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Title: Does stress increase risk of breast cancer? A 15-year prospective study.

Running title: Stress and risk of breast cancer: A 15-year prospective study

Authors

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Abstract

Objective: The possible impact of stress on cancer incidence remains controversial. We prospectively evaluated associations between life event stressors, social support, personality characteristics (optimism, anger control, anti-emotionality) and risk of developing primary breast cancer (BCa), in women at increased familial risk of BCa.

Methods: A prospective cohort, repeated measures design was employed. Recruitment was through the Kathleen Cuningham Foundation Consortium for Research into Familial breast cancer (kConFab), which collects genetic, epidemiological and clinical data from Australasian families with multiple BCa cases. Acute and chronic stressors for the prior three years, and psychosocial, clinical and epidemiological variables were measured at cohort entry and at three-yearly intervals. Cox proportional hazards regression analysis controlling for BCa risk factors and familial clustering, was undertaken. The primary outcome was histopathologically confirmed BCa (invasive or ductal carcinoma in situ (DCIS), including occult cases diagnosed during risk-reducing mastectomy).

Results: Of 3,595 consecutive women invited to participate, 3,054 (85.0%) consented. Of these, 2,739 (89.7%) from 990 families (range 1-16 per family) completed at least one assessment.
point. During the study 103 women were diagnosed with BCa. No stressor or psychosocial variable or interaction between them, was significantly associated with BCa in unadjusted or adjusted models (total acute stressors HR=1·03 (0·99-1·08), p=0·19; total chronic stressors HR=1·0 [0·90-1·11], p=0·98).

**Conclusions:** This study did not demonstrate an association between acute and chronic stressors, social support, optimism, anti-emotionality or anger control, and BCa risk. Women should focus on proven methods of BCa risk reduction.

**Key words:** breast cancer, high-risk, life events, prospective study, stress,
INTRODUCTION

Breast cancer (BCa) is the most common cancer and second most common cause of cancer death in women.\(^1\) Risk reduction, therefore, is a critical goal of health care, particularly for women at increased familial risk. Currently, risk reduction options for high-risk women, such as preventative surgery and risk-reducing medication, have potential morbidity, and uptake is variable.\(^2\) Less intensive strategies, such as regular exercise and avoiding prolonged use of hormone replacement therapy, offer some risk-reduction, but additional strategies are urgently required.\(^2\)

A persistent belief in the general community is that life event stressors have an adverse effect on cancer risk, particularly BCa.\(^3\) The literature on this topic is fraught with methodological limitations, including inadequate power and control of potential confounders, measurement error and an atheoretical approach that fails to consider potential moderators of stressor impact, such as stressor severity and chronicity, social support, personality and coping style.\(^4\) Moreover, the evidence is equivocal. The Chida et al (2008) meta-analysis\(^5\) concluded that life events do not, but a stress-prone personality, unfavorable coping styles and negative emotional responses do adversely effect cancer incidence, while noting evidence of publication bias. The Lin et al (2013) meta-analysis\(^6\) also concluded that life events as a whole are not associated with BCa risk, but high-intensity life events do increase risk of BCa. The Nielsen et al (2006) review\(^4\) suggested a fair test of the hypothesis has yet to be undertaken.
An important methodological challenge in this research is that exposure to high-intensity stressors, such as bereavement or divorce, is sporadic and cannot be imputed over time. Also, sufficient new BCa diagnoses within the study time frame are required. Retrospective registry-based studies linking recorded stressful life events (such as divorces and deaths) with cancer incidence, offer one solution to the large samples required, and have the advantage that exposure information is independent of subjects’ recall which may be biased. However, such studies are limited by the restricted range of stressful life events recorded. Case control studies (usually conducted on women with breast symptoms awaiting definitive diagnosis) are able to collect more detailed exposure data, but in a stressful pre-diagnostic environment, factors such as symptoms, family history, age and diagnostic communication might cue women regarding their breast cancer risk and bias stressful life event reporting. A longitudinal repeated measures design with initially unaffected participants is ideal since more detailed exposure data can be gathered, and recall will not be biased by existing symptoms or environmental factors. A large sample of women with a moderate to strong familial risk of BCa and/or \textit{BRCA1} and \textit{BRCA2} mutations, followed over time, can meet these requirements.

