Body size in early life and risk of epithelial ovarian cancer: results from the Nurses’ Health Studies

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Overweight and obesity have been associated with risk of cancer in women (Calle and Kaaks, 2004), but the findings for ovarian cancer are inconclusive. Some epidemiological studies have observed weak to moderate positive associations between adult body mass index (BMI) and ovarian cancer risk; others have found no association (Olsen et al, 2007). One potential explanation is that the timing of overweight and obesity during the lifecycle may be important. Although most studies have examined cancer risk in relation to recent BMI, studies examining BMI earlier in life, during late adolescence or young adulthood, have observed stronger positive associations (Fairfield et al, 2003; Engeland et al, 2004). Body fatness during childhood and early adolescence has been associated with breast cancer risk, independent of adult BMI (Berkey et al, 1999; Baer et al, 2005), suggesting that adiposity at young ages may affect risk of hormone-related cancers. Furthermore, there is evidence that height, a marker of early childhood growth and nutrition, is associated with ovarian cancer risk (Rodriguez et al, 2002; Engeland et al, 2003; Schouten et al, 2003, 2008), indicating that early life may be a critical time period for ovarian cancer initiation.

Therefore, we prospectively examined the associations of body size in early life – including body fatness at ages 5 and 10 years, BMI at the age of 18 years, birthweight, and height – with risk of epithelial ovarian cancer among participants in two large cohort studies, the Nurses’ Health Study (NHS) and the NHSII.

MATERIALS AND METHODS

Study design and population

The NHS began in 1976 and the NHSII in 1989, when 121 700 and 116 609 US female registered nurses, respectively, completed a mailed questionnaire about their lifestyle factors, health behaviours, and medical histories. Follow-up questionnaires have been sent to participants every 2 years since enrolment. Incident cases of epithelial ovarian cancer were reported on the biennial questionnaires through 2004 (NHS) and 2005 (NHSII). A gynaecologic pathologist reviewed the pathology reports and medical records to confirm the diagnosis and identify histological type, subtype, morphology, and stage (Tworoger et al, 2008).

Assessment of body size in early life and other covariates

Participants recalled their body fatness (also called ‘somatotype’), at ages 5 and 10 years using a nine-level figure drawing, where level 1 represents the most lean and level 9 represents the most overweight (Figure 1) (Stunkard et al, 1983). Among participants in the Third Harvard Growth Study, Pearsons correlations between recalled somatotype and measured BMI were 0.60 for the age of 5 years and 0.70 for the age of 10 years (Must et al, 1993). We averaged each participant’s reported somatotypes at ages 5 and 10 years to obtain an estimate of childhood body fatness. The levels 5 and above were combined in the analysis because of small numbers of participants in these categories.

Women reported their weight at the age of 18 years and their current height at enrolment; these were used to calculate BMI at the age of 18 years in kg m−2. In a sample of NHSII participants,
the Spearman's correlation between recalled and recorded weight at the age of 18 years was 0.87, and for BMI at the age of 18 years was 0.84 (Troy et al., 1995). Categories for BMI at the age of 18 years and height were chosen based on their distributions and previously used cutpoints.

Participants recalled their birthweight as <5.5, 5.5–6.9, 7.0–8.4, 8.5–9.9, and ≥10 pounds. The two highest categories (8.5 pounds and higher) were combined in the analysis to increase power. In a NHSII validation study (Troy et al., 1996), the correlation between self-reported birthweight and that obtained from state birth records was 0.74.

Age and other covariates were assessed on the questionnaires throughout the study.

### Statistical analysis

Participants contributed person-time from baseline (the questionnaire year that the exposure of interest was assessed) until the date of ovarian cancer diagnosis, report of other cancer (except nonmelanoma skin cancer), death, or 31 May 2004 (NHS) or 31 May 2005 (NHSII), whichever occurred sooner. We excluded women reporting a previous diagnosis of cancer except nonmelanoma skin cancer and those with a history of bilateral oophorectomy or pelvic irradiation. For analyses focusing on each body size measure, we excluded women who were oophorectomized or pelvic irradiation. For analyses focusing on each body size measure, we excluded women who were oophorectomized or pelvic irradiation.

