Joint effects of nulliparity and other breast cancer risk factors

S Opdahl*,1, MDK Alsaker^{1,2}, I Janszky^{1,3}, PR Romundstad¹ and LJ Vatten¹

Department of Public Health and Community Medicine, Faculty of Medicine, Norwegian University of Science and Technology, MTFS, Trondheim N-749 I, Norway; ²Department of Oncology, St. Olavs Hospital, Trondheim N-7006, Norway; ³Department of Public Health Sciences, Karolinska Institutet, Stockholm SE-171 77, Sweden

BACKGROUND: Pregnancy may reduce breast cancer risk through induction of persistent changes of the mammary gland that make the breast less susceptible to carcinogenic factors. It is not known to what extent the effects of parity are independent of other breast cancer risk factors.

METHODS: In a Norwegian cohort of 58191 women (2890 breast cancers), we assessed whether the effects of parity on postmenopausal breast cancer risk may be modified by menstrual and anthropometric factors. We calculated attributable proportions due to interaction as a measure of synergism.

RESULTS: Parity, height, body mass index (BMI), age at menarche and menopause were all associated with breast cancer risk in the expected directions. For BMI, follow-up was stratified into two age groups because of non-proportional hazards. We found that nulliparity and overweight may amplify each other's effect on breast cancer risk among women after 70 years of age (attributable proportion 0.21, 95% confidence interval 0.04-0.39). There was some indication that parity and age at menopause may antagonise each other's effect. Effects of parity were largely unaffected by age at menarche and height.

CONCLUSION: Nulliparity and overweight may have a synergistic effect on breast cancer risk in elderly women. If confirmed by others, the findings may help disentangle the interplay of different causes of breast cancer.

British Journal of Cancer (2011) 105, 731-736. doi:10.1038/bjc.2011.286 www.bjcancer.com

Published online 2 August 2011 © 2011 Cancer Research UK

Keywords: breast cancer risk; nulliparity; synergism; attributable proportion; epidemiology

The protective effect of pregnancies against breast cancer was first described as early as 1926 (Press and Pharoah, 2010), and was confirmed through a large number of subsequent studies (Kelsey et al, 1993). The underlying mechanisms may include differentiation of the mammary epithelial cells, reduced number of mammary stem cells, altered mammary response to oestrogen, and reduced levels of circulating hormones (Britt et al, 2007).

If the breast undergoes permanent changes during pregnancy, it is possible that the mammary gland's susceptibility to other important exposures could also be altered by pregnancy-related factors (Britt et al, 2007). However, it is unknown whether effects of parity can be modified by other risk factors for breast cancer. If such interactions were identified, our understanding of possible underlying mechanisms may be advanced, which could eventually lead to better preventive strategies.

Relatively few studies have simultaneously assessed the separate and joint effects of parity and other established risk factors, including age at menarche and menopause (La Veccia et al, 1992; Hirose et al, 2003), and body mass index (BMI) and height (Van den Brandt et al, 2000; Hirose et al, 2003). Only one of the studies was based on prospectively collected data (Van den Brandt et al, 2000), and the testing of heterogeneity of relative risks that was performed in previous studies may be of limited use in assessing causal inference related to interactions (Ahlbom and Alfredsson, 2005; Greenland et al, 2008).

Therefore, we have studied whether effects of parity on breast cancer risk may be modified by menstrual or anthropometric factors in a long-term follow-up of a large historical cohort.

MATERIALS AND METHODS

Study population and follow-up

The study population consists of women from three Norwegian counties (Nord-Trøndelag, Vestfold, and Aust-Agder) who were born between 1886 and 1928. They were invited to a breast cancer screening organised by the Norwegian Cancer Society between 1956 and 1959. The study has been described in detail elsewhere (Kvåle et al, 1987). Briefly, the participants were offered a clinical examination by a medical doctor, and information on reproductive history and demographic factors was collected during structured interviews, conducted by trained personnel. In the interview, information was recorded on age at menarche, number of full-term pregnancies, age at first full-term pregnancy, previous surgery or disease of the breasts or genital organs, age at menopause, marital status, place of residence, and the participant's own or her husband's occupation.

