Higher-status occupations and breast cancer: A life-course stress approach

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A B S T R A C T

Using the 1957–2011 data from 3682 White non-Hispanic women (297 incident breast cancer cases) in the Wisconsin Longitudinal Study, United States, we explore the effect of occupation in 1975 (at age 36) on breast cancer incidence up to age 72. Our study is motivated by the paradoxical association between higher-status occupations and elevated breast cancer risk, which presents a challenge to the consistent health advantage of higher social class. We found that women in professional occupations had 72122% higher incidence of breast cancer than women in lower-status occupations (Danø, 2003; Larsen et al., 2011). Moreover, the effect of higher-status occupations on breast cancer risk is only partly explained by reproductive histories, exogenous hormones, health behaviors, and socioeconomic differences in screening mammography (Danø et al., 2004; Larsen et al., 2011; Sprague, Trentham-Dietz, & Burnside, 2010). Because women with more socioeconomic resources are advantaged in terms of nearly all other health outcomes (Elo, 2009), the breast cancer risk associated with higher-status occupations presents a paradox that calls for particular attention to this disease. Existing research, however, is limited in several respects. Most studies were based on cross-sectional analyses of breast cancer rates across occupations or had a relatively short follow-up that did not capture the long latency period of breast cancer. Further, past studies considered a limited set of mediators potentially linking occupation to breast cancer and focused overwhelmingly on estrogen-related pathways, such as reproductive histories.

Introduction

Lower morbidity associated with higher socioeconomic status (SES) is one of the most consistent findings in social epidemiology (Elo, 2009), yet one paradox persists: the elevated risk of breast cancer among women holding higher-status occupations. Higher-status occupations are defined as professional and managerial occupations that are at the top of the U.S. Census classification system and are characterized by the highest levels of socioeconomic prestige indexes (Stevens & Cho, 1985). Women in professional and managerial occupations have 1.4–2.0 times greater risk of breast cancer diagnosis than women in lower-status occupations (Danø, 2003; Larsen et al., 2011; Pukkala et al., 2009). Moreover, the effect of higher-status occupations on breast cancer risk is only partly explained by reproductive histories, exogenous hormones, health behaviors, and socioeconomic differences in screening mammography (Danø et al., 2004; Larsen et al., 2011; Sprague, Trentham-Dietz, & Burnside, 2010).

Because women with more socioeconomic resources are advantaged in terms of nearly all other health outcomes (Elo, 2009), the breast cancer risk associated with higher-status occupations presents a paradox that calls for particular attention to this disease. Existing research, however, is limited in several respects. Most studies were based on cross-sectional analyses of breast cancer rates across occupations or had a relatively short follow-up that did not capture the long latency period of breast cancer. Further, past studies considered a limited set of mediators potentially linking occupation to breast cancer and focused overwhelmingly on estrogen-related pathways, such as reproductive histories. Because...
estrogen-related factors explain only part of the excess risk for breast cancer associated with higher-status occupations (Dano et al., 2004; Larsen et al., 2011), researchers need to continue the search for explanatory mechanisms, especially psychosocial stressors to which women in higher-status occupations are exposed because of the structural and cultural constraints of a gendered workplace (Ridgeway, 2001).

Using the 1957–2011 data from the Wisconsin Longitudinal Study (WLS), we explore the effect of occupation in 1975 (at age 36) on breast cancer incidence up to age 72. Our study extends previous research in several important ways. A long follow-up captures a time lag between exposures in higher-status occupation and breast cancer onset. Because the WLS includes women’s lifetime occupational histories, we explore not only the effect of occupation at a given time point but also the effect of the duration of occupational exposures. In addition, the WLS collected extensive information on job characteristics in 1975, which enables us to uncover specific aspects of higher-status occupations that are related to breast cancer.

**Life-course mechanisms linking occupation and breast cancer**

Biopsychosocial factors affecting the development of chronic conditions operate across the life course (Ben-Shlomo & Kuh, 2002). Consequently, the etiology of chronic diseases cannot be fully understood without incorporating earlier circumstances, in particular, exposures to health-related stressors (Pearlin, Schieman, Fazio, & Meersman, 2005). We adopt a life-course approach and explore how occupational experiences in young adulthood are related to breast cancer incidence over a 36-year period. To understand this relationship, we focus on two mechanisms—estrogen-related processes and social stress—that are not mutually exclusive and may supplement each other in explaining the effect of occupation.

