk. In addition to the benefit of fewer menstrual cycles resulting in decreased estrogen exposure in the breast tissue, exercise and healthy eating may contribute to decreased weight gain in adulthood. Adipose tissue is a major source of estrogen in postmenopausal women. Weight loss and low body mass index are associated with a decreased risk of breast cancer in postmenopausal women; however, this type of advice should be given cautiously. Recommending “thinner” to an adolescent girl may be associated with the development of eating disorders such as anorexia nervosa and bulimia (Martin & Ammerman, 2002). In addition, the burden of possible breast cancer development should not be added to adolescent worries, particularly if the timing of menarche can be altered only by radical shifts in lifestyle.

**Age at First Live Birth**

Chie et al. (2000) compared age at first pregnancy for breast cancer cases and controls and found a modest increased risk in breast cancer development (odds ratio = 1.07, confidence interval = 1.01–1.13) for each five-year increase in age at first full-term pregnancy. MacMahon et al. (1970) reported that women with their first full-term pregnancy before age 20 had a third of the breast cancer risk compared with women having their first full-term pregnancy after age 35. A short-term increased risk of breast cancer development may occur after pregnancy at any age; however, mammary cells become differentiated after this risk period, resulting in less susceptibility to carcinogenesis. This increased risk period is believed to last approximately 10 years (Bruzzi et al., 1988). An early pregnancy allows for mammary cell differentiation at an early age in a woman’s reproductive life, perhaps conferring a protective effect during later high-risk years. Brinton et al. (2002) found the protective effect of early pregnancy only with full-term pregnancy. Singletary (2003) suggested that this is because of cell differentiation in preparation for lactation in the later stages of pregnancy. Brinton et al. also reported that multiparous women and women who give birth around age 30 share a similar risk of breast cancer development. A full-term pregnancy after age 30 is associated with higher risk than nulliparity, possibly as a result of the increased risk period immediately after pregnancy. Brinton et al. speculated that already initiated cells may progress during the short-term high-risk period following late-age pregnancy. Because the protective effect of pregnancy is associated with maternal age of less than 20 years of age, it is unlikely to be a risk factor that is altered easily. However, the social trend toward later maternal age at pregnancy is continuing in North American societies (Lee et al., 2003), but changing reproductive choices, as suggested by Love et al. (2002), is unrealistic in any risk intervention strategy.

**Table 4. Advancing Age and Corresponding Increase in Breast Cancer Rates**

<table>
<thead>
<tr>
<th>Age (Years)</th>
<th>% Diagnosed With Breast Cancer</th>
<th>Actual Number of Cases per Interval</th>
</tr>
</thead>
<tbody>
<tr>
<td>0–39</td>
<td>0.4</td>
<td>1 in 228</td>
</tr>
<tr>
<td>40–59</td>
<td>4.0</td>
<td>1 in 24</td>
</tr>
<tr>
<td>60–79</td>
<td>7.0</td>
<td>1 in 14</td>
</tr>
<tr>
<td>Lifetime</td>
<td>12.0</td>
<td>1 in 8</td>
</tr>
</tbody>
</table>

*With each age interval passed without a breast cancer diagnosis, risk for that category should be subtracted from subsequent age intervals (Kelly, 2000).

Note: Based on information from Jemal, 2005.

**Past History of Breast Biopsy**

According to Page et al. (1978), women with a history of breast biopsy have an elevated risk of approximately twice the general population for future breast cancer development. This
is because of the underlying presence of benign breast disease, which has been found to be significantly associated with breast cancer development (Webber & Boyd, 1986). Breast biopsy history has been included in the Gail risk model as an important risk factor. Kelly (2000) argued against using the number of biopsies in a risk model because some, but not all, benign breast disease leads to biopsy, limiting its usefulness as a risk marker. Hughes, Mansel, and Webster (2000) wrote, "There is no reason to believe that the clinical presentations that induce a surgeon to perform a biopsy will be associated with high-risk pathology as most of the hyperplastic lesions with atypia are found incidentally at biopsy for a condition such as dominant nodularity" (p. 255). Is the fact that a woman had a biopsy important in risk assessment? Using the actual results of the biopsy may be more informative, but only if hyperplasia or atypical hyperplasia is present. Page et al. investigated the link between histologic changes present in breast tissue and breast cancer risk and concluded that benign breast disease is not necessarily associated with increased cancer risk; however, histologic changes defined as epithelial proliferative disease may distinguish high-risk groups from women with general population risk. Winer et al. (2001) noted that most breast biopsies result in nonproliferative disease findings. Using the number of biopsies in a risk model would lead to an overestimation of risk based on this information. Refining the concept of breast biopsy numbers is necessary for value in clinical decision making. Suggesting biopsies for large populations of at-risk women is unrealistic and cost prohibitive. Determining the presence of abnormal proliferative changes through less invasive methods that may lead to biopsy might improve the prediction value and specificity of this factor. Perhaps the incorporation of pathology findings (via biopsy, fine needle aspiration, lavage, or nipple aspiration) is more essential for enhanced risk assessment.