Information on anthropometric factors was added to the database following a compulsory mass examination for tuberculosis that was conducted between 1963 and 1975. That examination

^{*}Correspondence: Dr S Opdahl; E-mail: signe.opdahl@ntnu.no Received 9 March 2011; revised 1 July 2011; accepted 8 July 2011; published online 2 August 2011



included height measured to the nearest centimetre and weight measured to the nearest kilogram on regularly calibrated scales (Tretli, 1989).

At the population census in 1960, a unique 11-digit identity number was assigned to all Norwegian citizens. We used the identity number to link study participants to information on breast cancer incidence at the Cancer Registry of Norway, which also includes information on vital status and emigration, as recorded by the Population Registry at Statistics Norway. Invasive breast cancer was registered according to the International Classification of Diseases, 7th edition (code 170). Among 84 981 invited women who were alive at the census in 1960, 63 041 (74.2%) women had participated at the screening. Among 50 061 (79.4%) of the participating women, anthropometric information was also available.

We started breast cancer follow-up from the age of 55 years. After exclusion of women who died or emigrated before 55 years of age, 62 108 women were eligible for follow-up. Participants were followed from 1 January 1961, or from the age of 55 years, and in analyses that included anthropometric measures, women were followed from the year after the height and weight measurements, or from 55 years of age. End of follow-up was set to the date of breast cancer diagnosis, emigration, death, or to the end of follow-up on 31 December 2008, whichever occurred first.

Among eligible women, we excluded 759 women with a previous breast cancer diagnosis and 3158 women with missing information on the main study factors. A total of 58 191 women were therefore followed up for breast cancer occurrence. Among these women, 20748 were postmenopausal at the time of the interview and had reported age at menopause. A total of 451 women who reported a surgical menopause (bilateral oophorectomy and/or hysterectomy) were excluded from the analyses related to age at menopause. Thus, the analyses of age at menopause were restricted to 20 297 women with a natural menopause. Information on anthropometric factors was not available for all participants, and among 49624 eligible women with height and weight measurements, we similarly excluded women with prevalent breast cancer, women with missing information on parity or potentially confounding factors, and women with unreliable height measurements (wearing shoes, abnormal posture). Analyses that included anthropometric measures were therefore based on follow-up of 44 952 women. The number of exclusions and the number of women included in the analyses are summarised in Table 1.

Table I Summary of the number of women eligible, the exclusions made and the number of women included in analyses

	Exposure				
	Age at menarche	Age at natural menopause	Height and BMI		
Participants (n) Eligible for analyses (n) ^a	63 04 I 62 108	63 04 l 62 108	50 06 I 49 624		
Exclusions (n) Previous breast cancer diagnosis Missing information ^b Exposure not applicable ^c Surgical menopause Unreliable measurement	759 3158	759 3158 40 601 451	65 l 2332 1689		
Included in analyses (n)	58 191	20 297	44 952		

Abbreviation: BMI = body mass index. ^aWomen who could be followed from age 55 (excluding women who died or emigrated before age 55). ^bMissing information on parity or covariates. ^cWomen who were premenopausal at the time of the interview.

Parity and other risk factors

We categorised parity as nulliparous (no full-term pregnancy) or parous (one or more full-term pregnancies). In analyses of trend, age at menarche was categorised as ≤ 13 , 14, 15, ≥ 16 years, age at natural menopause as <45, 45-49, 50-54, ≥ 55 years, height as <155, 155-159, 160-164, 165-169, ≥ 170 cm, and BMI as <25, 25-29, ≥ 30 kg m⁻². When evaluating synergism, menstrual and anthropometric exposures were dichotomised at the median value (age at menarche <15 or ≥ 15 years, age at menopause <50 or ≥ 50 years, height <161 or ≥ 161 cm and BMI <26 or ≥ 26 kg m⁻²). Among parous women, number of full-term pregnancies was categorised as 1, 2, 3, ≥ 4 and age at first full-term pregnancy as <20, 20-24, 25-29, 30-34, ≥ 35 years.