A traditional approach to the etiology of breast cancer focuses on ovarian hormones, especially, estrogen (Kelsey, 1993). Influences that increase cumulative lifetime exposure to estrogen are considered important risk factors for breast cancer. Among these factors are reproductive history (later age at first birth and lower parity), health behaviors (regular alcohol use, sedentary lifestyle, and obesity for post-menopausal cancers), and components of the life-course estrogen cycle, including early age at menarche, late age at menopause, and hormone replacement therapy (Boyle & Poffetta, 2009; Friedenreich & Cust, 2008; Kelsey, 1993; Reeves et al., 2007; Vogel, 2008). Empirical studies highlight the importance of adopting a life-course approach to estrogen-related factors. For example, obesity increases the risk of post-menopausal breast cancer while decreasing the risk of pre-menopausal breast cancer (Reeves et al., 2007). Moreover, adiposity in early life has a long-term effect on breast cancer risk and is inversely related to the disease risk decades later (Sangaramoorthy, Shipps, Horn-Ross, Ko, & John, 2011).

Recently researchers have become interested in the social stress pathway to breast cancer and explored the prolonged exposure to steroid hormones produced by the adrenal cortex—glucocorticoids (GCs)—as an underlying physiological mechanism (Antonova, Aronson, & Mueller, 2011; McClintok et al., 2005). The effect of chronically elevated GCs, such as cortisol, is mediated by the activation of the glucocorticoid receptor (GR). GR is ubiquitously expressed in human breast tissue both in normal epithelium and cancerous cells (Antonova et al., 2011; McClintok et al., 2005). GR activation can directly promote mammary cell proliferation and inhibit apoptosis, which increases the risk of malignant transformations (Hermes et al., 2009).

Our study focuses on a cohort of women for whom particular types of employment presented exposure to a range of daily stressors. Participants in the WLS were born in 1939, launched their work and family trajectories in the 1950s and 1960s, and were the first cohort of White educated women to join the labor force in fairly large numbers (U.S. Census, 1970). U.S. women in professional and managerial occupations in the 1960s and 1970s faced socially structured stressors associated with gender stratification and cultural scripts of gender-appropriate behaviors (Kanter, 1977; Roussell, 1974). We consider the stress of female authority in managerial occupations and the stress of caring in professional occupations as gendered stress processes that can increase breast cancer risk via prolonged exposure of breast tissue to the anti-apoptotic and proliferative effects of chronically elevated cortisol.

**The stress of female authority**

Women of the WLS cohort who entered managerial occupations in the 1970s experienced prejudice and discrimination due to prevailing cultural attitudes that men made better leaders than women (Bartol, 1974; Kanter, 1977; Roussell, 1974). Neither men nor women preferred to work for a woman because women were seen as “temperamentally unfit” for management, which was consistent with the cultural stereotype of the woman boss as petty, controlling, and interfering (Bartol, 1974; Kanter, 1977). Roussell (1974) showed that high school departments headed by men were perceived as high in morale, whereas departments headed by women were perceived as high in “hindrance”—an indicator that the leader was seen as getting in the way of subordinates’ interests. Women in authority positions across a range of workplace settings found themselves socially isolated from subordinates and superiors and were more likely than men to report lack of communication and support from superiors and co-workers (Kanter, 1977; Roussell, 1974). Taken together, these findings suggest that authority positions exposed women to interpersonal tension and negative social interactions in the workplace (Korabik, 1995; Roussell, 1974).

**The stress of caring**

Traditional gender expectations in the 1950s and 1960s constrained career choices of highly educated women to primarily gender-appropriate areas, mostly teaching and nursing. In the U.S. in 1970, 25% of all professional women were nurses and 39% were teachers (U.S. Census, 1970). The proportions in our study are very similar, with 31% of professional women employed as nurses and 39% as teachers in 1975. Employees in caring occupations are required to act in their clients’ best interests and work in close contact with care recipients (Barron & West, 2007). Many workers feel responsible for clients’ well-being, which may lead to emotional and physical exhaustion, distress, and the inability to withdraw from work obligations (Barron & West, 2007). Moreover, workers in caring occupations perform emotional labor, which involves an expression of empathy and comfort as well as the suppression of negative feelings (Barron & West, 2007; Hochschild, 1983). Not only is emotional labor one of the major causes of occupational stress (Pugliesi, 1999), but also women are more psychologically and physically vulnerable than men to the adverse consequences of emotional labor and suppression (Barron & West, 2007).

