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Keywords: coffee; endometrial cancer; decaffeinated coffee; caffeine; tea; methylxanthines; cola

Intake of coffee, caffeine and other methylxanthines and risk of Type I vs Type II endometrial cancer

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Background: Coffee and other sources of methylxanthines and risk of Type I vs Type II endometrial cancer (EC) have not been evaluated previously.

Methods: Prospective cohort of 23356 postmenopausal women with 471 Type I and 71 Type II EC cases.

Results: Type I EC was statistically significantly associated with caffeinated (relative risk (RR) = 0.65 for 4 + cups per day vs \leq 1 cup per month: 95% confidence interval (CI): 0.47–0.89) but not decaffeinated (RR = 0.76; 95% CI: 0.50–1.15) coffee intake; there were no associations with tea, cola or chocolate, or for Type II EC. The inverse association with caffeinated coffee intake was specific to women with a body mass index 30 + kg m⁻² (RR = 0.56; 95% CI: 0.36–0.89).

Conclusion: Coffee may protect against Type I EC in obese postmenopausal women.

Following water and tea, coffee is the third most consumed beverage in the world (Bushman, 1998; La Vecchia and Tavani, 2007). A recent meta-analysis reported an inverse association of coffee intake with endometrial cancer (EC) risk (Je and Giovannucci, 2011). The presence of antioxidants and other chemopreventive compounds in coffee may explain its anticarcinogenic effect (Vivani, 1993; Cavin et al, 2002). However, it is not clear whether coffee per se, caffeine or other methylxanthines (e.g., theophylline and theobromine) are most relevant. Also unexplored is whether there is heterogeneity by Type I vs Type II EC, which may have different aetiologies (Bokhman, 1983; Doll et al, 2008; Mendivil et al, 2009). The aim of the present study was to evaluate the association of coffee consumption (with and without caffeine) and other sources of methylxanthines with risk of Type I vs Type II EC, overall and stratified on body mass index (BMI), smoking history and hormone therapy (HT) use.

MATERIALS AND METHODS

Details regarding the Iowa Women's Health Study (IWHS) have been published (Folsom *et al*, 1990). In brief, 41 836 women aged 55–69 years completed a self-administered survey at enrolment in 1986. The baseline survey included a 126-item semiquantitative food-frequency questionnaire (FFQ) (Willett *et al*, 1988), which included the average intake in the past year of the following items: caffeinated coffee; decaffeinated coffee; tea (excluding herbal teas); regular and sugar-free carbonated beverages with caffeine; chocolate; chocolate bars; and brownies. The FFQ was reliable and valid in this population (Munger *et al*, 1992).

Incident EC cases were identified through 2005 via annual linkage with the Iowa Cancer Registry. Cancer data were coded according to the International Classification of Diseases for Oncology (Fritz *et al*, 2000). Type I or Type II were classified