Statistical analyses

We used Cox proportional hazards models to compute hazard ratios (HRs) with 95% confidence intervals (CIs). The associations of menstrual and anthropometric factors with breast cancer risk were assessed both overall and within the two strata of parity. For each exposure, the category with the largest number of person years was chosen as the reference category. The *P*-values for trend across exposure categories were calculated by treating the categories as a continuous variable. We assessed heterogeneity of the HRs across strata of parity by likelihood ratio tests by comparing models with and without product terms between parity and each exposure variable (where exposure categories were treated as a continuous variable).

Statistical interaction in multiplicative models has no direct causal interpretation, whereas departure from additivity of risks may imply causal interaction (Ahlbom and Alfredsson, 2005; Greenland *et al*, 2008). Therefore, we calculated the attributable proportion due to interaction (API) with 95% CIs to evaluate whether effects of parity could be modified by menstrual or anthropometric factors (Hosmer and Lemeshow, 1992; Andersson *et al*, 2005). The API allows the assessment of effect modification on an additive scale using multiplicative models. An API with a value greater than 0 implies synergism, that is, the effect of the combined exposure is greater than expected from their separate effects, whereas a value below 0 indicates antagonism, that is, less influence by the joint exposure than expected from their separate effects (Andersson *et al*, 2005).

In all analyses, age was controlled for by using it as the time scale in the regression models. In addition, we adjusted for birth cohort (in 5-year categories), marital status (ever or never), county, urban or rural place of residence, and the participant's own or her husband's occupation (professional/private enterprise, manual work, domestic and other work). Age at menarche was considered a potentially confounding factor in analyses of age at natural menopause, height, and BMI, and these analyses were therefore also adjusted for age at menarche. In a separate analysis restricted to parous women, we controlled for potential confounding by number of full-term pregnancies and age at first full-term pregnancy, in the trend analyses of age at menarche, age at natural menopause, height and BMI.

Proportionality between hazards was checked by comparing log minus log plots of survival and by performing tests based on Schoenfeld residuals. Assumptions were met for all exposures except BMI, for which the log minus log survival curves were non-parallel from 70 years and onwards. Others have also found that the positive association of BMI among postmenopausal women may be more pronounced among older women (Yong *et al*, 1996; Galanis *et al*, 1998). On the basis of these previous observations and on the observed non-proportionality, analyses of BMI were stratified by follow-up time into two periods, one from 55–69 years of age, and the other with follow-up from 70 years of age. Each woman contributed person-years to one or both periods,



depending on her age at the start and end of follow-up. After stratification, the proportional hazards assumption was met in both age groups.

All analyses were conducted using Stata version 11 for Windows (Stata Corp., College Station, Texas).

Ethical considerations

The study was approved by the regional committee for medical research ethics and by the Norwegian Data Inspectorate.

RESULTS

A total of 58 191 women without breast cancer at baseline were followed for 1 400 436 person years with a mean follow-up of 24.1 years (standard deviation, s.d.: 9.7). During follow-up, 2890 women were diagnosed with invasive breast cancer. Mean age at diagnosis was 72.9 years (s.d. 9.4), and overall, 48 945 (84.1%) women died during follow-up. Characteristics of the participants at baseline are shown in Table 2. Compared with parous women, nulliparous women were older and had a lower BMI. They were also more likely to be unmarried and live in an urban community, and less likely to be employed in manual work (the participant's own or her husband's occupation).

The HRs for menstrual and anthropometric risk factors stratified by parity are presented in Table 3. Nulliparous women were at higher risk for breast cancer than parous women. For nulliparous and parous women combined, breast cancer risk was negatively associated with age at menarche, and positively associated with height, age at natural menopause, and BMI (results

Among both nulliparous and parous women, breast cancer risk increased with decreasing age at menarche. Breast cancer risk increased with increasing height among parous women, whereas the trend was less clear among nulliparous women.

For age at menopause, there was a strong positive association with breast cancer risk among parous women, but no clear trend among nulliparous women.

Using the entire follow-up period, there was a positive association of BMI with breast cancer risk in both strata of parity (results not shown). After stratification by age, BMI was not associated with breast cancer risk in the age group 55-69 years. However, for women 70 years and older, there was a clear positive association of BMI with breast cancer risk.

The HRs for age at menarche, height and BMI were homogenous across the two strata of parity, whereas the trend of increasing risk with increasing age at menopause was more pronounced among parous women.