In sum, the central argument of our study is that, in addition to estrogen-related factors, higher incidence of breast cancer among higher-status women may be explained by gendered occupational experiences. To evaluate the salience of the social stress pathway, we explore (a) whether women in professional and managerial occupations at age 36 had a higher breast cancer risk over the next 36 years relative to women in lower-status occupations and housewives, (b) whether the risk associated with higher-status
occupations accumulated with longer duration of psychosocial workplace exposures, and (c) the relative importance of the estrogen pathway and the social stress pathway as explanations for the effect of higher-status occupations on breast cancer risk.

Methods

The Wisconsin Longitudinal Study (WLS) is a long-term study of a random sample of men and women who graduated from Wisconsin high schools in 1957. Participants were interviewed in 1957 (5326 women), 1975 (4808 women), 1993 (4513 women), and 2004 (3792 women). The WLS sample retention is very high: 71% of women from the baseline sample participated in the 2004 wave, with vital status known for 94% of the original sample. Deceased participants were matched to the National Death Index (NDI) to ascertain the cause of death and age at death. Because the type of cancer was reported only in 2004, our analytic sample comprises 3682 women including (1) women who participated in 1975, 1993, and 2004 (both alive and deceased as of 2011), and (2) women who died of breast cancer any time after 1975 as established via the NDI. We conducted a detailed analysis of sample attrition and created two selection instruments based on the propensity score approach to adjust for potential selection bias, as described in the Methodological Appendix, Part A (available online as electronic Supplementary material).

Measures

The binary indicator of breast cancer incidence is coded 1 for all women diagnosed with breast cancer (alive and deceased) and 0 for women without breast cancer. We used two sources of information about breast cancer. First, women reported whether they had ever been diagnosed with breast cancer by a medical professional. Second, incident breast cancer cases include women who died of breast cancer before reporting it in the study. Out of the 297 incident breast cancer cases, 222 women were alive and 75 women were deceased as of 2011.

We considered two types of bias with respect to our measure of breast cancer. Type 1 bias may arise if some women who dropped out of the study were diagnosed with breast cancer but died of another cause. In this scenario, these breast cancer cases were neither reported in the study nor reflected in women’s death certificates. This proportion is likely to be very small and should be largely accounted for by our analysis of sample selection bias described in the Methodological Appendix, Part A. Type 2 bias may arise if a woman’s occupation affected her self-report of breast cancer. Results from Monte Carlo simulations described in the Methodological Appendix, Part B, suggest that our results are unlikely to be affected by this bias.

Occupation in 1975. Women reported their occupation and employer for every job spell held over the life course; these open-ended reports were assigned a three-digit Census occupational code. About 995 unique occupations were combined into major occupation categories following the U.S. Census classification system. Some of these major categories were further combined into broader groups to produce the final set of categories used in our analysis based on three-digit 1970 Census codes: housewife (985); professional (001–195); managerial (201–245); clerical, sales, service (260–395, 901–954, 980–984); crafts, operatives, laborers (401–824). Women reported the start and end dates of each job held between high school and retirement. A measure of job duration reflects a number of years that a woman spent in the job reported during the 1975 interview.

Job characteristics in 1975. The number of hours worked per week is represented with four dummy variables: fewer than 20, 20–29, 30–39, and 40 or more. The frequency of working under time pressure is coded from 1 = never to 5 = always. Women’s responsibility for things at work outside of her control is coded as 1 = never, 2 = rarely, 3 = some, 4 = frequently. High job autonomy is coded 1 if a woman reported not being supervised at work. An indicator of job satisfaction reflects the extent to which a woman was satisfied with her 1975 job coded from 1 = very dissatisfied to 4 = very satisfied. Job authority is measured with four items: “Do you have authority to hire and fire others?” “Can you influence pay received by others?” “Do you supervise the work of others?” “Do you decide what others do and how they do it?” This variable is dichotomized into high job authority (2+ tasks) and low job authority (0–1 tasks). All measures of job characteristics have been widely used in social sciences, and their validity and reliability are well-documented (Jencks, Perman, & Rainwater, 1988; Wolf & Fligstein, 1979).