The aim of this study was to prospectively evaluate associations between acute and chronic life event stressors, social support, personality characteristics (optimism, anger control, anti-emotionality) and the risk of developing primary BCa in women at increased familial risk of
BCa, independent of known risk factors and familial clustering.

The primary hypothesis was that exposure to high levels of acute and chronic stressors would be associated with an increased risk of developing BCa. Secondary hypotheses included that: there would be a dose-response relationship (or threshold effect), such that more severe and prolonged stressors would be associated with a greater increased risk of BCa; and that psychosocial factors (social support, optimism, anti-emotionality, anger repression) would be associated with, or moderate the impact of stressors on, risk of BCa.

METHODS

Study design

A prospective cohort, repeated measures design was employed, and complied with the Strobe statement for cohort studies. Acute and chronic stressors for the prior three years, and psychosocial, clinical and epidemiological variables were measured at entry into the psychosocial study cohort and at three-yearly intervals.

Recruitment Source

Recruitment was undertaken through the Kathleen Cuningham Foundation Consortium for Research into Familial breast cancer (kConFab) established in 1995 to co-ordinate the
collection of genetic, epidemiological and clinical data in Australasian families at high-risk of BCa.

kConFab eligibility criteria for families include having an identified deleterious mutation in 
*BRCA1*, *BRCA2* or another breast cancer predisposition gene, or ≥ one ‘high-risk’ family member, in addition to ≥ four cases of breast/ovarian cancer on one side of the family, ≥ two living family members with breast/ovarian cancer and ≥ four living first or second-degree unaffected female relatives aged 18 years and over.

Detailed family history and epidemiological data are collected at entry to kConFab and blood samples taken to enable mutation testing. The kConFab Clinical Follow-Up study updates new cancer events epidemiological and lifestyle risk data, screening behaviors and risk-reducing surgery and medication use at three-yearly intervals using a mailed questionnaire to participants, with subsequent confirmation of reported surgery and new cancer events from medical records.

**Eligibility Criteria for Psychosocial Study**

kConFab women aged 18-75 years, with no personal history of cancer (except non-melanoma skin cancer or in-situ carcinoma of the cervix), were invited to participate in the psychosocial study. Non-English-literacy, and physical or mental illness affecting memory recall, were exclusion criteria.
**Procedure**

The psychosocial study was introduced to women when they entered kConFab or the clinical follow-up study. Interested participants were mailed an information statement, consent form and questionnaire with a reply-paid envelope by the psychosocial study team; non-responders were contacted by mail or phone as needed. Semi-structured phone interviews elicited acute (AS) and chronic (CS) stressful life events over the three years previous to the interview date.\textsuperscript{13,14} Interviews and questionnaires were repeated three yearly, contiguous with the clinical follow-up study. This study was approved by the institutional ethics review boards of all participating sites (ethics number H02/050/02).

**Measures**

**Primary outcome**

The primary outcome was histopathologically confirmed BCa (invasive or ductal carcinoma in situ, including occult cases diagnosed during risk-reducing mastectomy).

**Key Independent Variables**

*Acute and Chronic Stressors:* The Life Events and Difficulties Schedule (LEDS),\textsuperscript{11} (a semi-structured interview protocol, was employed. Trained interviewers developed written clinical vignettes for each stressor elicited, including context and relevant personal and biographical
information. An independent rater rated stressors using precise, pre-defined criteria (IRR Kappa=0.71-0.78), to provide a constant, objective reference point to ensure consistency of ratings of threat/loss (i.e. the severity of the stressors) across participants. Each stressor was coded on multiple features, including:

- **Type** – Acute (AS) (discrete event) or chronic (CS) (continuing for at least 6 months);
- **Time - Date** (closest month); if CS duration (years) and current (yes/no);
- **Severity** (S) (mild, moderate, high, severe); and
- **Independence** from personal or familial risk of BCa or BCa diagnosis; BCa risk-related stressors (including family cancer diagnoses, recurrences and deaths, receipt of mutation testing results (for self or a family member) and risk-reducing surgery) were excluded from all analyses).

