Birth intervals and breast cancer risk

A Kauppiälä¹, P Kyyrönen², M Hinkula¹ and E Pukkala*²,³
¹Department of Obstetrics and Gynecology, Oulu University Hospital, FI-90250 Oulu, Finland; ²Finnish Cancer Registry, Institute for Statistical and Epidemiological Cancer Research, Pieni Roobertinkatu 9, FI-00130 Helsinki, Finland; ³School of Public Health, FI-33014 University of Tampere, Tampere, Finland

BACKGROUND: The interval between successive births (birth interval) may affect breast cancer risk, whereas interval from last birth to cancer onset may modify its behaviour.

METHODS: The study cohort consisted of 29 488 Finnish grand multiparous (GM) women, including 628 women with breast cancer. Conditional logistic regression for case-control design nested within the cohort was used to estimate proportional hazards (referred as relative risks, RR). Age at first birth and parity were co-variables.

RESULTS: Short interval (<1 year) between first and second birth increased the risk of advanced ductal breast cancer at ages <50 years (RR=5.29; 95% CI 2.00–14.0) as compared to interval 3+ years. The risk of advanced ductal cancer was also large (RR = 4.00; 95% CI 1.19–13.4) shortly (<3 years) after last birth as compared with the period 15+ years.

CONCLUSIONS: Short birth interval-associated excess breast cancer risk may be related to stimulatory effects of female steroid hormones produced during two closely connected pregnancies, or defective breast maturation owing to failures in breastfeeding.

British Journal of Cancer (2009) 101, 1213–1217. doi:10.1038/sj.bjc.6605300 www.bjcancer.com
Published online 8 September 2009
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Keywords: aetiology of breast cancer; risk factor; pregnancy; age at first birth; birth interval; interval from birth to cancer

Findings in our previous study of grand multiparous women (GM, at least five births) indicated that birth interval (time between two successive births) might affect the risk of breast cancer (Hinkula et al, 2001). This finding and those of others (Albrektsen et al, 2005) prompted us to evaluate the impact of individual birth intervals on risk. Specific attention was paid to the interval between first and second births because the first pregnancy had a significant role in the maturation of the breast resistance against carcinogenic influences (Russo et al, 1994a, 2005). The length of birth interval depends also on the duration of breastfeeding, which is an important determinant of breast cancer risk (Collaborative Group of Hormonal in Breast Cancer, 2002; Ursin et al, 2005; Lord et al, 2008).

Pregnancies have dual effects on breast cancer: an immediate increase in the risk after childbirth is followed by a long-term protection (Woods et al, 1980; Kvåle and Heuch, 1987; Kelsey et al, 1993). In addition, cancers appearing within 2–6 years after birth are more frequently advanced than those in women having a long interval between birth and cancer onset (Kroman et al, 1997; Wohlfarth et al, 2001; Phillips et al, 2004; Rosenberg et al, 2004; Albrektsen et al, 2005, 2006).

This study population comprises of only GM women, most of whom belong to the religious minority, the Laestadian movement within the Lutheran church, which forbids the use of artificial contraception. Hence conception in this population takes place in physiological circumstances, thereby representing a specific advantage for our study. Here, we aimed to explore the impact of individual birth intervals on breast cancer risk at different stages. Another aim was to assess how the relative risk changes in relation to time since last birth in a Finnish cohort of GM women.

MATERIALS AND METHODS

The data of the Finnish national Population Register comprised 29 488 women born 1935 or later and registered as having at least five biological children by the end of 1997. This database involves complete linkages between parents and the children born in October 1953 or later, but precluding our obtaining full parity history for women born before 1935. The exactly registered birthdays of each child form the basis for birth interval calculations.

Breast cancer cases diagnosed among the GM women between the fifth childbirth and 31 December 2006 were identified in an automatic record linkage with the files of the national population-based Finnish Cancer Registry, using personal identifiers as the key. There were 628 such breast cancers. The Finnish Cancer Registry files include clinical stage and cancer morphology. The registry receives cancer notifications from hospitals, pathological and cytological laboratories, and also from physicians outside hospitals; its coverage is almost 100% (Teppo et al, 1994).