Estrogen-related variables. Reproductive history in 1975 and 1993 is represented with a variable coded 1 if a woman had at least one child, age at first birth, and the number of biological children. Facial adropisity in adolescence was coded from pictures in 1957 high school yearbooks by six coders on a scale ranging from 1 (the lowest) to 11 (the highest). The scale has very high reliability and validity (Reithier, Hauser, & Swallen, 2009). Body mass index (BMI) in 1993 is categorized as healthy weight (BMI < 25), overweight (BMI 25–29.9), and obese (BMI ≥ 30). Alcohol use in 1993 reflects the number of days on which a woman consumed alcohol in a month prior to the interview and the number of drinks for each day on which alcohol was consumed. Participation in light exercise (such as walking) and vigorous exercise (such as jogging) in 1993 are coded from 1 = less than once a month to 4 = 3 or more times a week. The life-course estrogen cycle is reflected with ages at menarche and menopause, hysterectomy/oophorectomy (1 = yes), and hormone replacement therapy ever (1 = yes).

Control variables include education (in years), logged household annual income, and marital status (1 = married). Family history of breast cancer is coded 1 for women who reported that their mother or sister was diagnosed with breast cancer.

Our variables have 23% of missing values on average. Multiple imputation analysis was carried out in Stata 12.1 using two imputation models. The first model (for the analysis presented in Table 2) included all variables from Table 2. The second model (for the analysis presented in Table 3) included all variables from Table 3. Both imputation models also included a binary indicator of a breast cancer diagnosis, the cumulative baseline hazard, and characteristics of family background in 1957 described in the Methodological Appendix, Part A. Five completed data sets were generated under each imputation model, the survival analysis was conducted separately on each data set, and five sets of results were pooled into a single multiple-imputation inference.

Analytic plan

We begin by comparing means/proportions for all study variables by 1975 occupation categories (Table 1). To estimate the effect of occupation on breast cancer, we use a semi-parametric Cox survival model. The hazard function for woman i at time t is modeled as:

$$h(t) = h_0(t) \exp(X_i \beta + Z_i \gamma)$$  (1)

where $h_0(t)$ is the hazard of breast cancer incidence evaluated at exact age t, $h_0$ is a nonparametric baseline hazard, $\beta$ and $\gamma$ are vectors of parameters containing the effects of variables on the breast cancer hazard, $X_i$ is a vector of occupation categories, and $Z_i$ is a vector of mediating and control variables. The test of the proportionality...
assumption indicates that the effect of each occupation category is constant over time, thus, satisfying the assumption. To illustrate a significant interaction term, we plot predicted hazard functions (Fig. 1).

Table 2 shows results from models predicting breast cancer incidence for women diagnosed after 1975. Table 3 presents findings for women diagnosed after 1993 because health behaviors and certain other estrogen-related variables were reported for the first time only in 1993. To explore the potential effects of duration of occupational exposures, we centered the number of years that each woman spent in her 1975 job at the median (8 years), limited our sample to women in professional and managerial occupations, and estimated an interactive effect of job authority and job duration. To illustrate a significant interaction term, we plot predicted hazard functions (Fig. 1).

Finally, we apply a decomposition technique for a formal test of mediation in a survival setting (VanderWeele, 2011) to examine the extent to which estrogen-related and social stress variables mediate the effect of occupation. If we denote the focal predictor as \( X \), the mediator as \( M \), and a vector of control variables as \( Z \), then

\[
\log h(t|X, M, Z) = \log(h_0) + \gamma_1 X + \gamma_2 M + \gamma_3 Z
\]

(2)

Further, if we denote the effect of \( X \) on \( M \) as \( \delta_X \), the total effect of the focal predictor reflecting a change in the log hazard of breast...
cancer with a one-unit change in the predictor from $X$ to $X^*$ can be decomposed into direct and indirect effects as follows:

$$
\log h(t, X, M) - \log h(t, X^*, M^*) = \left[ \gamma_1^* (X - X^*) \right] + \left[ \gamma_2^* \beta_1 (X - X^*) \right]
$$

where $\gamma_1^*$ is the direct effect and $\gamma_2^* \beta_1$ is the indirect effect.

### Results

#### Descriptive analysis

Table 1 shows that compared to housewives, a significantly greater proportion of women in professional and managerial occupations were diagnosed with breast cancer (0.095 and 0.145, respectively, $p < 0.001$). In contrast, women in blue-collar occupations had a significantly lower prevalence of breast cancer (0.051, $p < 0.05$), whereas women in clerical, sales, and service occupations were similar to housewives in terms of breast cancer risk.