Examples are: death of a spouse/child (AS: severe); buying/selling a home (AS: moderate); handicapped child requiring fulltime care (CS: severe); ongoing acrimonious dispute with neighbours over noise (CS: moderate).

**Psychosocial Variables**

Psychosocial variables included those previously identified as predictors or moderators of the impact of stressors on BCa risk (social support and personality factors), using psychometrically proven measures.
Social Support: The eight-item Duke-UNC Functional Social Support Questionnaire measures satisfaction with social support.\textsuperscript{15} Total scores range from eight to 40; higher scores indicate greater social support.

Optimism: The 12-item Life Orientation Test (LOT) scale\textsuperscript{16} measures dispositional optimism. Total scores range from 0 to 32; high scores indicate greater optimism.

Anti-Emotionality: The four-item Anti-Emotionality Scale\textsuperscript{17} assesses absence of emotional behaviour. Scores range from one to four; higher scores indicate more anti-emotional behaviour.

Anger Control: The seven-item subscale of the Courtauld Emotional Control Scale\textsuperscript{18} measures anger control. Total scores range from seven to 28; higher scores indicate greater control.

Potential Covariates/Confounders

Current Anxiety and Depression: We controlled for current anxiety and depression, which may influence self-report of stressors and psychosocial variables, in all models containing psychosocial variables. The seven-item subscales of the Hospital Anxiety and Depression Scale\textsuperscript{19} measure anxiety and depression over the past week. Scores range between zero and 21; high scores indicate greater current anxiety and depression.
As known at last time-point: mutation status (BRCA1 mutation, BRCA2 mutation, no BRCA1 or BRCA2 mutation, untested/unknown); number of 1st and 2nd degree relatives with a) BCa and b) ovarian cancer; age at menarche; number of live births; parity by breast feeding (nulliparous, parous and breast fed, parous and not breast fed); smoking history (current, former, never); exercise (sufficiently active, insufficiently active, sedentary, unknown).  

Time-varying: bilateral salpingo-oophorectomy; benign breast disease; hormone replacement therapy use (never, not within last three years, within last three years, unknown); oral contraception use (never, not within three years, within last three years, unknown); body mass index; and anxiety and depression (where applicable).

STATISTICAL METHODS

Cox proportional hazards regression analysis, with the time scale anchored to the age at the start and end of the time period covered by each LEDS interview, was performed to examine the association between stressors recorded in each three-year period and diagnosis of BCa in the next three-year period. Mutation status was entered as a stratification variable to allow for differing baseline hazard functions. The analysis was adjusted for potential confounders and included a random family effect to accommodate covariance due to familial clustering. Women undergoing risk-reducing mastectomy were censored at that date. Number and severity of stressors, and each
psychosocial variable, were evaluated in separate models. Statistical tests were two-sided with a Type 1 error rate of 5% for main effects.

Simulation studies indicate that ten events per parameter estimated provides reliable estimates of the hazard ratios. Over the study period, 103 women were diagnosed with BCa; therefore, a model with four stressor variables and four psychosocial variables (our final model) can be evaluated reliably. With insufficient power to evaluate all the covariables, they are not individually interpreted. Analysis was conducted by a qualified biostatistician (JC).

RESULTS

Of 3,595 eligible women consecutively invited to participate in the psychosocial study between May 2001 and December 2010, 3,054 (85·0%) consented. Of these 2,739 (89·7%) from 990 families (range 1-16 per family) completed at least one LEDS interview and were included in the analyses. The mean follow-up was 7·2 years, with 645 (23·5%) completing one data time-point, 916 (33·4%) completing two, 980 (35·8%) completing three and 200 (7·3%) completing four (attenuated with study closure in December 2011). During the study period 103 women from 97 families were diagnosed with BCa.

On average, women were aged 45·0 years at study entry, 6% were BRCA1 and 6% were BRCA2 mutation positive, 34% were confirmed BRCA1 or BRCA2 mutation negative and 54% were
without a known \textit{BRCA1} or \textit{BRCA2} mutation within their family and therefore not tested. On average the women reported scores at the upper end of the range in social support and optimism, and in the lower end of the range in anti-emotionality and anger control (see Supplementary Table 1).