Most adult characteristics were not associated with early life body size (Table 2). In both cohorts, greater body fatness during childhood and BMI at the age of 18 years were associated with later menarche and greater current BMI, and taller height was associated with later menarche. In the NHS, women with greater childhood body fatness and taller women were slightly younger at baseline, and women who were heavier in childhood and at the age of 18 years were slightly less likely to use postmenopausal hormones. In the NHSII, women with greater childhood body fatness, greater BMI at the age of 18 years, and taller height were less likely to be parous, and those with greater BMI at the age of 18 years also had shorter duration of oral contraceptive use.

In the NHS, greater body fatness at ages 5 and 10 years were associated with decreased risk of ovarian cancer (Table 3), although the association was only significant for the age of 10 years (RR for level 5 vs level 1 = 0.69, 95% CI: 0.48–0.99, P for trend = 0.01). Averaging ages 5 and 10 years, the RR for childhood body fatness level ≥5 compared to level 1 was 0.81 (95% CI: 0.53–1.24, P for trend = 0.04). In contrast, there was some suggestion of a positive association for average childhood body fatness in the NHSII (RR for level ≥5 vs level 1 = 2.09, 95% CI: 0.98–4.48), although this was not statistically significant (P for trend = 0.10) (Table 3). The associations for body fatness at ages 5 and 10 years individually and average childhood body fatness were significantly different by cohort (P for heterogeneity = 0.03, 0.01.

### RESULTS

Most adult characteristics were not associated with early life body size (Table 2). In both cohorts, greater body fatness during childhood and BMI at the age of 18 years were associated with earlier menarche and higher current BMI, and taller height was associated with later menarche. In the NHS, women with greater childhood body fatness and taller women were slightly younger at baseline, and women who were heavier in childhood and at the age of 18 years were slightly less likely to use postmenopausal hormones. In the NHSII, women with greater childhood body fatness, greater BMI at the age of 18 years, and taller height were less likely to be parous, and those with greater BMI at the age of 18 years also had shorter duration of oral contraceptive use.

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The interaction with menopausal status ($P = 0.09$), although neither these nor other factors except age were age-standardised in 5-year intervals for each cohort. Average childhood body fatness was calculated by taking the average of each participant's body fatness at ages 5 and 10 years, using a nine-level figure drawing. Baseline for each analysis was the year when the body size measure was assessed (see Table 1). Mother or sister had ovarian cancer according to the participant's response on the questionnaire; family history was evaluated using data from 1992 for NHS and 1993 for NHSII because it was not available in previous cycles.

### Table 2: Characteristics in 1988 (Nurses' Health Study) and 1989 (Nurses' Health Study II) by Body Size Measures and Study Population