The assessments of synergism between parity and age at menarche, age at natural menopause, height, and BMI are presented in Table 4. We found that the effects of age at menarche and height did not substantially differ for nulliparous and parous women. There was some indication that nulliparity and late age at menopause could decrease each other's effect, although the statistical power to detect such an antagonistic interaction was limited (API -0.24, 95% CI -0.55, 0.08). For the combined effects of nulliparity and high BMI, the results indicated independent effects in the age group 55-69 years, and synergistic effects after 70 years of age (API 0.21, 95% CI 0.04, 0.39).

Adjustments for potentially confounding factors had no material influence on the results. Among parous women, controlling for number of full-term pregnancies and age at first full-term pregnancy did not influence the associations of age at menarche, age at natural menopause, height, or BMI with breast cancer risk (results not shown).

Table 2 Characteristics at baseline of 58 191 women at risk of breast cancer, by parity

	Nulliparous women $(n = 10765)^a$	women	Total (n = 58 191)
Mean age in 1961 (s.d.) ^c Mean age at height and weight measurement (s.d.) ^d	51.7 (11.0) 58.8 (10.4)		` '
Mean age at menarche (s.d.) Mean age at natural menopause (s.d.) Mean height, cm (s.d.) Mean body mass index, kg m ⁻² (s.d.)	` '	\ /	` '
Marital status (%) Never married Ever married		548 (1.2) 46 878 (98.8)	6229 (10.7) 51 962 (89.3)
County (%) Nord-Trøndelag Aust-Agder Vestfold	2816 (26.2)	9993 (21.1)	19 343 (33.2) 12 809 (22.0) 26 039 (44.7)
Community of residence (%) Urban Rural			12 374 (21.3) 45 817 (78.7)
Occupation (own or husband's, %) Professional, private enterprise Manual Domestic and other work	2890 (26.8)	22 048 (46.5)	16 164 (27.8) 24 938 (42.9) 17 089 (29.4)

^aWomen with no full-term pregnancy. ^bWomen with one or more full-term pregnancies. ^cAge in 1961 is not equivalent to age at start follow-up for all participants. ^dAge at height and weight measurement is not equivalent to age at start follow-up for all participants.

DISCUSSION

In this prospective cohort study, we found that overweight may enhance the effect of nulliparity on breast cancer risk in elderly women. The results suggest that one out of five breast cancer cases among nulliparous and overweight women may be attributed to a synergistic effect of these factors.

Although there has been extensive research on the association of menstrual and anthropometric factors on breast cancer risk (Kelsey et al, 1993; Friedenreich, 2001), most studies have not shown results stratified by parity. In a meta-analysis of seven cohort studies, interaction between parity and anthropometric factors was assessed on a multiplicative scale, and showed no evidence for interaction in relation to breast cancer risk (Van den Brandt et al, 2000). Also, the results of a case-control study from Japan suggested no interaction on a multiplicative scale between parity and menstrual and anthropometric factors (Hirose et al, 2003). In contrast to our results, the Japanese investigators found no effect of age at menarche, but a strong positive association of age at menopause among nulliparous women. In a case-control study from Italy, the associations of menstrual factors with breast cancer risk within groups of parity were similar to the results of the Japanese study (La Veccia et al, 1992). Our results partly confirm these earlier observations, we found no evidence that the relative risks for breast cancer associated with age at menarche, height and BMI were different between parous and nulliparous women, as the tests for interaction were not statistically significant on a multiplicative scale. However, absence of association modification on a multiplicative scale does not exclude effect modification in a causal sense (Greenland et al, 2008). In the case of parity and BMI, our results suggest that the two factors do not act independently, but rather amplify each other's effect. On the



Table 3 Risk factors for breast cancer by parity, in postmenopausal women from three counties in Norway