Descriptive data for acute and chronic stressors, frequency and severity, are shown in Table 1. Most stressors, both acute and chronic, were mild or moderate in severity. The distribution of stressors was similar across those with and without BCa.

Unadjusted and adjusted hazard ratios and p-values for stressor variables and psychosocial variables are shown in Table 2. The total number of AS and CS were not associated with BCa risk (Model 1). Model 2 included the total number of AS and CS by severity (mild, moderate, high, severe). The risk of BCa was similar across severity for AS. For CS, the hazard ratio was somewhat greater for high and severe CS than for mild and moderate CS, although the confidence intervals are wide. A similar pattern is observable in Model 3, where the number of AS and CS were collapsed into two categories: low-moderate and high-severe. Although none of the stressor variables reached statistical significance, the hazard ratio (HR) for the high-severe CS was 1·28 (95% Confidence Interval (CI) 0·88-1·85) in the adjusted model (HR=1·31; CI 0·92-1·87 unadjusted model). As the study was powered to explore a threshold of at least one high-severe stressor, Model 4 examined the threshold effect of exposure to high-severe acute stressors.
and chronic stressors in the absence of mild-moderate stressors, although similarly to Model 3, neither reached statistical significance.

Models 5-8 evaluated the association between each time-varying psychosocial variable (adjusted for anxiety and depression) and BCa diagnosis. None of the psychosocial variables reached statistical significance.

Model 9, the full model, included the total number of low-moderate AS, high-severe AS, low-moderate CS, and high-severe CS, and the time-varying psychosocial variables. None of the stressors or psychosocial variables were significantly associated with BCa in the unadjusted or adjusted models. Interactions between the stressor variables and the psychosocial variables were tested and none reached statistical significance after adjusting for covariables. The results of Model 9 are shown in Table 2 and Supplementary Figures 1 and 2.

**DISCUSSION**

In this rigorous, prospective study evaluating the impact of stressors and psychosocial variables on BCa risk, we found no significant associations, although there was a trend for significance for high severe chronic life stress (HR: 0.91-2.04, p=0.13). The study addressed methodological issues which have limited interpretation of prior studies, including: objective, consistent ratings
of stressors, differentiating acute and chronic stressors of different types (e.g. illness, financial); assessing stressors prior to diagnosis to avoid recall bias; and controlling for BCa risk factors, anxiety and depression at time of self-reported psychosocial data collection, and potential behavioural consequences of stressors such as smoking. Stressors related to BCa risk were excluded from all analyses.

The results may not generalise to women in the general population, not at increased familial risk. However, internationally, most rigorous studies have also failed to find an association between psychosocial factors and cancer, such as a large prospective study of life events in 11,000 women from the general population with 313 incident BCas, (Surtees et al, 2010), the UK Million Women study, a recent registry-based Danish study exploring prolonged job strain and risk of cancer in 6,571 women, a prospective study of 10,519 Finnish women measuring self-perceived stress levels in Finland, and a prospective cohort study of 30,277 Japanese residents.

In contrast, the Finnish Twin Cohort, involving 10,808 women (180 incident BCas) found that severe events, (including divorce/separation, death of a husband/close relative/friend), were associated with increased risk of BCa. Thus, the literature on the general population remains somewhat conflicted. None of these studies used the gold-standard approach to measuring
stressors used here. It is also possible that moderating variables not measured, such as coping strategies tailored to different stressors, may influence impact of stressors on BCa risk.

**Clinical Implications**

Over the past 40 years, women have been exposed to strong messages about the importance of ‘thinking positively’ and reducing stress in their lives, which can add to the burden of guilt in those who develop cancer, who feel they have somehow failed. Our results, based on rigorous methodology, add to the growing literature providing reassurance to women at increased risk of BCa, who are concerned that the (often unavoidable) stressors in their lives may increase their risk of BCa. Of course, reducing the impact of stress on general health and quality of life remains a worthwhile goal for women.