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Intake ^b	Person-years	Cases	Age-and energy- adjusted RR	P-trend	Multivariable- adjusted RR (95% CI) ^c	P-trend	Cases	Age- and energy- adjusted RR	P-trend	Multivariable- adjusted RR (95% Cl) ^c	P-trend	P -heterogeneity
Total coffee intake (cups)	-				-						_	
Never or ≼once per month <1 cup per week 1 cup per day 2-3 cups per day 4 + cups per day	37 203 41 565 48 627 136 795 109 730	64 64 55 188 100	1.00 0.89 0.65 0.80 0.54	0.00021	1.00 (reference) 0.95 (0.66, 1.36) 0.75 (0.52, 1.09) 0.95 (0.71, 1.28) 0.71 (0.51, 0.99)	0.11	7 8 13 26 17	1.00 0.98 1.39 1.02 0.87	0.63	1.00 (reference) 0.98 (0.36, 2.72) 1.31 (0.51, 3.35) 1.01 (0.43, 2.36) 0.84 (0.33, 2.12)	0.64	06.0
Caffeinated coffee (cups)												
Never or ≼once per month <1 cup per week 1 cup per day 2-3 cups per day 4 + cups per day	106 367 68 439 35 148 91 456 72 510	168 86 37 121 59	1.00 0.80 0.67 0.84 0.52	0.00023	1.00 (reference) 0.83 (0.63, 1.08) 0.70 (0.48, 1.02) 0.92 (0.72, 1.18) 0.65 (0.47, 0.89)	0.033	18 7 16 11	1.00 1.66 1.19 1.07 0.97	0.66	1.00 (reference) 1.56 (0.81, 3.01) 1.08 (0.43, 2.74) 1.08 (0.55, 2.13) 0.85 (0.37, 1.93)	0.58	62.0
Decaffeinated coffee (cups)					-		-					
Never or ≼once per month <1 cup per week 1 cup per day 2-3 cups per day 4 + cups per day	163 125 80 239 36 994 63 125 30 436	224 90 42 87 28	1.00 0.81 0.82 1.00 0.68	0.19	1.00 (reference) 0.85 (0.66, 1.09) 0.82 (0.58, 1.15) 1.06 (0.82, 1.36) 0.76 (0.50, 1.15)	0.53	31 16 12 5	1.00 1.04 0.97 0.89 0.89	0.86	1.00 (reference) 1.15 (0.62, 2.14) 1.05 (0.46, 2.41) 1.01 (0.50, 2.04) 1.08 (0.41, 2.80)	0.93	0.76
Joint intake of caffeinated and decaff	feinated coffee	e intake	(cups)		-		-					
Never or ≼once per month Decaffeinated only, 1–3 cups per day Decaffeinated only, 4 + cups per day Caffeinated only, 1–3 cups per day Caffeinated only, 4 + cups per day Caffeinated 1 + cups per day decaffeinated 1 + cups per day	83 259 67 797 23 749 23 749 95 043 65 062 39 009	132 97 25 125 57 35	1.00 0.90 0.67 0.56 0.56	N/A	1.00 (reference) 1.02 (0.78, 1.33) 0.81 (0.52, 1.27) 0.95 (0.73, 1.22) 0.73 (0.52, 1.02) 0.69 (0.47, 1.01)	A/A						
Tea (cups), not herbal												
Never or <once month<br="" per="">1–3 cups a month 1–4 cups a week 5+ cups a week</once>	161 860 62 881 75 012 74 166	207 70 95 99	1.00 0.87 0.99 1.05	0.72	1.00 (reference) 0.87 (0.66, 1.15) 0.89 (0.69, 1.15) 0.95 (0.74, 1.22)	0.55	27 13 15 16	1.00 1.26 1.35	0.32	1.00 (reference) 1.28 (0.65, 2.52) 1.23 (0.65, 2.34) 1.26 (0.65, 2.43)	0.46	0.36
Cola, regular or low calorie (glass, bo	ttle or can)											
Never or <once month<br="" per="">1–3 cups a month 1 + per week</once>	234 311 57 412 82 196	295 66 110	1.00 0.93 1.10	0.52	1.00 (reference) 0.99 (0.75, 1.30) 1.08 (0.86. 1.36)	0.55	39 15 17	1.00 1.68 1.41	0.15	1.00 (reference) 1.65 (0.89, 3.07) 1.42 (0.79, 2.56)	0.16	0.28