**Family History of Breast Cancer**

A family history of breast cancer is associated with a significant increase in breast cancer risk; however, only 5%–10% of breast cancers are believed to have strong hereditary origins (Winer et al., 2001). In addition, Winer et al. wrote that "family history is a heterogeneous risk factor with different implications depending on the number of relatives with breast cancer, the exact relationship, the age at diagnosis, and the number of affected relatives" (p. 1652). A person with multiple relatives diagnosed with breast cancer at an early age is at greater risk than a woman with one relative diagnosed at a postmenopausal age. Kelly (2000) listed the following indications that hereditary cancers may be present: young age at diagnosis, one person diagnosed with several different cancers, cancers present in two or more generations, and three or more cancers found in close relatives. Complicating the family history is that shared environment might contribute to disease development in all family members, independently of any inherited genetic mutation.

Two tumor suppressor genes have been identified that are associated with true genetic risk of breast cancer development. Located on chromosome 17 is *BRCA1*, and on chromosome 13 is *BRCA2* (Winer et al., 2001). Mutations in either of these genes correlate with a 50%–85% lifetime chance of developing breast cancer. Additionally, these mutations can be passed down by either the mother or father. The large size of *BRCA1* and *BRCA2* makes genetic testing prohibitively expensive and unreasonable for large populations (Winer et al.). The cost of testing for a *BRCA* mutation was more than $2,500 in 2000 (Kelly, 2000). Also, all *BRCA1* and *BRCA2* mutations are not the same. Researchers have been unable to determine whether mutations in different locations on the gene convey the same level of risk. At this time, a positive genetic test means that a person might be at increased risk for breast cancer development; however, a negative test cannot rule out the possibility of another unknown mutation. Counseling a woman in regard to genetic testing involves a complex and complete screening process, including the discussion of breast cancer prevention strategies available in the event of a positive test. Other considerations regarding genetic counseling include the need for privacy and availability of qualified genetic counselors to guide future decisions affected by the presence of *BRCA1* and *BRCA2* mutations.

**Atypical Hyperplasia**

Recent studies have demonstrated a significant relationship between the presence of atypical hyperplasia in breast tissue or fluid samples and increased breast cancer risk (Fabian et al., 2000; Wrensch et al., 2001). Cytologic and histologic attributes associated with atypical hyperplasia include (a) an increase in cellular mitotic activity, (b) nuclear enlargement, (c) irregular nuclear borders, (d) nuclear hyperchromasia, (e) involvement of two or fewer ductal sections, and (f) foci measuring less than 2 mm (Rosen, 2001). Cells may be obtained by a number of methods, including breast biopsy, fine needle aspiration, ductal lavage, and nipple aspiration; however, results may vary based on the method of cell extraction chosen. Dupont and Page (1985) reexamined breast biopsies of 3,303 women after 17 years and found that women with atypical hyperplasia had a relative risk for invasive breast cancer of 5.3, with an increased relative risk of 11 for women with atypical hyperplasia and a positive family history. Inspired by an early study (Papanicolaou, Holmlund, Bader, & Falk, 1958), Sartorius, Smith, Morris, Benedict, and Friesen (1977) developed a nipple aspiration device to obtain breast fluid from 1,706 women. Fluid was obtained in approximately 50% of the cohort, and study results indicated a significant relationship between the presence of atypia and underlying breast cancer. Fabian et al. used fine needle aspiration to examine cells for the presence of atypical hyperplasia and determined that cytomorphologic findings of atypical hyperplasia are useful in evaluating short-term breast cancer risk. In several studies, abnormal cellular cytology in breast fluid was associated with an increased risk of breast cancer (Wrensch et al., 1992, 2001; Wrensch, Petrikis, King, Lee, & Miike, 1993). King, Chew, Petrikis, and Ernst (1983) documented the high correlation between atypical hyperplasia found in nipple aspirate fluid and atypical proliferative disease found in breast biopsy. This study confirmed the feasibility of using any of the available methods (biopsy, fine needle aspiration, ductal lavage, or nipple aspiration) to examine abnormalities associated with higher breast cancer risk. If cytologic and histologic methods of obtaining cells yield equally accurate information, choosing less invasive and costly procedures (e.g., fine needle aspiration, nipple aspiration) would allow for broader use of this marker for risk assessment. Dooley et al. (2001) concluded that ductal lavage is safe and well tolerated by most women, as well as a source of many breast epithelial cells for analysis. O'Shaughnessy (2001) stated that ductal lavage was a promising risk assessment tool. In addition, a number of breast cancer specialists recommended incorporating breast fluid findings into the breast cancer risk profile (Goodman, 2002).
Current Models of Breast Cancer Risk Assessment