**Limitations of the Study**

The study was powered assuming a 5.7% rate of BCa and an 18% exposure rate of at least one high-severe AS or CS over an average follow-up of ten years in a cohort of 3100 women. This roughly equates to having 80% power to detect a HR as small as 1.73. We actually observed a BCa rate of 3.8% in our cohort of 2,739 women with an average follow-up of 7.2 years. Of the 6217 interviews included in analysis, 2675 (43.0%) had at least one high-severe AS or CS (39.0% with at least one AS plus 13.6% with at least one CS less 9.6% with at least one AS and CS). Therefore, we had 80% power to detect a HR as small as: 1.76 for at least one AS, and 2.24
for at least one CS (Model 4). Thus, the power to detect HRs of the magnitude of those observed in Model 9 was modest. The very right-skewed HR for high severe chronic events may have been significant if power was higher.

Our response rate of 85% was high, but women with more or less stress in their lives may have been differentially attracted to the study. We controlled for a very comprehensive list of potential confounders but some aspects (such as coping style) were not assessed. It is possible that our choice of women from multiple-case BCa families was flawed and that the impact of stressors may be inconsequential in comparison to the impact of genetic influences.

Conclusions
Given the lack of evidence found for an association between stressors and psychosocial factors and BCa, we suggest that women at increased risk of BCa focus on proven methods of risk reduction such as close monitoring, risk-reducing medication or, for those at high risk, surgery.

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References


how these compare to expert-endorsed risk factors. Cancer Causes & Control 2014; 25(7): 771-785.


22. Surtees PG, Wainwright NW, Luben RN, Khaw KT, Bingham SA. No evidence that social stress is associated with breast cancer incidence. Breast Cancer Research and


Table 1. Distribution of the average number of acute and chronic stressors per woman (averaged over each woman’s follow-up time)

<table>
<thead>
<tr>
<th></th>
<th>Mild-moderate stressors</th>
<th>High-severe stressors</th>
<th>All stressors</th>
</tr>
</thead>
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<tr>
<td></td>
<td>Affected (n=103)</td>
<td>Unaffected (n=2636)</td>
<td>Affected (n=103)</td>
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<tr>
<td>Acute Stressors (Mean)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>0</td>
<td>0 (0)</td>
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<td>42 (41)</td>
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<tr>
<td>&gt;0-5</td>
<td>25 (24)</td>
<td>495 (19)</td>
<td>61 (59)</td>
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<td>&gt;5-10</td>
<td>49 (48)</td>
<td>1371 (52)</td>
<td>0 (0)</td>
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<tr>
<td>&gt;10-15</td>
<td>24 (23)</td>
<td>620 (24)</td>
<td>-</td>
</tr>
<tr>
<td>&gt; 15</td>
<td>5 (5)</td>
<td>146 (6)</td>
<td>-</td>
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<tr>
<td>Chronic Stressors (Mean)</td>
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<td></td>
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<tr>
<td>0</td>
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<td>234 (9)</td>
<td>77 (75)</td>
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<td>1104 (42)</td>
<td>25 (24)</td>
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<td>&gt;2-4</td>
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<td>791 (30)</td>
<td>1 (1)</td>
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<td>&gt;4</td>
<td>19 (18)</td>
<td>507 (19)</td>
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Table 2. Unadjusted and Adjusted Hazard Ratios (HR) and 95% Confidence Intervals (CI) of a breast cancer diagnosis