With respect to job characteristics in 1975, women in managerial occupations had the highest job authority but also worked longer hours and were more frequently held responsible for things outside their control. Professional and managerial women worked under time pressure more frequently but also reported higher job satisfaction than lower-status women. Women in professional occupations were less likely than managerial women but more likely than lower-status women to supervise others and decide what/how others did at work. Women in managerial occupations were much more likely than all other women to have high job authority (72%), although job authority was still more prevalent among professional women (27%) than women in lower-status occupations. Finally, lower-status women spent about two fewer years in their 1975 job than professional and managerial women.

With respect to estrogen-related variables, women in higher-status occupations had several characteristics of an unfavorable profile of breast cancer risk: later age at first birth, lower parity, more regular use of alcohol, lower adiposity in adolescence, and, among professional women only, later age at menopause and higher use of hormone replacement therapy. Yet, two characteristics reflected a favorable risk profile: higher physical activity (professional women only) and a lower risk of obesity in midlife.

### Survival analysis

Model 1 in Table 2 reveals that, compared to housewives, women in professional occupations had 72% higher risk (HR = 1.72, 95% CI: 1.25, 2.36) and women in managerial occupation had 57% higher risk (HR = 1.57, 95% CI: 1.02, 2.42) of a breast cancer diagnosis after 1975. In contrast, women in lower-status occupations were similar to housewives in terms of breast cancer risk.

Models 2 and 3 explore the estrogen-related pathway. As indicated in Model 2, higher adiposity in adolescence was associated with a lower breast cancer risk (HR = 0.82, 95% CI: 0.71, 0.95); yet, the effect of higher-status occupation changed only trivially compared to Model 1. Model 3 includes reproductive history and shows that nulliparity and later age at first birth increased the risk of breast cancer. The mediation decomposition analysis reveals that reproductive variables mediate 23% of the association between professional occupations and breast cancer hazard, and this
The mediating effect is significant at the 0.01 level. Yet, the elevated breast cancer risk of professional women remained statistically significant and large in magnitude (HR = 1.59, 95% CI: 1.15, 2.20). In contrast, adjustment for reproductive variables does not change the effect of managerial occupations.

Table 3

Cox proportional hazard models predicting breast cancer incidence based on occupation in 1975 among women diagnosed after 1993 (N = 3646).