Overview

For the purposes of this article, a breast cancer risk assessment model refers to mathematical models that calculate actual risk of breast cancer development as well as genetic tests (e.g., BRCAPRO) that examine known breast cancer gene mutations (e.g., BRCA1, BRCA2). The most commonly employed breast cancer risk assessment models currently are the Gail model and the Claus model (mathematical models) and BRCAPRO, which is used to evaluate the possible presence of genetic mutations associated with increased risk of breast cancer development. The Tyrer-Cuzick model has been developed to address concerns and limitations of currently used models. This model incorporates the likelihood of the presence of genes predisposing one to breast cancer, as well as personal risk factors (Tyrer, Duffy, & Cuzick, 2004). However, this model has not been validated independently (Amir et al., 2003). Euhus (2001) stated that an understanding of the principles used in each of these models is essential for healthcare professionals engaged in risk management counseling. MacDonald (2002) suggested that all healthcare providers will come in contact with a woman who has a family history of breast cancer at some point, given the prevalence of this disease. Risk assessment models are not used uniformly in clinical practice, making the accuracy of each woman’s risk assessment a function of her provider’s knowledge. Regarding healthcare providers, Kelly (2000) reported, “Many have a general knowledge of breast cancer risks, but few make it specialty, have the time to keep up with all the latest developments in this area, or are aware of all whose risk might be increased” (p. 174).

Gail Model

Gail et al. (1989) developed a mathematical model for risk assessment of invasive and in situ breast cancer using information from 284,780 Caucasian women participating in the Breast Cancer Detection Demonstration Project from 1973–1980. This was a first attempt to refine population characteristics and base risk assessment on subgroups of women with varying risk factors, including age, age at menarche, number of prior breast biopsies, age at first live birth, and number of first-degree relatives affected with breast cancer. Relative risk was calculated for each of these risk factors; those relative risks (i.e., the probability of developing breast cancer in a given population) then were used to calculate absolute risk at five years from the time of assessment and a lifetime risk up to the age of 90. This model has been modified to include African Americans as well as Caucasians and uses invasive cancer as the only defined “breast cancer event” (Euhus, Leitch, Huth, & Peters, 2002). In addition, the presence of atypical hyperplasia has been added as a risk factor (Euhus, Leitch, et al.). The modified Gail model was used to qualify women for enrollment eligibility by the NSABP to assess the effectiveness of tamoxifen in preventing breast cancer development. Women with a five-year Gail score of more than 1.7% were designated “high risk” and qualified for participation in the tamoxifen study. In addition, this model was used for selection of candidates for the Study of Tamoxifen and Raloxifene trial comparing the effectiveness of tamoxifen versus raloxifene (Euhus, 2001).

Strengths of the Gail model include its attempt to adapt risk assessment from the general population to be more applicable to specific subgroups. In a study by Euhus, Leitch, et al. (2002), the Gail model was used in specialized clinic settings, although it is criticized widely for not accounting for adequate family history information. The Gail model was developed prior to extensive genetic testing and now is thought to be most applicable to women without a strong family history suggestive of an inherited genetic mutation (Sakorafas, Krespi, & Pavlakis, 2002).

Criticisms of the Gail model are wide and varied, but it is limited by the characteristics of the data set used for its development. Kelly (2000) reported that the Gail model was problematic because (a) relative risk is not an accurate way to obtain absolute risk, (b) the number of biopsies included in the calculation is too simplistic (the pathology information obtained from the biopsy is more informative than the fact that a biopsy was performed), (c) all relevant family history is not included (i.e., grandparents and paternal history relatives are excluded), and (d) risk is overestimated in young women. Bondy and Newman (2003) found that the model has not been validated in African American women and stated their concern relative to enrollment and recruitment of African Americans in the ongoing NSABP trials. In addition to complaints regarding lack of validation for African Americans, no attempt has been made to validate the Gail model in other ethnic populations. The addition of atypical hyperplasia may enhance model accuracy; perhaps this would replace the number of biopsies with more useful biologic information.