<table>
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<th>Variable</th>
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<th>Adjusted</th>
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<td>95% CI</td>
<td>p-value</td>
<td>HR</td>
<td>95% CI</td>
<td>p-value</td>
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<td>No. Acute Stressors (AS)</td>
<td>1.03</td>
<td>0.98-1.08</td>
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<td>1.03</td>
<td>0.99-1.08</td>
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<td>No. Chronic stressors (CS)</td>
<td>1.00</td>
<td>0.90-1.10</td>
<td>0.95</td>
<td>1.00</td>
<td>0.90-1.11</td>
<td>0.98</td>
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<td>MODEL 2</td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>No. AS - mild</td>
<td>1.03</td>
<td>0.98-1.09</td>
<td>0.27</td>
<td>1.04</td>
<td>0.98-1.10</td>
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<tr>
<td>No. AS - moderate</td>
<td>1.03</td>
<td>0.92-1.15</td>
<td>0.63</td>
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<td>1.03</td>
<td>0.81-1.29</td>
<td>0.84</td>
<td>1.00</td>
<td>0.78-1.28</td>
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<td>0.87</td>
<td>0.52-1.44</td>
<td>0.58</td>
<td>0.95</td>
<td>0.57-1.56</td>
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<td>No. CS - mild</td>
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<td>0.88-1.11</td>
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<td>0.99</td>
<td>0.88-1.12</td>
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<td>0.93</td>
<td>0.73-1.18</td>
<td>0.53</td>
<td>0.93</td>
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<td>1.21</td>
<td>0.75-1.94</td>
<td>0.44</td>
<td>1.18</td>
<td>0.73-1.90</td>
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<td>1.58</td>
<td>0.72-3.48</td>
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<td>MODEL 3</td>
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<td>No. AS - mild-moderate</td>
<td>1.03</td>
<td>0.98-1.08</td>
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<td>1.04</td>
<td>0.99-1.09</td>
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<td>0.79-1.20</td>
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<td>0.88-1.08</td>
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<td>0.88-1.85</td>
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<td>MODEL 4</td>
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<tr>
<td>One or more AS - high-severe</td>
<td>1.33</td>
<td>0.88-2.00</td>
<td>0.18</td>
<td>1.25</td>
<td>0.82-1.91</td>
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<td>0.91-2.46</td>
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<td>MODEL 5 Social Support</td>
<td>1.01</td>
<td>0.98-1.04</td>
<td>0.31</td>
<td>1.00</td>
<td>0.97-1.03</td>
<td>0.97</td>
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<tr>
<td>MODEL 6 Optimism</td>
<td>1.04</td>
<td>1.00-1.08</td>
<td>0.08</td>
<td>1.03</td>
<td>0.99-1.08</td>
<td>0.18</td>
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<tr>
<td>MODEL 7 Anti-Emotionality</td>
<td>0.83</td>
<td>0.57-1.20</td>
<td>0.32</td>
<td>0.84</td>
<td>0.57-1.25</td>
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<tr>
<td>MODEL 8 Anger</td>
<td>1.00</td>
<td>0.94-1.07</td>
<td>0.92</td>
<td>1.00</td>
<td>0.93-1.07</td>
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<tr>
<td></td>
<td>No. AS - mild-moderate</td>
<td>No. AS - high-severe</td>
<td>No. CS - mild-moderate</td>
<td>No. CS - high-severe</td>
<td>Social Support</td>
<td>Optimism</td>
</tr>
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<td>0.98-1.09</td>
<td>0.21</td>
<td>1.05</td>
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<td>0.77-1.21</td>
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<td>0.73-1.19</td>
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<td>0.90-1.12</td>
<td>0.94</td>
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<td>0.85-1.95</td>
<td>0.23</td>
<td>1.36</td>
<td>0.91-2.04</td>
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</table>

AS=Acute stressors; CS=Chronic stressors
Mutation status (BRCA1, BRCA2, negative BRCA1/2, untested/unknown) was specified as a stratification variable in all unadjusted and adjusted analyses.

# Adjusted for (as known at last data collection): No. of 1st & 2nd degree relatives with a) BCa and b) ovarian cancer; age at menarche, no. live births, parity by breast feeding (nulliparous, parous and breast fed, parous and not breast fed), smoking history (current, former, never), exercise (sufficiently active, insufficiently active, sedentary, unknown); and adjusted for (time-varying): bilateral salpingo-oophorectomy (BSO), benign breast disease, HRT use (never, not within last 3 years, within last 3 years, unknown), oral contraception use (never, not within 3 years, within last 3 years, unknown), body mass index. Models 5 through 9 additionally control for time-varying anxiety and depression.
Author/s:

Title:
Does stress increase risk of breast cancer? A 15-year prospective study

Date:
2018-08-01

Citation:

Persistent Link:
http://hdl.handle.net/11343/283886