<table>
<thead>
<tr>
<th>Variables</th>
<th>Model 1</th>
<th>Model 2</th>
<th>Model 3</th>
<th>Model 4</th>
</tr>
</thead>
<tbody>
<tr>
<td>Occupation in 1975:</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Housewifea</td>
<td>1.00</td>
<td>1.00</td>
<td>1.00</td>
<td>1.00</td>
</tr>
<tr>
<td>Professional</td>
<td>2.22*** (1.55, 3.19)</td>
<td>2.07*** (1.43, 2.99)</td>
<td>1.89*** (1.30, 2.75)</td>
<td>1.91*** (1.31, 2.77)</td>
</tr>
<tr>
<td>Managerial</td>
<td>1.89** (1.18, 3.05)</td>
<td>1.50 (0.89, 2.53)</td>
<td>1.49 (0.88, 2.52)</td>
<td>1.50 (0.88, 2.55)</td>
</tr>
<tr>
<td>Clerical, sales, service</td>
<td>0.90 (0.63, 1.29)</td>
<td>0.870 (0.60, 1.25)</td>
<td>0.895 (0.62, 1.29)</td>
<td>0.883 (0.61, 1.27)</td>
</tr>
<tr>
<td>Crafts, operatives, laborers</td>
<td>0.82 (0.49, 1.39)</td>
<td>0.78 (0.46, 1.33)</td>
<td>0.84 (0.50, 1.43)</td>
<td>0.87 (0.51, 1.48)</td>
</tr>
<tr>
<td>High job authority in 1975</td>
<td>1.45* (1.02, 2.07)</td>
<td>1.49* (1.05, 2.13)</td>
<td>1.49* (1.05, 2.13)</td>
<td></td>
</tr>
<tr>
<td>Estrogen-related variables:</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Adiposity in 1957</td>
<td></td>
<td></td>
<td></td>
<td>0.81* (0.67, 0.97)</td>
</tr>
<tr>
<td>Reproductive history in 1993:</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>At least one birth</td>
<td>0.32 (0.09, 1.05)</td>
<td>0.30* (0.09, 0.99)</td>
<td>0.30* (0.09, 0.99)</td>
<td></td>
</tr>
<tr>
<td>Birth × age at first birth</td>
<td>1.05* (1.01, 1.09)</td>
<td>1.05* (1.01, 1.09)</td>
<td>1.05* (1.01, 1.09)</td>
<td></td>
</tr>
<tr>
<td>Number of children</td>
<td>0.95 (0.69, 1.30)</td>
<td>0.94 (0.69, 1.29)</td>
<td>0.94 (0.69, 1.29)</td>
<td></td>
</tr>
<tr>
<td>Health behaviors in 1993:</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Healthy weight (BMI &lt; 25)a</td>
<td>1.00</td>
<td></td>
<td></td>
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<tr>
<td>Overweight (BMI 25–29.9)</td>
<td>1.06 (0.70, 1.61)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Obese (BMI ≥ 30)</td>
<td>1.19 (0.87, 1.63)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Daily number of drinks</td>
<td>1.00 (0.98, 1.02)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Days of the month when drinks</td>
<td>1.07 (0.93, 1.23)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Light physical activity</td>
<td>1.07 (0.94, 1.22)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Vigorous physical activity</td>
<td>0.97 (0.91, 1.05)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Life-course estrogen cycle:</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Age at menarche</td>
<td>0.93 (0.85, 1.03)</td>
<td>0.93 (0.85, 1.03)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Age at menopause</td>
<td>1.03* (1.01, 1.05)</td>
<td>1.03* (1.01, 1.05)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Hysterectomy/oophorectomy</td>
<td>0.81 (0.57, 1.15)</td>
<td>0.82 (0.57, 1.16)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Hormone replacement therapy</td>
<td>1.36 (1.03, 1.79)</td>
<td>1.30 (1.00, 1.73)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Family history of breast cancer</td>
<td>1.12 (0.57, 1.15)</td>
<td>1.12 (0.57, 1.15)</td>
<td>1.13 (0.57, 1.16)</td>
<td></td>
</tr>
<tr>
<td>Attrition propensity score</td>
<td>0.84 (0.64, 1.09)</td>
<td>0.84 (0.64, 1.09)</td>
<td>0.83 (0.63, 1.09)</td>
<td>0.87 (0.66, 1.15)</td>
</tr>
<tr>
<td>Model fit:</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Log likelihood (df)b</td>
<td>−1772 (7)</td>
<td>−1770 (8)</td>
<td>−1760 (16)</td>
<td>−1755 (23)</td>
</tr>
<tr>
<td>AICc</td>
<td>3559</td>
<td>3557</td>
<td>3552</td>
<td>3557</td>
</tr>
<tr>
<td>BICd</td>
<td>3608</td>
<td>3604</td>
<td>3653</td>
<td>3702</td>
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</table>

Note: Each cell contains hazard ratios and 95% confidence intervals (in parentheses). All models control for education, household income, and marital status. *p < 0.05. **p < 0.01. ***p < 0.001 (two-tailed test).

a Reference group.
b df = degrees of freedom.
c AIC = Akaike information criterion.
d BIC = Bayesian information criterion.

Models 4 and 5 evaluate the social stress pathway. Weekly work hours, time pressure, responsibility for things at work outside own control, job autonomy, and job satisfaction are not significantly related to breast cancer incidence (Model 4). In a sensitivity analysis we estimated the effect of each job characteristic separately and found that none of them was significant even without other variables in the model. Interestingly, the effect of higher-status occupations even increases compared to Model 3 because responsibility for things outside one’s own control, time pressure, and job satisfaction are more prevalent among higher-status women and are related negatively (although not significantly) to breast cancer risk.

When job authority is added in Model 5, breast cancer risk associated with managerial occupations declines and becomes not significant (HR = 1.42, 95% CI: 0.87, 2.30). Women in managerial occupations had higher job authority than other women. In turn, high job authority is associated with a 1.57 greater hazard of breast cancer than low authority (95% CI: 1.12, 2.18). The decomposition analysis indicates that job authority mediates 55% of the relationship between managerial occupations and breast cancer incidence. This mediating effect of job authority is significant at the 0.001 level. In contrast to women in managerial occupations, job authority does not explain the elevated risk of breast cancer among professional women likely because their jobs involved less authority. Only 27% of professional women reported high levels of job authority.
authority compared to 72% of women in managerial occupations (Table 1). Job authority mediates 12% of the effect of professional occupations on breast cancer hazard, and this indirect effect is not statistically significant.