Claus Model

In 1993, Claus, Risch, and Thompson published information on a model that incorporated extensive family history of cancer development. These data were obtained from the Cancer and Steroid Hormone Study, consisting of interviews of 4,730 confirmed breast cancer cases and 4,688 controls. The final model included breast cancer information on not only mothers and sisters but aunts and grandmothers as well. The development of the Claus model supported the notion that inherited genetic mutations might increase the risk of breast cancer and was a hint of a genetic component that would be elucidated further in the following five years (Euhus, 2001). The Claus model also addressed an inadequacy of the Gail model. The strength of the Claus model is its ability to incorporate the age of affected family members at diagnosis into the analysis. Since the discovery of BRCA1 and BRCA2 mutations, this information has taken on more importance, given that a woman with early onset of the disease is more likely to carry one of these mutations. However, the Claus model does have its own limitations: It does not include known breast cancer risk factors that are unrelated to family history of breast cancer, such as those included in the Gail model (Euhus). Therefore, the Claus model cannot be used among women without a family history of breast cancer. Because of the small sample size of African Americans in the original data set, final risk assessments did not include race. Other ethnicities were not addressed, probably because of the limited amount of information available for analysis. This model may be most helpful for women with a strong family history of breast cancer. Comparisons between the Gail and Claus model are shown in Table 5.
Table 5. Variables Used in the Gail and Claus Models

<table>
<thead>
<tr>
<th>Variable</th>
<th>Gail</th>
<th>Claus</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age</td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
<td>First-degree family history (i.e., mother, sisters, and daughters)</td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
<td>Second-degree family history (i.e., aunts and grandmothers)</td>
<td>No</td>
<td>Yes</td>
</tr>
<tr>
<td>Age at onset in relatives</td>
<td>No</td>
<td>Yes</td>
</tr>
<tr>
<td>Age at menarche</td>
<td>Yes</td>
<td>No</td>
</tr>
<tr>
<td>Age at first live birth</td>
<td>Yes</td>
<td>No</td>
</tr>
<tr>
<td>Number of breast biopsies</td>
<td>Yes</td>
<td>No</td>
</tr>
<tr>
<td>Atypical hyperplasia</td>
<td>Yes</td>
<td>No</td>
</tr>
<tr>
<td>Race and ethnicity</td>
<td>Yes</td>
<td>No</td>
</tr>
</tbody>
</table>

Note: Based on information from McTiernan et al., 2001.

**BRCA PRO**

Unlike the Gail and Claus models of breast cancer risk assessment, BRCA PRO is used to determine the probability of having a genetic mutation (specifically BRCA1 or BRCA2) associated with an increased risk of developing breast cancer. Although other genetic risk models exist, BRCA PRO is considered the most comprehensive (Allman, Gilligan, & Redlich, 2005). It is described as mathematically “intense” and uses Bayes theorem to answer the questions: “Given this pattern of affected and unaffected relatives, what is the probability that this individual carries a mutation in one of the BRCA genes? Given this BRCA gene mutation probability, what is the probability that this individual will develop breast cancer?” (Euhus, 2001, p. 228). The reliability of the calculation grows as more information is added to the model about the age and history of relatives with breast and ovarian cancer. Euhus wrote that the key to the usefulness of this model lies in knowing the underlying frequency of mutated genes in the population to which a patient belongs (e.g., European American, Eastern European Jewish).

BRCA PRO was found to be relatively accurate in predicting the presence of BRCA mutations in samples where the probability of penetrance was either very high (> 95%) or very low (< 5%) (Berry et al., 2002). BRCA PRO is a sensitive tool, missing only 15% of mutations present; however, Berry et al. did not determine whether this tool is useful in predicting which mutation carriers will develop breast cancer. Additional studies found that BRCA PRO more accurately identified possible mutations than experienced risk counselors (Euhus, Smith, et al., 2002). Limitations of the model include its underestimation of women’s risk when familial clustering is unrelated to BRCA gene mutation (Euhus, 2001). Allain et al. (2002) listed lack of verification of family history as another limitation of this tool. BRCA PRO does not evaluate risk factors unrelated to family history (e.g., reproductive risk factors, presence of atypical hyperplasia). See Table 6 for a comparison of the three breast cancer risk assessment models.