Statistical methods

For each breast cancer case, 50 controls were randomly selected among the GM cohort members who were at risk for breast cancer at the time of cancer onset of the case and fulfilled the matching criteria; a tolerance of ±1 year was allowed on the date of birth. The proportional hazards method was applied to this case–control
data. A woman was a non-case until she became a case, so controls for each case were selected among non-cases irrespective of whether they later became cases. The point estimates, hazard ratios were obtained in SAS 9.1 using the PHREG procedure with the option TIES = DISCRETE, which requests the discrete logistic model. This method is same as the method of the conditional logistic regression analysis if the controls were selected among the individuals who were at risk of becoming cases. Proportional hazard were referred here as relative risks (RR). Possible interactions were evaluated using the TPHREG procedure, a test release of the PHREG procedure that incorporates the CLASS statement in SAS 9.1. No interactions were found between the study variables.

The RRs were also calculated for ductal and lobular, and for local and advanced (regional or distant metastases) breast cancers separately. Further stratification was based on the age at the diagnosis of cancer (<50 years and 50+ years). Pregnancy at the age of 50 years or later is extremely rare. The younger women represent thus the years of reproduction and the older ones that of ovarian quiescence. The distribution of patients into different subcategories is presented in Table 1.

The likelihood ratio test was used to evaluate statistical significance of the parameters of interest. The trend test was analysed using linear trend test for the classified variables. Each of the four intervals between first and fifth births was classified into three categories (<1, 1–2 and 3+ complete years). Parity (5, 6, 7, 8, 9 and 10+ children) and the age at first birth (<20, 20–24, 25–29, 30+ years) were added in each model. The time from the latest birth to the date of onset of cancer was stratified into four categories (<3, 3–6, 7–14, 15+ complete years).

**RESULTS**

Overall, the RR did not differ significantly between different birth intervals, whereas short interval (<3 years) between last birth and cancer onset was associated with a significantly increased risk as compared to those with the interval of 15+ years (Table 2). Increase in the number of births and decline in age at first birth diminished the risk of breast cancer.

For cancers diagnosed before 50 years of age, the RR for short interval (<1 year) between first and second birth was increased twofold when compared with birth interval of 3+ years, and the interval from last birth to cancer onset among the cases was three times more often short (<3 years) than among the controls (Table 3). An increase in parity from 5 to 8+ nearly halved the RR in both age groups. In the group of 50+ year-old GM women, a decline in age at first birth decreased significantly the RR of breast cancer.

The most significant findings appeared in advanced ductal cancer diagnosed before age 50 years (Table 4). Short (<1 year) interval between first and second birth was associated with more than fivefold increased RR when compared with the 3+ years category. The RR of this cancer type appearing within 3 years after last birth was fourfold as compared with the 15+ years category.

**Table 2** Number of breast cancer cases (N) among grand multiparous women, and model-based multivariable relative risks (RR) with 95% confidence intervals (95% CI), by study variables

| Variable | N  | RR  | 95% CI |
|----------|----|-----|--------|
| Interval between births |    |     |        |
| 1st and 2nd |    |     |        |
| <1 year | 60 | 1.06 | 0.76–1.49 |
| 1–2.99 years | 472 | 0.91 | 0.72–1.14 |
| 3+ years | 96 | 1.00 | Ref. |
| 2nd and 3rd |    |     |        |
| <1 year | 25 | 0.78 | 0.50–1.21 |
| 1–2.99 years | 449 | 1.05 | 0.86–1.27 |
| 3+ years | 154 | 1.00 | Ref. |
| 3rd and 4th |    |     |        |
| <1 year | 18 | 1.08 | 0.66–1.78 |
| 1–2.99 years | 397 | 1.06 | 0.89–1.27 |
| 3+ years | 213 | 1.00 | Ref. |
| 4th and 5th |    |     |        |
| <1 year | 12 | 0.91 | 0.51–1.64 |
| 1–2.99 years | 326 | 1.03 | 0.86–1.22 |
| 3+ years | 290 | 1.00 | Ref. |
| Time since last birth* |    |     |        |
| <3 years | 32 | 2.36 | 1.31–4.27 |
| 3–6.99 years | 45 | 1.54 | 0.98–2.41 |
| 7–14.99 years | 164 | 1.25 | 0.96–1.63 |
| 15+ years | 387 | 1.00 | Ref. |
| Number of birthsb |    |     |        |
| 5 | 407 | 1.00 | Ref. |
| 6 | 119 | 0.79 | 0.63–0.97 |
| 7 | 41 | 0.66 | 0.47–0.93 |
| 8+ | 61 | 0.58 | 0.42–0.80 |
| Age at first birthc |    |     |        |
| <20 years | 212 | 1.00 | Ref. |
| 20–24 years | 303 | 0.97 | 0.81–1.16 |
| 25–29 years | 95 | 1.45 | 1.12–1.89 |
| 30+ years | 18 | 1.47 | 0.88–2.46 |