	Nulliparous women ^a			Parous women ^b					
	No. of person-years	No. of cases	Rate ^c	Hazard ratio (95% confidence interval)	No. of person-years	No. of cases	Rate	Hazard ratio (95% confidence interval)	P _{LR test} d
Parity	248 382	636	256	1.26 (1.12, 1.42)	1152054	2254	196	I (reference)	
Age at menarch	he, years ^e								
I3	68 436	202	295	1.24 (1.02, 1.64)	337 12	731	217	1.12 (1.01, 1.24)	
14	83 659	201	240	l (reference)	374 377	732	196	l (reference)	
15	54746	142	259	1.08 (0.87, 1.34)	258 930	475	183	0.94 (0.83, 1.05)	
≥16	41 541	91	219	0.92 (0.72, 1.18)	181 635	316	174	0.89 (0.78, 1.02)	
P_{trend}^{f}		, ,	2.7	0.04	101 055	3.0	.,.	< 0.001	0.88
Height, cm ^g									
< 155	21718	56	258	0.90 (0.65, 1.23)	98 269	167	170	0.73 (0.61, 0.86)	
155-159	40 665	111	273	0.99 (0.77, 1.27)	226 554	436	192	0.85 (0.75, 0.96)	
160-164	52 839	142	269	I (reference)	291 489	642	220	I (reference)	
165-169	30 739	87	283	1.09 (0.83, 1.42)	171 802	359	209	0.98 (0.86, 1.11)	
≥ 170	13 004	36	277	1.11 (0.77, 1.61)	53 799	137	255	1.22 (1.01, 1.47)	
P _{trend}	13001	30	2//	0.24	33777	137	233	< 0.0001	0.23
Age at menopa	iuse, years ^g								
<45	17557	37	211	0.85 (0.57, 1.26)	51 452	63	122	0.62 (0.47, 0.84)	
45-49	38 682	102	264	1.02 (0.77, 1.36)	122 937	192	156	0.78 (0.64, 0.94)	
50-54	39 408	104	264	I (reference)	131 848	272	206	I (reference)	
≥55	2548	7	275	1.01 (0.47, 2.17)	10 560	31	294	1.34 (0.93, 1.96)	
P _{trend}	23 10	,	2/3	0.55	10 300	31	271	< 0.0001	0.04
Breast cance	er risk among wo	men follow	ed from ag	ge 55 to 69 years					
Body mass inde	ex, kg m ^{-2s}								
< 25	32 180	64	199	I (reference)	161 590	233	144	I (reference)	
25-29	24 434	43	176	0.88 (0.59, 1.30)	149 325	240	161	1.13 (0.94, 1.36)	
≥30	8638	19	220	1.09 (0.64, 1.85)	65 233	108	166	1.17 (0.93, 1.49)	
P_{trend}				0.99		,,,,		0.14	0.60
Breast cance	er risk among wo	men follow	ed from ag	ge 70 years or older					
Body mass inde			•	•					
< 25	43019	106	246	I (reference)	180 234	384	213	I (reference)	
25-29	37 388	139	372	1.50 (1.16, 1.94)	199 634	494	247	1.17 (1.02, 1.34)	
≥30	13619	61	448	1.76 (1.27, 2.44)	87 60 1	287	328	1.56 (1.33, 1.83)	
/									

^aWomen with no full-term pregnancy. ^bWomen with one or more full-term pregnancies. ^cIncidence per 100 000 person-years. ^dP-value from likelihood ratio test (comparison of model with and without interaction term). ^eAdjusted for age, birth cohort, county of residence, urban or rural community of residence, marital status, occupation (own or husband's). ^fP-value for trend across exposure categories. ^gAdjusted for age, birth cohort, county of residence, urban or rural community of residence, marital status, occupation (own or husband's) and age at menarche.

other hand, parity and age at menopause showed a possible antagonistic interaction, both on the multiplicative and, more importantly, on the additive scale.

The interaction that we observed may advance our understanding of how different risk factors can act together. If confirmed by others, the finding might also be relevant for preventive strategies (Thompson, 1991), as our results suggest that maintaining a healthy body weight could potentially prevent a relatively higher proportion of breast cancer cases among nulliparous than among parous women.