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				Γype I (N =	= 471)				Typ	e II (N=71)		
Chocolate (bars or pieces)												
Never or <once month<br="" per="">1-3 bars per pieces a month 1+ per week</once>	171 617 122 065 80 237	232 143 96	1.00 0.87 0.90	0.28	1.00 (reference) 0.87 (0.70, 1.09) 0.94 (0.73, 1.21)	0.47	30 20 21	1.00 1.01 1.80	0.071	1.00 (reference) 1.00 (0.55, 1.80) 1.79 (0.98, 3.26)	0.085	0.062
Candy bars												
Never or <once month<br="" per="">1–3 bars a month 1+ per week</once>	208240 113664 52015	269 141 61	1.00 0. <i>97</i> 0.93	0.60	1.00 (reference) 0.98 (0.79, 1.21) 0.96 (0.71, 1.29)	0.76	33 26 12	1.00 1.58 1.80	0.044	1.00 (reference) 1.46 (0.85, 2.50) 1.71 (0.84, 3.48)	0.087	0.090
Brownies (one)												
Never or <once month<br="" per="">1-3 servings a month 1+ per week</once>	229481 110854 33584	297 135 39	1.00 0.95 0.92	0.52	1.00 (reference) 1.02 (0.82, 1.26) 0.98 (0.68, 1.40)	1.00	43 23 5	1.00 1.21 0.97	0.71	1.00 (reference) 1.12 (0.65, 1.92) 1.00 (0.38, 2.58)	0.82	0.83
Caffeine (mg per day)										-		
<29.7 29.7–158.3 158.4–385.0 > 385.0	92717 93302 93896 94004	138 132 107 94	1.00 0.95 0.77 0.68	0.0015	1.00 (reference) 0.93 (0.72, 1.18) 0.80 (0.61, 1.04) 0.80 (0.61, 1.05)	0.059	13 22 23 13	1.00 1.74 1.84 1.09	0.76	1.00 (reference) 1.65 (0.82, 3.29) 1.80 (0.90, 3.59) 0.98 (0.43, 2.23)	0.84	0.38
Abbreviations: CI = confidence interval; HT = hormone ^a Type I defined as ICD-O codes 8000, 8010, 8041, 814. ^b Frequency of use ('never or less than once per month', (3) tea (1 cup), not herbal teas; (4) Coke, Pepsi or other o (2) tea (1 cup), not herbal teas; (9) condy bars, for example chaited for age. diabetes. duration of HT use. hvoer	therapy; ICD = Int 0, 8210, 8262, 826; 71–3 per month', cola with sugar; (5) le, Snickers, Milky trension, ace at m	:ernational C 3, 8380, 8480 1 per week', caffeine-free Way, Reeses	lassification of I , 8560 and 8570 , 2–4 per week', Coke, Pepsi or); and (10) brow	Diseases; RR = , and Type II of 5-6 per week', other cola with mies (1). Total	relative risk. Jefined as ICD-O code: , '1 per day', '2-3 per da 1 sugar, (6) low calorie co coffee is the sum of ca odor mass index, wanist.	s 8050, 8260, y', 4–5 per da ola, for exam ffeinated plu to-hip ratio. s	8310, 8323, . y/, '6 + per d ole, Tab with s decaffeinat	8441, 8460, 8950, 8951 a lay) was asked for the fol r caffeine; (7) low calorie v ted coffee intake.	nd 8980. Iowing items: caffeine-free o a. total enero	(1) caffeinated coffee (1 c cola, for example, Pepsi fi xv and alcohol use.	up); (2) decaf ee; (8) chocol	einated coffee (1 cup); ate (bars or pieces), for

based on registry codes (see Table 1 footnote) as described previously (Uccella *et al*, 2011); there was no central pathology review. Deaths were ascertained by follow-up surveys, annual linkage with Iowa death certificates and linkage to the National Death Index.

Women with history of cancer before baseline, except nonmelanoma skin cancer (n = 3830); hysterectomy before baseline (n = 14350); extreme dietary intake (<600 or > 5000 kcal per day) or incomplete FFQ questionnaires (≥ 30 blank food items) (n = 3096); or who were not postmenopausal at baseline (n = 569) were excluded from the present analysis (not mutually exclusive), yielding a final sample size of 23356 study participants.

Each woman accumulated person-years of follow-up from baseline to date of EC diagnosis, move from Iowa, death or administrative censoring on 31 December 2005. Relative risks (RR) and 95% confidence intervals (95% CIs) were estimated using Cox proportional hazards regression, and modelling age was used as the time variable (Korn *et al*, 1997). All Cox model attributes included as covariates are listed in corresponding table footnotes, and were selected *a priori* based on their suspected or known associations with endometrial cancer. Separate analyses were carried out for Type I and Type II EC. Tests for trend were carried out by ordering the intake quartiles from lowest to highest and including the resulting variable as a 1 d.f. linear term in the Cox regression models.