*Likelihood ratio test: P = 0.004, P-trend = 0.006.  
+Likelihood ratio test: P = 0.001, P-trend = 0.0001.  
*Likelihood ratio test P = 0.01, P-trend = 0.02.

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**Table 1** Number (N) and percentage of women with breast cancer in 1974–2006 among 29,488 women in Finland born 1935+ and registered to have at least five biological children, by histology, clinical stage at diagnosis, and age at diagnosis

| Breast cancer category | All ages | Age <50 years | Age 50+ years |
|------------------------|---------|--------------|--------------|
|                        | N | Percentage of all women | N | Percentage of sub-category | N | Percentage of sub-category |
| All                    | 628 | 100 | 180 | 29 | 448 | 71 |
| Histology              |    |     |     |     |     |     |
| Ductal cancer          | 484 | 77 | 140 | 29 | 344 | 71 |
| Lobular cancer         | 101 | 16 | 21 | 21 | 80 | 80 |
| Other types/unknown    | 43 | 7 | 19 | 44 | 24 | 56 |
| Clinical stage         |    |     |     |     |     |     |
| Local                  | 326 | 52 | 67 | 21 | 259 | 80 |
| Advanced               | 264 | 42 | 105 | 40 | 159 | 60 |
| Unknown                | 38 | 6 | 8 | 21 | 30 | 79 |
Table 3 Number of breast cancer cases (N) and model-based relative risks (RR) with 95% confidence interval (95% CI) in <50 and 50+ year-old grand multiparous women, by age at follow-up and by study variables

| Variable | Age at follow-up <50 years | Age at follow-up 50+ years |
|----------|-----------------------------|-----------------------------|
|          | N   | RR  | 95% CI      | N   | RR  | 95% CI      |
| Interval between births |       |     |              |       |     |              |
| 1st and 2nd birth |       |     |              |       |     |              |
| <1 year | 20  | 2.03 | 1.08–3.83 | 40  | 0.81 | 0.54–1.21 |
| 1–2.99 years | 136 | 1.29 | 0.82–2.03 | 336 | 0.80 | 0.61–1.04 |
| 3+ years | 24  | 1.00 | Ref        | 72  | 1.00 | Ref        |
| 2nd and 3rd birth |       |     |              |       |     |              |
| <1 year | 5   | 0.71 | 0.28–1.85 | 20  | 0.80 | 0.49–1.31 |
| 1–2.99 years | 129 | 1.23 | 0.85–1.78 | 320 | 0.98 | 0.78–1.24 |
| 3+ years | 46  | 1.00 | Ref        | 108 | 1.00 | Ref        |
| 3rd and 4th birth |       |     |              |       |     |              |
| <1 year | 3   | 0.85 | 0.26–2.80 | 15  | 1.14 | 0.66–1.96 |
| 1–2.99 years | 116 | 1.26 | 0.89–1.78 | 281 | 1.00 | 0.81–1.24 |
| 3+ years | 61  | 1.00 | Ref        | 152 | 1.00 | Ref        |
| 4th and 5th birth |       |     |              |       |     |              |
| <1 year | 6   | 1.68 | 0.70–4.03 | 6   | 0.64 | 0.28–1.45 |
| 1–2.99 years | 86  | 0.86 | 0.61–1.23 | 240 | 1.12 | 0.91–1.37 |
| 3+ years | 88  | 1.00 | Ref        | 202 | 1.00 | Ref        |
| Time since last birth |       |     |              |       |     |              |
| <3 years | 31  | 3.27 | 1.42–7.51 | 1    | 1.38 | 1.32–144 |
| 3–6.99 years | 40  | 2.13 | 1.07–4.26 | 5   | 1.50 | 0.58–3.90 |
| 7–14.99 years | 86  | 1.75 | 1.00–3.05 | 78  | 1.10 | 0.80–1.51 |
| 15+ years | 23  | 1.00 | Ref        | 364 | 1.00 | Ref        |
| Number of births |       |     |              |       |     |              |
| 5      | 110 | 1.00 | Ref        | 297 | 1.00 | Ref        |
| 6      | 37  | 0.90 | 0.60–1.34 | 82  | 0.73 | 0.57–0.94 |
| 7      | 12  | 0.63 | 0.33–1.21 | 29  | 0.66 | 0.45–0.99 |
| 8+     | 21  | 0.55 | 0.29–1.02 | 40  | 0.59 | 0.40–0.87 |
| Age at first birth |       |     |              |       |     |              |
| <20 years | 53  | 1.00 | Ref        | 159 | 1.00 | Ref        |
| 20–24 years | 92  | 1.04 | 0.73–1.49 | 211 | 0.93 | 0.75–1.15 |
| 25–29 years | 29  | 1.22 | 0.73–2.04 | 66  | 1.52 | 1.12–2.06 |
| 30+ years | 6   | 1.41 | 0.54–3.64 | 12  | 1.51 | 0.82–2.81 |
| Total   | 180 | 1.00 | Ref        | 448 |      |             |