|               | Average childhood body fatnessa | BMI at the age of 18 years, kg m$^{-2}$ | Height, m | Birthweight, lbs |
|---------------|--------------------------------|-----------------------------------------|-----------|-----------------|
|               | $< 5$ | $> 5$ | $< 20$ | $> 25$ | $< 1.6$ | $> 1.75$ | $< 5.5$ | $> 8.5$ |
| **Nurses' Health Study** (NHS) |    |        |        |        |        |        |        |
| Sample size at baseline | 19 371 | 4633 | 29 077 | 8212 | 24 877 | 5165 | 3312 | 6868 |
| Means |    |        |        |        |        |        |        |
| Age (years) | 55.4 | 53.9 | 54.4 | 54.6 | 55.0 | 53.8 | 53.6 | 54.9 |
| Age at menarche (years) | 12.8 | 12.2 | 12.8 | 12.2 | 12.3 | 12.9 | 12.4 | 12.6 |
| Parity (among parous women) | 3.2 | 3.0 | 3.1 | 3.1 | 3.2 | 3.0 | 3.1 | 3.1 |
| Duration of oral contraceptive use (months) | 24.6 | 24.0 | 25.2 | 23.2 | 22.9 | 23.2 | 25.9 | 24.7 |
| Current body mass index (kg m$^{-2}$) | 24.4 | 28.1 | 23.4 | 30.8 | 25.7 | 25.1 | 25.5 | 26.1 |
| **Percentages** |    |        |        |        |        |        |        |
| Parous | 93.0 | 92.3 | 92.5 | 90.8 | 92.4 | 90.5 | 92.6 | 92.5 |
| Premenopausal | 31.6 | 31.7 | 27.8 | 27.1 | 25.2 | 25.5 | 34.2 | 33.3 |
| Postmenopausal | 68.2 | 68.2 | 70.1 | 70.6 | 69.3 | 69.7 | 65.0 | 65.8 |
| Family history of ovarian cancerd | 2.6 | 2.8 | 2.5 | 2.9 | 2.2 | 2.7 | 3.0 | 2.6 |
| History of tubal ligation | 17.4 | 17.9 | 17.0 | 18.3 | 17.0 | 15.5 | 19.3 | 18.2 |
| Current postmenopausal hormone user (among postmenopausal women) | 25.3 | 22.3 | 27.8 | 20.9 | 25.2 | 26.8 | 23.4 | 24.8 |

### Table 2: Characteristics in 1988 (Nurses' Health Study) and 1989 (Nurses' Health Study II) by Body Size Measures and Study Population

|               | Average childhood body fatnessa | BMI at the age of 18 years, kg m$^{-2}$ | Height, m | Birthweight, lbs |
|---------------|--------------------------------|-----------------------------------------|-----------|-----------------|
|               | $< 5$ | $> 5$ | $< 20$ | $> 25$ | $< 1.6$ | $> 1.75$ | $< 5.5$ | $> 8.5$ |
| **Nurses' Health Study II (NHSII)** |    |        |        |        |        |        |        |
| Sample size at baseline | 19 220 | 6984 | 44 078 | 11 745 | 21 440 | 8922 | 2834 | 11 467 |
| Means |    |        |        |        |        |        |        |
| Age (years) | 34.6 | 34.8 | 34.3 | 34.3 | 34.5 | 34.0 | 35.0 | 33.7 |
| Age at menarche (years) | 12.8 | 12.0 | 12.7 | 12.0 | 12.2 | 12.7 | 12.3 | 12.5 |
| Parity (among parous women) | 2.1 | 2.0 | 2.1 | 2.0 | 2.0 | 2.0 | 2.1 | 2.1 |
| Duration of oral contraceptive use (months) | 46.9 | 43.0 | 45.6 | 39.3 | 42.7 | 44.1 | 45.7 | 43.3 |
| Current body mass index (kg m$^{-2}$) | 22.1 | 27.6 | 21.4 | 31.3 | 24.3 | 23.9 | 24.1 | 24.5 |
| **Percentages** |    |        |        |        |        |        |        |
| Parous | 69.9 | 62.5 | 70.8 | 56.7 | 70.2 | 64.8 | 68.0 | 67.3 |
| Premenopausal | 99.0 | 98.6 | 99.0 | 98.3 | 98.7 | 98.7 | 98.7 | 98.8 |
| Family history of ovarian cancerd | 1.5 | 1.9 | 1.5 | 1.8 | 1.5 | 1.6 | 1.8 | 1.5 |
| History of tubal ligation | 16.5 | 16.1 | 15.1 | 14.9 | 17.3 | 13.6 | 17.1 | 14.5 |

### Notes

- All factors except age were age-standardised in 5-year intervals for each cohort.
- Average childhood body fatness was calculated by taking the average of each participant's body fatness at ages 5 and 10 years, using a nine-level figure drawing.
- Baseline for each analysis was the year when the body size measure was assessed (see Table 1).
- Mother or sister had ovarian cancer according to the participant's response on the questionnaire; family history was evaluated using data from 1992 for NHS and 1993 for NHSII because it was not available in previous cycles.