We formally determined if risk ratios for the exposure variables differed by type of EC using a competing risk form of Cox proportional hazards regression (Lunn and McNeil, 1995). We also examined associations between exposure variables and subtypespecific EC risk within strata defined by BMI, smoking status and use of HT. All statistical tests were two-sided, and analyses were carried out using SAS (SAS Institute Inc., Cary, NC, USA) and R software systems.

RESULTS

At study baseline, there were 23 356 women in the at-risk cohort, of whom 5218 (22.3%) were obese (BMI \ge 30 kg m⁻²) and

6843 (29.3%) drank 4 + cups per day of coffee (caffeinated or decaffeinated). The correlation of coffee intake with EC risk factors is shown in Table 2.

During the 20-year follow-up period, we identified a total of 542 incident cases of EC, 471 Type I and 71 Type II. The mean age at diagnosis of Type I EC was 71.8 years (range, 57.2–89.5 years) and Type II EC was 72.8 years (range, 60.2–89.3 years).

There was an inverse association of caffeinated coffee consumption with risk of Type I EC after multivariate adjustment (RR = 0.65 for 4 + cups per day compared with ≤ 1 cup per month; *P*-trend = 0.033), but there were no statistically significant trends with intake of total coffee, decaffeinated coffee, tea, colas or other sources of methylxanthines, although the highest intake of total coffee and decaffeinated coffee did have RRs < 0.8 (Table 1). Compared with women who did not drink either caffeinated or decaffeinated coffee, those who drank 4+ cups per day of caffeinated coffee only (RR = 0.73; 95% CI: 0.52-1.02) or 1 + cups per day of both types of coffee (RR = 0.69; 95% CI: 0.47–1.01) had lower EC risk, whereas the association was weaker and not statistically significant for women who drank 4+ cups per day of decaffeinated coffee only (RR=0.81; 95% CI: 0.52-1.27). Caffeine intake showed a suggestive inverse associated with risk (RR = 0.80 for > 385 mg per day compared with < 29.7 mg per day; P-trend = 0.059). In contrast, coffee and other sources of methylxanthines were not associated with risk of Type II EC.

We next examined coffee intake with risk of Type I EC within strata defined by BMI ($30 + vs < 30 \text{ kg m}^{-2}$), smoking history (ever/never) and HT use (ever/never); the sample size was too small to conduct these analyses for risk of Type II EC. As shown in Table 3, the inverse associations for total and caffeinated coffee, caffeine and perhaps decaffeinated coffee were only observed among obese women and not among women with a BMI < 30 kg m^{-2} . There was no striking or consistent heterogeneity in the associations for coffee or caffeine intake when stratified on smoking status (Supplementary Table 1) or HT use (Supplementary Table 2).

Table 2. Correlation of coffee intake with selected	endometrial cancer risk factors	s, Iowa Women's Health	Study (1986)
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		Ir	take of coffee		
Variable	Never or ≼1 per month (N=2340)	<1 cup per week (N =2638)	1 cup per day (N = 3040)	2–3 cups per day (N =8495)	4+ cups per day (N =6843)
Mean \pm s.d.					
Age (years) Body mass index (kg m ⁻²) Waist-to-hip ratio Total energy (kcal per day) Pack years of smoking	62.1 ± 4.2 27.6 ± 5.6 0.843 ± 0.086 1785 ± 613 5.7 ± 15.0	62.7 ± 4.2 27.3 ± 5.5 0.837 ± 0.082 1718 ± 600 5.9 ± 14.4	62.9 ± 4.2 27.0 ± 5.0 0.838 ± 0.083 1785 ± 602 5.6 ± 14.0	62.3 ± 4.2 26.8 ± 4.9 0.832 ± 0.081 1804 ± 584 8.3 ± 16.4	61.4 ± 4.1 26.5 ± 5.0 0.828 ± 0.086 1871 ± 648 15.7 ± 21.4
Percent distribution					
Adult-onset diabetes (ever) Hypertension (ever) Any alcohol use Age at menarche >12 years Age at menopause >50 years Never used HT	7.1% 36.7% 22.9% 55.8% 63.2% 74.4%	6.9% 38.5% 35.9% 58.9% 62.0% 73.9%	6.3% 38.3% 39.6% 60.7% 64.3% 73.1%	5.3% 34.9% 50.0% 59.1% 64.1% 73.1%	4.8% 30.1% 54.9% 57.9% 59.7% 73.3%
Smoking history					
Current Former Never	7.6% 12.8% 79.6%	8.6% 15.8% 75.6%	8.0% 15.8% 76.3%	12.7% 20.7% 66.6%	27.7% 22.8% 49.4%