We explored the accumulation of breast cancer risk with duration of occupational exposures and found a significant interactive effect between job authority and duration in the 1975 job. Fig. 1 shows that the effect of job authority on the breast cancer hazard accumulates with longer duration among professional and managerial women who had high authority (HRjob authority × duration = 1.15, 95% CI: 1.06, 1.27). For example, women with high job authority who worked for 15 years in this job had a significantly greater risk of breast cancer than women with similar duration but low job authority. Women with low job authority and short job duration (5 years) had the lowest hazard.

Table 3 presents results from models based on women diagnosed with breast cancer after 1993, thus, excluding 36 women. Model 1 indicates that the effect of professional (HR = 2.22, 95% CI: 1.55, 3.19) and managerial (HR = 1.89, 95% CI: 1.18, 3.05) occupations becomes even stronger when women with a diagnosis before age 54 are excluded. Model 2 confirms that, consistent with Model 5 of Table 2, the effect of managerial occupations is reduced substantially and becomes not significant after adjustment for job authority, whereas the effect of professional occupations does not change.

Model 3 includes reproductive history and characteristics of life-course estrogen cycle. Later age at first birth, later age at menopause, and hormone replacement therapy are related to a higher risk of breast cancer. The decomposition analysis indicates that 20% of the effect of professional occupations is conveyed indirectly via these estrogen-related variables (p < 0.01). The effect of managerial occupations, already not significant in Model 2, was not altered after adjustment for estrogen-related variables. Job authority and estrogen-related variables are both significant predictors of breast cancer net of each other, which emphasizes the additive effects of these two mechanisms. Finally, Model 4 indicates that higher adiposity in 1957 reduces breast cancer risk, whereas health behaviors in 1993 are unrelated to post-1993 breast cancer. The effects of higher-status occupations changed only trivially after adjustment for all estrogen-related health behaviors in Model 4.

Discussion

Drawing on a life-course biosocial stress framework, our study documents long-term effects of higher-status occupations on women’s elevated risk of a breast cancer diagnosis. We find that women who were in professional and managerial occupations in 1975 at age 36 had a substantially higher risk of a breast cancer diagnosis up to age 72 compared to housewives and women in lower-status occupations. To explain the long-reaching effect of occupation in 1975, we assess estrogen-related and social stress explanations and find that these mechanisms have additive effects, which points to their complementary nature in explaining the elevated breast cancer risk of professional and managerial women. With respect to the estrogen-related pathway, about 20% of the elevated breast cancer risk among professional women was explained by their later age at first birth, lower parity, more regular alcohol use, higher use of hormone replacement therapy, and later menopause. Yet, the effect of professional occupations remains large and significant net of estrogen-related variables. Moreover, estrogen variables have little effect on the association between managerial occupations and breast cancer risk. These patterns are consistent with other studies that show that the effect of higher-status occupation decreases only modestly and remains large in magnitude and statistically significant after adjustment for reproductive histories and other estrogen-related variables (Dana et al., 2004; Larsen et al., 2011).

With respect to the social stress pathway, our findings point to women’s job authority as a potentially important source of job-related stress (Schieman & Reid, 2009). Job authority mediated 55% of the effect of managerial occupations on breast cancer risk, which reduced the direct effect of managerial occupations to non-significance. Women in managerial occupations had higher job authority than other women. In turn, high job authority is associated with a 1.55 higher risk of breast cancer than low authority, net of all estrogen-related and control variables. Further, our findings reveal that the risk of breast cancer associated with job authority accumulates with longer duration in the professional/managerial occupations. This pattern points to the importance of chronic stress resulting in gradual accumulation of deleterious exposures and incremental transformations of bodily systems (Ben-Shlomo & Kuh, 2002; Pearlin et al., 2005). Consistent with the life course perspective, we emphasize the concept of duration (Pearlin et al., 2005) and show how social risk factors for breast cancer operate over extended periods of time resulting in the compounding of cumulative damage with the longer exposure to stressors.

Exercising job authority was particularly stressful for women in the context of gender inequality embedded in the occupational structure of the 1960s and 1970s, when women in managerial positions often faced prejudice, tokenism, discrimination, social isolation, and resistance from subordinates, colleagues, and superiors (Rousell, 1974; Kanter, 1977). This chronic stress may have been an important long-term link between higher-status jobs in young adulthood and the risk of breast cancer. Animal models suggest potential mechanisms through which interpersonal stress of higher status is related to chronic hyperactivity of glucocorticoid responses. Among female rats, social isolation and disruption of supportive ties increase the risk of developing mammary tumors fourfold, with the primary mechanism being dysregulation of the GC system (McClintok et al., 2005). In young adulthood, months before tumor initiation, socially isolated rats developed an enhanced GC response to stressors with markedly delayed reduction of corticosterone to normal levels. Both aspects of this stress reactivity process were related to an increased risk of mammary tumors in middle and old age (Hermes et al., 2009).