### Results

One major difference between the NHS and NHSII cohorts is the menopausal status of participants when childhood body size was assessed (NHS: 32% premenopausal in 1988, NHSII: 99% premenopausal in 1989). To explore whether this could explain the observed difference in the association of childhood body fatness with risk, we combined the data from both cohorts and stratified by menopausal status in the cycle before diagnosis (Table 4). There was some suggestion of a weak positive association between childhood body fatness and risk of ovarian cancer in premenopausal women (pooled RR for level $> 5$ vs level $1 = 1.38$, 95% CI: 0.70 – 2.71, $P$ for trend = 0.92) and a weak inverse association in postmenopausal women (comparable RR = 0.85, 95% CI: 0.54 – 1.31, $P$ for trend = 0.09), although neither these nor the interaction with menopausal status ($P = 0.37$) were statistically significant. Results for BMI at the age of 18 years were similar in premenopausal women, but there was no evidence of an inverse association in postmenopausal women ($P$ for interaction = 0.11).

Alternatively, the observed variation in the associations for childhood body fatness could be explained by age differences between participants in the two cohorts (NHS: mean age = 54.3 in 1988, NHSII: mean age = 34.3 in 1989); therefore, we conducted a preliminary analysis combining both cohorts and stratifying by age, while adjusting for menopausal status. There were nonsignificant positive associations between childhood body fatness and ovarian cancer risk in women less than the age of 50 years (pooled RR for level $> 5$ vs level $1 = 1.77$, 95% CI: 0.85 – 3.69, $P$ for trend = 0.39) and in women between the ages 50 and 59 years (comparable RR = 1.30, 95% CI: 0.64 – 2.65, $P$ for trend = 0.34), whereas the association was inverse in women at the age of 60 years and older (comparable RR = 0.67, 95% CI: 0.39 – 1.16, $P$ for trend = 0.01). The interaction between childhood body fatness and age was statistically significant ($P = 0.001$). When we jointly stratified by age and menopausal status, the positive association between childhood body fatness and ovarian cancer risk appeared stronger in premenopausal women under the age of 45 years (pooled RR for childhood body fatness level $> 5$ vs level $1 = 2.51$, 95% CI: 0.94 – 6.73) than in those at the age of 45 years and older (comparable RR = 0.83, 95% CI: 0.30 – 2.28). Conversely, the inverse association in postmenopausal women was stronger among those at the age of 60 years and older (pooled RR for...
childhood body fatness level $\geq 5$ vs level 1 = 0.67, 95% CI: 0.39–1.16) than in those younger than the age of 60 years (comparable RR = 1.63, 95% CI: 0.73–3.66).

Additional adjustment for current BMI as a continuous variable had virtually no impact on the childhood body fatness associations in the NHS or for postmenopausal women overall (data not shown). However, the positive association for childhood body fatness in the NHSII and in premenopausal women was attenuated when including current BMI (RR for childhood body fatness level $\geq 5$ vs level 1 in premenopausal women = 1.23, 95% CI: 0.61–2.47, $P$ for trend = 0.73). Adjustment for age at menarche had no substantial impact on the associations (data not shown).

Height was positively associated with ovarian cancer risk in both cohorts (pooled RR for $\geq 1.75$ vs $<1.65$ m = 1.43, 95% CI: 1.05–1.96, $P$ for trend = 0.001), and the test for heterogeneity was not significant ($P = 0.22$); however, the association appeared stronger in the NHSII (comparable RR = 2.35, 95% CI: 1.19–4.63, $P$ for trend = 0.01) than in the NHS (comparable RR = 1.27, 95% CI: 0.88–1.82, $P$ for trend = 0.01) (Table 5). The positive association was slightly stronger among premenopausal than postmenopausal women, although there were no significant interactions with menopausal status or age (data not shown). Birthweight was not significantly associated with risk of ovarian cancer in either cohort.