A possible synergism between nulliparity and obesity suggests that there might be pathways where both factors are necessary for the development of breast cancer (Ahlbom and Alfredsson, 2005; Greenland et al, 2008). A high BMI is associated with higher production of oestrogen in adipose tissues, altered mammary stromal environment and higher insulin levels, and it has been suggested that these factors may mediate the positive association of BMI with breast cancer risk in postmenopausal women (Friedenreich, 2001; McCready et al, 2010). Results from a recent study suggested that insulin may mediate most of the increased risk associated with a high BMI (Gunter et al, 2009). Insulin is

believed to increase breast cancer risk through mitogenic, and not mutagenic mechanisms, and may therefore act at a late stage in cancer development (Vineis *et al*, 2010). A synergism between BMI and nulliparity seems consistent with current models for breast cancer development (Siemiatycki and Thomas, 1981; Vineis *et al*, 2010), as nulliparous women may have a higher number of breast cells with a malignant potential (Britt *et al*, 2007), and overweight may promote cell growth (Vineis *et al*, 2010).

The positive association of BMI with breast cancer risk, and the synergism between BMI and nulliparity, were restricted to older postmenopausal women. A stronger association of BMI with breast cancer risk among older postmenopausal women has been described previously, and might be explained by a higher cumulative exposure due to longer duration of obesity, or that a certain induction period may be necessary (Yong *et al*, 1996; Galanis *et al*, 1998).

Late age at menopause is believed to increase breast cancer risk through a longer period of exposure to ovarian sex steroids (Kelsey *et al*, 1993). It has been suggested that nulliparous women may be more vulnerable to the carcinogenic effects of oestrogens (Russo and Russo, 2006), and therefore, the indication for a possible

	No. of person-years	No. of cases	R ate ^a	Hazard ratio (95% CI)
Age at menarche and parity ^b				
Late menarche (≥ 15 years) and parous ^c	440 565	791	180	I (reference)
Early menarche (< 15 years) and parous	711 489	1463	206	1.15 (1.05, 1.25)
Late menarche (≥ 15 years) and nulliparous ^d	96 286	233	242	1.30 (1.10, 1.54)
Early menarche (<15 years) and nulliparous API (95% CI) ^e	152 096	403	265	1.42 (1.23, 1.64) -0.02 (-0.18, 0.15)
Height and parity ^f				,
Short (< 161 cm) and parous	387 264	751	194	I (reference)
Tall (≥ 161 cm) and parous	454 648	990	218	1.18 (1.07, 1.30)
Short (<161 cm) and nulliparous	74511	195	262	1.16 (1.07, 1.30)
Tall (≥161 cm) and nulliparous	84 454	237	281	1.50 (1.27, 1.78)
API (95% CI)	TCF FO	237	201	-0.01 (-0.20, 0.19)
Age at menopause and parity ^f				
Early menopause (<50 years) and parous	174 389	255	146	I (reference)
Late menopause (≥50 years) and parous	142 408	303	213	1.39 (1.17, 1.66)
Early menopause (<50 years) and nulliparous	56 239	139	247	1.55 (1.21, 1.98)
Late menopause (≥50 years) and nulliparous	41 957	111	265	1.57 (1.20, 2,05)
API (95% CI)				-0.24 (-0.55, 0.08)
Breast cancer risk among women followed fr <i>BMI and parity</i> ^f	om age 55 to 69 years			
Normal BMI ($<26 \mathrm{kg}\mathrm{m}^{-2}$) and parous	200 041	293	146	I (reference)
High BMI (\geq 26 kg m ⁻²) and parous	176 106	288	164	1.13 (0.96, 1.34)
Normal BMI ($<26 \mathrm{kg}\mathrm{m}^{-2}$) and nulliparous	39 062	73	187	1.17 (0.86, 1.59)
High BMI ($\geq 26 \text{ kg m}^{-2}$) and nulliparous	26 190	53	202	1.29 (0.92, 1.80)
API (95% CI)	20170	33	202	-0.01 (-0.38, 0.36)
Breast cancer risk among women followed fr	om age 70 years or older			
BMI and parity ^f				
Normal BMI ($<$ 26 kg m ^{-2}) and parous	228 59 1	507	222	I (reference)
High BMI ($\geqslant 26 \text{ kg m}^{-2}$) and parous	238 878	658	275	1.23 (1.09, 1.39)
Normal BMI ($<26 \mathrm{kg}\mathrm{m}^{-2}$) and nulliparous	53 035	141	266	1.25 (1.00, 1.55)
High BMI (≥26 kg m ⁻²) and nulliparous	40 992	165	403	1.88 (1.53, 2.30)
API (95% CI)				0.21 (0.04, 0.39)