P-trend = 0.04. LR test P = 0.05. P-trend = 0.008. LR test P = 0.006. P-trend = 0.001. LR test P = 0.009. P-trend = 0.04.

These factors operate separately; there was not a single breast cancer GM woman, with birth interval < 1 year and interval from last birth to cancer < 3 years. A long interval (15+ years) between last birth and cancer onset may protect against breast cancer at age 50+ years (Table 4).

DISCUSSION

This study finds that among GM women, breast cancer is associated with birth interval. Most important was the new finding that short birth interval between first and second birth was significantly associated with increased risk of advanced ductal cancer in young GM mothers. The risk of ductal breast cancer in clinically advanced stage is also high during the 3 years following the last birth.

Previous studies on the risk factors in breast cancer have comprised predominantly nulliparous women and women with few children. We widened the perspective by evaluating this topic in a homogenous group of Finnish GM women, whose breast cancer risk is low, 45% having below the average incidence (Hinkula et al, 2001).

The registers used in this study, the National Population Register for births, and the Finnish Cancer Registry, are reliable and virtually complete (Teppo et al, 1994; Pukkala, 2009). Because the most important results in this study are from ages < 50 years, the lack of information on postmenopausal hormone therapy or body weight does not weaken the validity of the findings.

The RR of advanced ductal breast cancer of GM women below 50 years of age was more than five times higher if the interval between first and second birth was < 1 year (= pregnancy interval < 3–4 months) as compared to 3+ years. The mechanism of this detrimental effect is unclear. The fact that it appeared only after the first pregnancy might reflect an association with breast maturation, because first pregnancy may have a central role in the differentiation of breast cells more resistant to carcinogenic influences (Lambe et al, 1994; Russo and Russo, 1994b; Chie et al, 2000; Russo et al, 2005; Wagner and Smith, 2003; Siwko et al, 2008). The suggestion that incomplete differentiation may...
breastfeeding, and especially no breastfeeding is associated with breastfeed or have only for a very short time. Short-term a short interval between first and second birth, do not have GM women, with increased breast cancer risk in association with of placental hormones (estrogens and progesterone) and prolactin celluar changes owing to the potential for breast cancer would initiate, in the beginning of second pregnancy, abnormal and progesterone (from corpus luteum of the second pregnancy) In such a case, the joint actions of prolactin, (induced by suckling) first childbirth is thus possible but supposedly a rare phenomenon. Fecundation of lactating GM mother within 3–4 months after her weakens especially from the fourth postpartum month onward. menstrual cycle and early fecundation after the birth. The declines rapidly, which allows early resumption of ovulatory nursing mothers, in whom each breastfeeding induces a transient principal hormone for milk biosynthesis, remains elevated only in prolactin level rises strongly from the eighth week onwards risk of breast cancer (Tworoger et al, 2003; Albrektsen et al, 2005). The most recent prospective study suggests that prolactin might increase the risk of breast cancer (Tworoger et al, 2007). During pregnancy, prolactin level rises strongly from the eighth week onwards reaching the peak concentration at term. Thereafter prolactin, the principal hormone for milk biosynthesis, remains elevated only in nursing mothers, in whom each breastfeeding induces a transient peak in its level. In non-lactating woman prolactin concentration declines rapidly, which allows early resumption of ovulatory menstrual cycle and early fecundation after the birth. The contraceptice effect of prolactin is not absolute; the efficacy weakens especially from the fourth postpartum month onward. Fecundation of lactating GM mother within 3–4 months after her first childbirth is thus possible but supposedly a rare