The observed associations for body size in early life were similar for invasive cases alone and by histological type, although these analyses were limited by small case numbers. No significant interactions were observed between any of the body size measures and parity, oral contraceptive use, postmenopausal hormone use, or family history of breast or ovarian cancer (data not shown).

**DISCUSSION**

Results from this study indicate that body size in early life may be related to the risk of epithelial ovarian cancer. Body fatness during childhood was associated with ovarian cancer risk, although the association differed by cohort; greater body fatness at ages 5 and 10 years was associated with a slightly lower risk among NHS women, but a suggestion of an increased risk among NHSII women. These differences could be explained by differences in the menopausal status or age of participants. Body mass index at the age of 18 years and birthweight were not associated with risk; however, height was positively associated with risk.

To our knowledge, this is the first study to investigate the association of childhood fatness with ovarian cancer risk. Previous studies have examined the relation between overweight and obesity...
in adulthood and ovarian cancer risk or mortality, with mixed results (Olsen et al., 2007). One possible reason for these inconsistencies pertains to the timing of body size assessment. Several studies have observed stronger associations for BMI in adolescence or young adulthood than for recent BMI (Engeland et al., 2003; Anderson et al., 2004). The results from our study indicate that body size at even younger ages may be an important predictor of ovarian cancer risk.

A second potential explanation is that the relation between body size at young ages and ovarian cancer risk may differ by menopausal status or age. In a pooled analysis of 12 cohorts, BMI at baseline was not associated with risk overall or among postmenopausal women, but there was a positive association in premenopausal women (Schouten et al., 2008). Our findings also suggest that the positive association for childhood body fatness may be limited to premenopausal women, particularly younger premenopausal women.

### Table 4

| Premenopausal (NHS and NHSII) | Postmenopausal (NHS and NHSII) |
|-------------------------------|--------------------------------|
| **Cases** | **Person-years** | **Multivariatea RR (95% CI)** | **Cases** | **Person-years** | **Multivariatea RR (95% CI)** |
| Average body fatness at the ages of 5 and 10 yearsb | | | 123 | 250 962 | 1.0 (REF) |
| 1 | 25 | 239 481 | 1.0 (REF) | 96 | 232 530 | 0.89 (0.68–1.17) |
| 1.5–2 | 44 | 422 079 | 1.16 (0.71–1.90) | 59 | 167 233 | 0.79 (0.58–1.08) |
| 2.5–3 | 32 | 345 738 | 0.96 (0.57–1.64) | 38 | 142 234 | 0.60 (0.42–0.87) |
| 3.5–4.5 | 31 | 281 222 | 1.12 (0.65–1.90) | 24 | 60 158 | 0.85 (0.54–1.31) |
| ≥5 | 13 | 85 055 | 1.38 (0.70–2.71) | 0.92 | | |
| **P for trendd** | 0.09 | | | | | |