**Using Atypical Hyperplasia to Enhance Assessment Models**

Most women who develop breast cancer do not have a known genetic mutation that indicates increased risk for the disease. How can more specific biologic information be obtained to refine breast cancer risk assessment? Perhaps examining breast epithelial cells (via lavage, nipple aspirate fluid, or periareolar fine needle aspiration) will illuminate cellular changes leading to cancer development. Daly and Ross (2000) stated that an understanding of the biologic progression from healthy breast epithelium to malignancy has been impeded by a lack of access to at-risk tissue for surveillance. Studies show atypical hyperplasia’s contribution to increased risk in breast cancer development to be four- to fivefold in atypical hyperplasia, rising to anywhere from 11- to 18-fold in women with atypical hyperplasia and family history of breast cancer (Dupont & Page, 1985; Singletary, 2003). These relative risks are higher by a substantial margin than relative risks of currently accepted breast cancer risk factors such as age at menarche or age at first pregnancy. Increased emphasis should be placed on obtaining biologic markers of breast cancer risk that will allow for more accurate assessment of who is truly at risk for disease development. O’Shaughnessy (2001) wrote that more specific tools, such as ductal lavage to obtain cytologic information, are necessary to stratify women into useful risk assessment categories. Promising studies indicate that evaluating breast epithelium may yield important clues as to who may be at great risk for breast cancer (Fabian et al., 2000; Wrensch et al., 2001). This additional to risk assessment has become more feasible because data from less invasive means (nipple aspiration) provide

Table 6. Advantages and Disadvantages of the Gail, Claus, and BRCA PRO Models

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>Gail</th>
<th>Claus</th>
<th>BRCA PRO</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Advantages</strong></td>
<td>Accurately predicts the number of expected cases of breast cancer in large-scale clinical trials; incorporates nonfamily risk factors</td>
<td>Uses information from first- and second-degree relatives; incorporates age at diagnosis of affected family members</td>
<td>Most comprehensive estimate of genetic mutation risk; highly sensitive</td>
</tr>
<tr>
<td><strong>Disadvantages</strong></td>
<td>All relevant family history of breast cancer is not included; the model may overestimate risk in young women.</td>
<td>Does not include breast cancer risk factors other than family history</td>
<td>Underestimates risk in women with familial clustering unrelated to BRCA1 and BRCA2 mutations; does not evaluate risk factors unrelated to family history of breast cancer</td>
</tr>
<tr>
<td><strong>High-risk definition</strong></td>
<td>High risk is defined as a score of more than 1.7% within a five-year time period.</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Most appropriate population</strong></td>
<td>Women without a strong family history of breast cancer</td>
<td>Women with a strong family history of breast cancer</td>
<td>Women with a strong family history of breast or ovarian cancer</td>
</tr>
</tbody>
</table>
a degree of pathologic information on par with breast biopsy (King et al., 1985). In the past, cytologic information has been available only for a limited number of at-risk women, which has made the inclusion of atypical hyperplasia information sporadic in risk assessment models. Incorporating these findings into regular risk assessment may help to further specify who requires more aggressive, invasive follow-up. At present, assessment of atypical ductal hyperplasia may be one of the risk assessment tools with the most potential.

Conclusion

The mathematical Gail and Claus models may benefit from the addition of a serum- or tissue-based biomarker of breast cancer risk. As these models are used currently, certain women’s risk of breast cancer development may be overestimated or underestimated. Risk factors used in these models are largely unmodifiable, either practically or ethically. In addition, many of the risk factors used for assessment contribute very small relative risks, making their importance in risk models questionable. The definition of who is at high risk for breast cancer development should be expanded and articulated. The development of breast cancer prevention options makes this articulation even more critical. Fisher et al.’s (1998) conclusion that tamoxifen was most beneficial in women with atypical hyperplasia suggested an important link between cytologic findings and benefit from prevention strategies. Studying cytologic and histologic proliferative patterns such as atypical hyperplasia may lead to the next step in refining risk assessment.

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References


ONF Continuing Education Examination

Strengths and Limitations of Breast Cancer Risk Assessment

Credit Hours: 1.6
Passing Score: 80%
Test ID # 05-323-04
Test processing via ONS Web site: FREE
Test processing via mail-in form: $15

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1. Modification of breast cancer risk assessment techniques has become necessary because of
   b. Clearer delineation of the environmental causes of breast cancer.
   c. New interventions that must be used immediately upon diagnosis of breast cancer.
   d. Novel diagnostic techniques that carry a lower risk during the workup for breast cancer.