phenomenon. In such a case, the joint actions of prolactin, (induced by suckling) and progesterone (from corpus luteum of the second pregnancy) would initiate, in the beginning of second pregnancy, abnormal cellular changes owing to the potential for breast cancer development. Alternatively, the long-term joint stimulatory actions of placental hormones (estrogens and progesterone) and prolactin during two closely consecutive pregnancies may serve as initiators for malignant transformation of epithelial breast cells. In spite of our lack of breastfeeding data, we hypothesise that GM women, with increased breast cancer risk in association with a short interval between first and second birth, do not have breastfed or have only for a very short time. Short-term breastfeeding, and especially no breastfeeding is associated with shorter birth interval (Rutstein 2005). An increase in breast cancer risk has been reported among premenopausal American women, who nursed their first (RR = 1.37) or second (RR = 1.44) child <1 month (Byers et al, 1985).

Because lactation participates in the differentiation of mammary epithelium in its terminal phase (Russo and Russo, 1994b; Russo et al, 2001, 2008), deficient breastfeeding of the first child might leave the breast cells susceptible to carcinogenic influences. This would be in accordance with the finding that lactation has a significant independent protective effect in breast cancer (Byers et al, 1985; Collaborative Group of Hormonal Factors in Breast Cancer, 2002; Ursin et al, 2005), particularly before menopause (Byers et al, 1985; Newcomb et al, 1994; Lord et al, 2008). Insufficient breastfeeding resulting in defective breast maturation might thus be the primary cause for breast cancer of young GM women in this specific subgroup.

The picture of breast cancer risk increase associated with birth intervals appeared to be more complicated than the previous one that long birth interval solely would affect the risk (Kvåle et al, 1987; Albrektsen et al, 2005, 2006). In the light of the present results the picture of adverse effects associated with birth intervals seems to be more comapre than the view that only long birth intervals affect risk (Kvåle and Heuch, 1987; Albrektsen et al, 2005, 2006).

The RR of breast cancer of young GM women was highest shortly after their last birth, which is in accordance with earlier observations of mothers with few pregnancies (Kvåle and Heuch, 1987; Leon et al, 1995; Wohlfarth et al, 2001; Albrektsen et al, 2005, 2006). Transiently increased breast cancer risk after the last childbirth in this study showed no association with the number of pregnancies. The detrimental effect of short interval extended to GM women aged 50+ years. Transient increase risk after latest birth may even last for 15 years (Lambe et al, 1994; Liu et al, 2002).

Young age at first birth and increasing parity are established protective factors in breast cancer risk (La Vecchia et al, 1989; Layde et al, 1989; Lambe et al, 1996; Merrill et al, 2005). In this study, after adjusting for other study variables, age at first birth and parity were significant factors in GM women aged 50 + years. Among GM women <50 years, our previous study of the same cohort (Hinkula et al, 2001) that did not include birth intervals and age at last birth showed, that age at first birth – but not parity – is a significant risk factor. In contrast, in this study parity is a significant risk factor among younger GM women – but not age at first birth. Thus, the new age-related variables seem to weaken the age at first birth effect and strengthen the role of parity as an independent risk factor in this age period.

Elevated risk of advanced ductal breast cancer of young GM women with short interval between first and second birth might have an association with female steroid hormones and prolactin secreted during two closely consecutive pregnancies or with defective maturation of the breast owing to inappropriate nursing.

ACKNOWLEDGEMENTS

We thank Irma H Russo and Jose Russo for their valuable advise in the preparation of the article.

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