### Table 5

| Nurses’ Health Study (NHS) | Nurses’ Health Study II (NHSII) | NHS and NHSII combined |
|----------------------------|--------------------------------|-------------------------|
| **Cases** | **Person-years** | **Multivariatea RR (95% CI)** | **Cases** | **Person-years** | **Multivariatea RR (95% CI)** | **Cases** | **Person-years** | **Multivariatea RR (95% CI)** |
| Height, m | | | | | | | |
| <1.6 | 148 | 572 360 | 1.0 (REF) | 18 | 314 588 | 1.0 (REF) | 166 | 886 961 | 1.0 (REF) |
| 1.6 to <1.65 | 217 | 734 892 | 1.16 (0.94–1.43) | 36 | 448 775 | 1.41 (0.80–2.49) | 253 | 1 183 667 | 1.19 (0.98–1.45) |
| 1.65 to <1.7 | 198 | 710 542 | 1.11 (0.89–1.37) | 35 | 452 681 | 1.39 (0.78–2.47) | 233 | 1 163 223 | 1.14 (0.93–1.39) |
| ≥1.75 | 37 | 116 768 | 1.27 (0.88–1.82) | 16 | 129 593 | 2.35 (1.19–4.63) | 53 | 246 361 | 1.43 (1.05–1.96) |
| **P for trendd** | 0.01 | | | | | | |
| Birthweight, lbs | | | | | | | |
| <5.5 | 13 | 34 925 | 0.99 (0.55–1.78) | 2 | 36 519 | 0.68 (0.16–2.84) | 15 | 71 444 | 0.93 (0.54–1.60) |
| 5.5–6.9 | 69 | 165 868 | 1.08 (0.79–1.48) | 29 | 303 063 | 1.19 (0.74–1.92) | 98 | 468 931 | 1.11 (0.85–1.44) |
| 7.0–8.4 | 101 | 248 159 | 1.0 (REF) | 45 | 527 664 | 1.0 (REF) | 146 | 775 823 | 1.0 (REF) |
| ≥8.5 | 43 | 72 412 | 1.32 (0.92–1.90) | 12 | 149 557 | 0.83 (0.44–1.58) | 55 | 221 969 | 1.17 (0.85–1.60) |
| **P for trendd** | 0.58 | | | | | | |

### Notes

- a Multivariate analyses adjusted for age (continuous), parity (continuous), duration of oral contraceptive use (continuous), tubal ligation history (yes/no), and body mass index at the age of 18 years (<1.6, 1.6 to <1.65, 1.65 to <1.7, 1.7 to <1.75, and ≥1.75 m). b Average childhood body fatness calculated by taking the average of each participant’s body fatness at the ages 5 and 10 years. c Multivariate analyses for body fatness modelled as continuous variable. d Multivariate analyses for height adjusted for same factors except height (continuous) and BMI at the age of 18 years modelled as medians of categories. e Multivariate analyses for height adjusted for same factors except height (continuous), parity (continuous), duration of oral contraceptive use (continuous), tubal ligation history (yes/no), and body mass index at the age of 18 years (continuous). f Multivariate analyses for birthweight adjusted for same factors except height (continuous), parity (continuous), duration of oral contraceptive use (continuous), tubal ligation history (yes/no), and body mass index at the age of 18 years (continuous). g P value from multivariate model with birthweight modelled as continuous variable. h P value from multivariate model with BMI at the age of 18 years modelled as medians of categories. i P value from multivariate model with BMI at the age of 18 years modelled as medians of categories. j P value from multivariate model with BMI at the age of 18 years modelled as medians of categories. k P value from multivariate model with BMI at the age of 18 years modelled as medians of categories. l P value from multivariate model with BMI at the age of 18 years modelled as medians of categories. m P value from multivariate model with BMI at the age of 18 years modelled as medians of categories. n P value from multivariate model with BMI at the age of 18 years modelled as medians of categories. o P value from multivariate model with BMI at the age of 18 years modelled as medians of categories. p P value from multivariate model with BMI at the age of 18 years modelled as medians of categories. q P value from multivariate model with BMI at the age of 18 years modelled as medians of categories. r P value from multivariate model with BMI at the age of 18 years modelled as medians of categories. s P value from multivariate model with BMI at the age of 18 years modelled as medians of categories. t P value from multivariate model with BMI at the age of 18 years modelled as medians of categories. u P value from multivariate model with BMI at the age of 18 years modelled as medians of categories. v P value from multivariate model with BMI at the age of 18 years modelled as medians of categories. w P value from multivariate model with BMI at the age of 18 years modelled as medians of categories. x P value from multivariate model with BMI at the age of 18 years modelled as medians of categories. y P value from multivariate model with BMI at the age of 18 years modelled as medians of categories. z P value from multivariate model with BMI at the age of 18 years modelled as medians of categories.
that the observed associations for childhood body fatness were modest and the trends were only marginally significant. However, strong associations between childhood body fatness and risk of breast cancer have been observed in previous studies, suggesting that it may predict the risk of hormone-related cancers in women. Further, previous studies have observed significant associations of both adolescent and adult BMI with ovarian cancer risk, lending plausibility to our findings.