2. Currently, most women who develop breast cancer exhibit how many risk factors?
   a. 0
   b. 1
   c. 2–3
   d. 4 or more

3. When assessing a woman’s risk of developing breast cancer using current risk assessment models, which of the following would indicate increased risk?
   a. Menarche at 13 years of age
   b. History of radiation therapy for Hodgkin disease
   c. Being 55 years of age
   d. Never having had children

4. Currently, a woman is considered at high risk of developing breast cancer if she
   a. Used hormone replacement therapy.
   b. Carries a mutated BRCA1 or BRCA2 gene.
   c. Reached menopause after the age of 55.
   d. Has a history of undergoing breast biopsy.

5. When helping a woman at extremely high risk for developing breast cancer evaluate her options, which prevention option that is associated with the greatest reduction in this risk should be noted?
   a. Prophylactic oophorectomy
   b. Lifestyle changes
   c. Selective estrogen receptor modulator therapy
   d. Prophylactic mastectomy

6. For a woman with a strong family history of breast cancer, which breast cancer risk assessment model would be most appropriate to use?
   a. Study of Tamoxifen and Raloxifene
   b. Tyer-Cuzick
   c. Claus
   d. Gail

7. Which commonly used risk factor is believed to play the most significant role in the development of breast cancer?
   a. Family history of breast cancer
   b. Age at first live birth
   c. Personal history of breast biopsy
   d. Increasing age

8. A history of breast biopsy is considered a risk factor for developing breast cancer because
   a. Abnormal breast cells released during biopsy have the propensity to spread into local tissue.
   b. Benign breast disease that leads to biopsy is significantly associated with cancer development.
   c. Stress associated with breast biopsy procedures stimulates breast cell malignant transformation.
   d. The majority of breast biopsy results leads to findings of proliferative breast disease.

9. Which of the following methods for obtaining breast epithelial cells is most feasible for use in a large breast cancer screening program?
   a. Incisional biopsy
   b. Nipple aspiration
   c. Excisional biopsy
   d. Nipple scraping

10. The Gail breast cancer risk assessment model would be most appropriate for evaluating women
    a. With a family history of cancers in two or more generations.
    b. Across a wide variety of ethnic and minority groups.
    c. Who appear to exhibit several noninherited risk factors.
    d. Younger than 40 years of age and premenopausal.

11. For a woman with multiple family members diagnosed with breast and ovarian cancer, which assessment model would be most helpful in estimating her breast cancer risk?
    a. Gail
    b. Claus
    c. BRCA PRO
    d. Tyer-Cuzick

12. Which of the following factors has been found to most significantly increase a woman’s relative risk of developing breast cancer?
    a. Atypical hyperplasia
    b. Age at menarche
    c. Nulliparity
    d. History of breast biopsy
13. When developing a breast health educational program for adolescent girls, which recommendation would be most appropriate to include?
   a. Maintain a thin body through a high-protein diet.
   b. Plan to breastfeed any children for at least one year.
   c. Take a multivitamin with minerals every day.
   d. Regularly engage in enjoyable physical activity.

14. The breast cancer risk factor that currently shows the most potential in the refinement of risk assessment tools is
   a. Genetic mutations beyond BRCA1 and BRCA2.
   b. Atypical hyperplasia.
   c. Breast cell response to tamoxifen exposure.
   d. Number of breast biopsies.

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1. a 2. a 3. a 4. a 5. a 6. a 7. a 8. a 9. a 10. a
   b  b  a  a  b  b  b  b  b  b
   c  c  c  c  c  c  c  c  c  c
   d  d  d  d  d  d  d  d  d  d

11. a 12. a 13. a 14. a 15. a 16. a 17. a 18. a 19. a 20. a
    b  b  a  a  b  b  b  b  b  b
    c  c  c  c  c  c  c  c  c  c
    d  d  d  d  d  d  d  d  d  d

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**Program Evaluation**

1. How relevant were the objectives to the CE activity’s goal?

2. How well did you meet the CE activity’s objectives (see page 605)?
   - Objective #1
   - Objective #2
   - Objective #3

3. To what degree were the teaching/learning resources helpful?

4. Based on your previous knowledge and experience, do you think that the level of the information presented in the CE activity was
   Too basic  Appropriate  Too complex

5. How long did it take you to complete the CE activity? __________ minutes

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