Further, studies among baboons suggest that in some contexts, dominant individuals exhibit an unfavorable profile of GC hyperactivity. Cortisol dysregulation among higher-status primates is associated with low social control, lack of social support, and situations where dominant individuals have to repeatedly reassert their rank (Gesquiere et al., 2011). This scenario is consistent with interpersonal tension and resistance experienced by incumbents of authority positions who are not perceived as legitimate (Ridgeway, 2001). Thus, the GC-related effects of higher status may have also extended to this cohort of women in managerial occupations. These mechanisms remain speculative in our study because we do not have direct measures of women’s perceptions of gender discrimination or biomarkers of the GC system. We view our findings as the first step suggesting that it is worthwhile to explore women’s experiences in higher-status occupations with respect to breast cancer and a call for more attention to stressful occupational exposures as explanations for the puzzling effect of managerial occupations.

Our findings provide indirect support for the stress of caring model among professional women. The effect of professional occupations on the elevated risk of breast cancer was not explained by all estrogen-related variables and job characteristics available in our study. Because professional women were predominantly employed in caring occupations, mostly teaching and nursing, characteristics reflecting the stressful side of carework may be an important underlying mechanism. Emotional suppression and
emotional exhaustion among women teachers is associated with higher
stress to breast cancer in one study because these stress processes
operate on different levels: social, psychological, physiological, and
molecular. Therefore, important directions for future research
include developing interdisciplinary collaborations, integrating
multiple levels of analysis, and collecting longitudinal data on
chronic strains and biomarkers among women in higher-status
occupations.

Appendix A. Supplementary data

Supplementary data related to this article can be found at http://
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Limitations and future directions

Our study has several important limitations. The analyses are
based on one cohort of women who were born in 1939; thus, our
findings may be most relevant to women born in the first half of
the 20th century. Yet, because of the persistence of gender inequality
and dominant gender beliefs, our study is still likely to reflect
experiences of women in current and future cohorts (Ridgeway,
2001). Further, the WLS contains only White non-Hispanic partic-
ipants. Conditions of minority women in higher-status positions
may be even more stressful than those of White women. Recent
biological research suggests that the cortisol-mediated effects of
social stress on breast cancer may be stronger among Black women
compared to White women (McClintock et al., 2005). An important
step for future research is to explore race differences in health
implications of higher-status occupations.

Some estrogen-related variables were not measured in our
study, in particular, oral contraceptives and breastfeeding. Evidence
for the role of these two variables in breast cancer etiology is mixed
and varies by context and duration (Kelsey, 1993; Nichols et al.,
2007). Moreover, breastfeeding rates were very low in the 1970s
among women of all social classes (Wolf, 2003). Although it is
unlikely that inclusion of these variables would substantially alter
our findings, it is still important to incorporate a wide range of
indicators of life-course estrogen cycle to assess more precisely
their relative importance compared to the social stress pathway.
Further, the WLS did not collect information on health behaviors in
1975. Therefore, our conclusions about the lack of explanatory
power of health behaviors in midlife (in 1993) are limited because
lifestyle assessed contemporaneously with occupation in 1975 may
have been more consequential for breast cancer than behaviors in
midlife.

In this study we focus on chronic stress and do not include acute
stressful life events, such as loss of loved ones or disease in the
family, which can have potential—albeit modest—effects on breast
cancer (Kruk & Aboul-Enein, 2004). For a more nuanced under-
standing of the social stress pathway, future research should
distinguish between chronic strains and stressful events (Pearlin
et al., 2005). Moreover, a myriad of other aspects of higher-status
jobs are not measured in the WLS. We could not include mea-
sures of emotional labor, emotional suppression, gender discrimi-
nation, women’s actual relationships with co-workers and superiors,
and women’s perceptions of social support and social strain at work in
1975. Yet, we are unaware of any data set that combines the strengths of
the WLS with more comprehensive measures of stressful experiences in
higher-status occupations. Moreover, it is impossible to document all pathways linking
social stress to breast cancer in one study because these stress processes
operate on different levels: social, psychological, physiological, and
molecular. Therefore, important directions for future research