The associations of body size in childhood with ovarian cancer risk could be mediated through endogenous hormones (Risch, 1998; Lukanova and Kaaks, 2005), although epidemiological data on the relation of endogenous hormones with ovarian cancer risk are sparse and inconsistent (Eliassen and Hankinson, 2008). Obesity during adolescence has been associated with polycystic ovary syndrome (PCOS), which is characterised by elevated levels of luteinising hormone (LH) and possibly increased risk of ovarian cancer (Schouten et al, 1996). Paradoxically, although, adiposity in pre-menopausal women without PCOS (Cramer et al, 2002) and postmenopausal women (Malacara et al, 2001) has been related to lower LH levels. Obesity in pre-adolescent and adolescent girls is associated with hyperinsulinemia and increased production of androgens (Stoll, 1998), which may be related to ovarian cancer risk (Risch, 1998). Also, lower levels of progesterone could increase risk (Risch, 1998). Obesity in adolescence is associated with increased risk of ovulatory infertility in adulthood (Stoll, 1998), which leads to decreased progesterone levels, and infertility has been associated with a modest increase in ovarian cancer risk (Tworoger et al, 2007a). Clearly, more research is needed to clarify the biological mechanisms underlying these associations.

The observed positive association between height and ovarian cancer risk is consistent with results from previous studies in several populations (Rodriguez et al, 2002; Engeland et al, 2003; Schouten et al, 2003, 2008). Interestingly, the positive association for height has been restricted to or stronger among younger or premenopausal women (Kuper et al, 2002; Lukanova et al, 2002; Engeland et al, 2003; Schouten et al, 2008), consistent with our findings. It is also possible that the difference in the observed association for height in the NHS and the NHSII could be due to a cohort effect.

Despite consistency across epidemiological studies, the biological mechanisms explaining the observed associations for height are unclear. Adult height may be a marker for genetic factors or of energy intake, caloric restriction, or exposure to sex and growth hormones in early life (Schouten et al, 2003). Height consistently

has been associated with risk of other cancers (Gunnell et al, 2001), and the growth hormone/insulin-like growth factor (IGF) axis is a potential pathway (Gunnell, 2000). However, epidemiological studies of circulating levels of IGF and ovarian cancer risk are inconclusive (Tworoger et al, 2007b), and thus other mechanisms may be involved.

There are almost no epidemiological data on the relation of birthweight to risk of ovarian cancer. In a small retrospective medical record review, birthweight was not associated with mortality from ovarian cancer, but greater weight gain in the first year of life was associated with increased mortality (Barker et al, 1995). The investigators hypothesised that patterns of gonadotropin release are established in utero and during infancy, and that this could influence ovarian cancer pathogenesis later in life. A population-based case–control study found no overall association between birthweight and ovarian cancer risk, although among women younger than the age of 55 years, there was a decreased risk of ovarian cancer for those who weighed <5.5 pounds at birth compared to those who weighed 5.5–9 pounds (Rossing et al, 2008). Our findings do not support an association of birthweight with ovarian cancer risk, although this should be confirmed in other populations.

Our study has several limitations. Although we combined data from two large cohort studies, we had limited power to examine interactions or variation by histological type. Another limitation is the reliance on recall of body size in early life. Important strengths of our study include its large sample size, its prospective design, the confirmation of ovarian cancer cases, and the detailed information on menopausal status and ovarian cancer risk factors.

In summary, this study suggests that body fatness during childhood and adult height may be related to ovarian cancer risk, and that these associations may vary by menopausal status or age. This is the first study to examine the associations of body fatness during childhood with ovarian cancer risk, and it is one of the largest, most comprehensive studies of the other body size measures to date. Further research should confirm these findings in other populations as well as examine the underlying biological mechanisms.

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