Table 4 Assessment of synergism between parity and other breast cancer risk factors among postmenopausal women from three counties in Norway

Abbreviations: API = attributable proportion due to interaction; BMI = body mass index, CI = confidence interval. ^aIncidence per 100 000 person-years. ^bAdjusted for age, birth cohort, county of residence, urban or rural community of residence, marital status, occupation (own or husbands). ^cParous women defined as women with one or more full-term pregnancies. ^dNulliparous women defined as women with no full-term pregnancy. ^eAttributable proportion due to interaction. ^fAdjusted for age, county of residence, urban or rural community of residence, marital status, occupation (own or husbands) and age at menarche.

antagonism between a late age at menopause and nulliparity that we observed, was unexpected. The finding should also be interpreted with caution because of limited statistical power. In general, an antagonism between two factors may indicate competition between factors in pathways where only one of the factors is a necessary cause of the disease (Greenland *et al.*, 2008).

The high attendance and prospective nature of our study minimise the possibility for bias in selection or information, although non-differential misclassification of the collected information would be expected and most likely result in conservative estimates of effect. Reporting of new cancers to the Cancer Registry of Norway is mandatory by law, and the data of the Cancer Registry are linked to the national Cause of Death Registry, which ensures a nearly complete registration of cancer cases (Lund, 1981). The women in this study are too old to have experienced any organised mammography screening programme (Kalager *et al*, 2009), and therefore, most if not all cancer cases, were clinically detected.

We had no information on family history of breast cancer, alcohol intake, physical activity, or use of exogenous hormones (oral contraceptives or hormone replacement therapy), all of which could have potentially confounded the results. However, given the birth year of the youngest women, the use of oral contraceptives in the cohort is likely to be negligible, and hormone replacement therapy may be relevant only for a small proportion of the youngest participants (Bergsjø, 1984). Although we adjusted for several potential confounders in the multivariable analyses, we

cannot exclude the possibility of uncontrolled confounding. Nevertheless, any remaining confounder potentially able to influence our results considerably would need to (1) be strongly associated with breast cancer risk and the exposures that we studied, and (2) be unrelated to the potential confounders that were included in our models.

In the evaluation of synergism, we carried out five tests of interactions. Although these tests were dependant, caution is needed when interpreting our findings. In general, considerably higher statistical power is required to detect interactions than to detect single exposure—disease associations. Our study included a large number of elderly nulliparous women and an exceptionally long follow-up. Nevertheless, the statistical power to detect interactions was limited.

In conclusion, our findings suggest a possible synergistic effect of nulliparity and BMI in elderly women on breast cancer risk. If confirmed by others, this finding may advance our understanding of breast cancer, and might eventually lead to better preventive strategies.

ACKNOWLEDGEMENTS

The study was financially supported by the Norwegian Research Council. We thank the Cancer Registry of Norway for providing the data, the Norwegian Cancer Society for carrying out the original study, and the attending women for participating in the study.



REFERENCES

- Ahlbom A, Alfredsson L (2005) Interaction: a word with two meanings creates confusion. Eur J of Epidemiol 20: 563 564
- Andersson T, Alfredsson L, Kallberg H, Zdravkovic S, Ahlbom A (2005) Calculating measures of biological interaction. Eur J Epidemiol 20: 575-579
- Bergsjø P (1984) Comments on the use of oestrogens 1979-83. In Drug Consumption in Norway, Øydvin K (ed), pp 114-115. Norsk medisinaldepot: Oslo
- Britt K, Ashworth A, Smalley M (2007) Pregnancy and the risk of breast cancer. Endocr Relat Cancer 14: 907 933
- Friedenreich CM (2001) Review of anthropometric factors and breast cancer risk. Eur J Cancer Prev 10: 15-32
- Galanis DJ, Kolonel LN, Lee J, Le Marchand L (1998) Anthropometric predictors of breast cancer incidence and survival in a multi-ethnic cohort of female residents of Hawaii, United States. Cancer Causes Control 9: 217-224
- Greenland S, Lash TL, Rothman KJ (2008) Concepts of interaction. In *Modern Epidemiology*, Rothman KJ, Greenland S, Lash TL (eds), pp 71-83. Lippincott Williams & Wilkins: Philadelphia
- Gunter MJ, Hoover DR, Yu H, Wassertheil-Smoller S, Rohan TE, Manson JE, Li J, Ho GYF, Xue X, Anderson GL, Kaplan RC, Harris TG, Howard BV, Wylie-Rosett J, Burk RD, Strickler HD (2009) Insulin, insulin-like growth factor-1, and risk of breast cancer in postmenopausal women. J Natl Cancer Inst 101: 48 60
- Hirose K, Tajima K, Hamajima N, Takezaki T, Inoue M, Kuroishi T, Miura S (2003) Impact of established risk factors for breast cancer in nulligravid Japanese women. *Breast Cancer* **10:** 45-53, doi: 10.1007/BF02967625
- Hosmer DW, Lemeshow S (1992) Confidence interval estimation of interaction. *Epidemiology* **3:** 452-456
- Kalager M, Haldorsen T, Bretthauer M, Hoff G, Thoresen SO, Adami HO (2009) Improved breast cancer survival following introduction of an organized mammography screening program among both screened and unscreened women: a population-based cohort study. Breast Cancer Res 11: R44
- Kelsey JL, Gammon MD, John EM (1993) Reproductive factors and breast cancer. Epidemiol Rev 15: 36-47

- Kvåle G, Heuch I, Eide GE (1987) A prospective study of reproductive fators and breast cancer. *Am J Epidemiol* 126: 831 841
- La Veccia C, Negri E, Bruzzi P, Dardanoni G, Decarli A, Franceschi S, Palli D, Talamini R (1992) The role of age at menarche and age at menopause on breast cancer risk: Combined evidence from four case-control studies. Ann Oncol 3: 625 – 629
- Lund E (1981) Pilot study for the evaluation of completeness of reporting to The Cancer Registry. In *Incidence of Cancer in Norway 1978*, pp 11–14. The Cancer Registry of Norway: Oslo
- McCready J, Arendt LM, Rudnick JA, Kuperwasser C (2010) The contribution of dynamic stromal remodelling during mammary development to breast carcinogenesis. *Breast Cancer Res* 12: 205
- Press DJ, Pharoah P (2010) Risk factors for breast cancer. A reanalysis of two case-control studies from 1926 and 1931. *Epidemiology* 21: 566-572 Russo J, Russo IH (2006) The role of estrogen in the initiation of breast cancer. *J Steroid Biochem Mol Biol* 102: 89-96
- Siemiatycki J, Thomas DC (1981) Biological models and statistical interactions: an example from multistage carcinogenesis. Int J Epidemiol 10: 383-387
- Thompson WD (1991) Effect modification and the limits of biological inference from epidemiological data. *J Clin Epidemiol* 44: 221 232
- Tretli S (1989) Height and weight in relation to breast cancer mobidity and mortality. A prospective study of 570,000 women in Norway. *Int J Cancer* 44: 23 30
- Van den Brandt PA, Spiegelman D, Yaun SS, Adami HO, Beeson L, Folsom AR, Fraser G, Goldbohm RA, Graham S, Kushi L, Marshall JR, Miller AB, Rohan T, Smith-Warner SA, Speizer FE, Willett WC, Wolk A, Hunter DJ (2000) Pooled analysis of prospective cohort studies on height, weight, and breast cancer risk. *Am J Epidemiol* 152: 514–527
- Vineis P, Schatzkin A, Potter JD (2010) Models of carcinogenesis: an overview. *Carcinogenesis* 31: 1703 1709
- Yong LC, Brown CC, Schatzkin A, Schairer C (1996) Prospective study of relative weight and risk of breast cancer: the Breast Cancer Detection Demonstration Project follow-up study, 1979 to 1987–1989. *Am J Epidemiol* 143: 985–995

This work is published under the standard license to publish agreement. After 12 months the work will become freely available and the license terms will switch to a Creative Commons Attribution-NonCommercial-Share Alike 3.0 Unported License.