Abbreviation: HT = hormone therapy.

Table 3. Association of coffee and caffeine with risk of Type I endometrial cancer, stratified by BMI, Iowa Women's Health Study, 1986–2005

		BMI <	30 kg m ^{- 2}	l		BMI 30	+ kg m ⁻²		
	1	,							T
			Multivariable-				Multivariable-		
			adjusted RR ^a			-	adjusted RR ^a		
Level of intake	Person-years	Cases	(95% CI)	P-trend	Person-years	Cases	(95% CI)	P-trend	P -interaction
Total coffee intake									
Never or \leqslant once per month	27 242	25	1.00 (reference)	0.75	9961	39	1.00 (reference)	0.010	0.054
<1 cup per week	31 426	28	1.09 (0.62, 1.94)		10 140	36	0.82 (0.52, 1.31)		
1 cup per day	37 451	29	1.00 (0.57, 1.77)		11 176	26	0.60 (0.36, 0.99)		
2–3 cups per day	108 294	110	1.33 (0.83, 2.14)		28 501	78	0.72 (0.49, 1.07)		
4+ cups per day	88 005	59	0.99 (0.59, 1.66)		21 725	41	0.53 (0.34, 0.84)		
Caffeinated coffee						<u> </u>			
Never or ≼once per month	80 699	74	1.00 (reference)	0.80	25 668	94	1.00 (reference)	0.0079	0.063
<1 cup per week	53 029	46	1.02 (0.70, 1.49)		15 410	40	0.66 (0.45, 0.97)		
1 cup per day	27 728	23	0.97 (0.59, 1.59)		7420	14	0.51 (0.28, 0.91)		
2–3 cups per day	72716	75	1.21 (0.86, 1.69)		18740	46	0.71 (0.50, 1.02)		
4+ cups per day	58 245	33	0.77 (0.50, 1.19)		14 264	26	0.56 (0.36, 0.89)		
Decaffeinated coffee									
Never or ≼once per month	125 409	110	1.00 (reference)	0.95	37 716	114	1.00 (reference)	0.32	0.58
<1 cup per week	63 908	54	0.94 (0.67, 1.32)		16 331	36	0.73 (0.50, 1.08)		
1 cup per day	28 922	24	0.93 (0.59, 1.46)		8072	18	0.71 (0.43, 1.19)		
2–3 cups per day	49781	45	1.06 (0.74, 1.51)		13 344	42	1.05 (0.73, 1.50)		
4+ cups per day	24 397	18	0.90 (0.53, 1.53)		6039	10	0.58 (0.30, 1.11)		
Caffeine (mg per day)									
<29.7	71 320	63	1.00 (reference)	0.66	21 397	75	1.00 (reference)	0.038	0.19
29.7–158.3	71 693	71	1.11 (0.78, 1.58)		21 609	61	0.80 (0.56, 1.12)		
158.4–385.0	74716	64	1.00 (0.69, 1.44)		19179	43	0.67 (0.46, 0.99)		
>385.0	74 687	53	0.94 (0.64, 1.38)		19317	41	0.70 (0.47, 1.04)		

Abbreviations: $\mathsf{BMI}\,{=}\,\mathsf{body}$ mass index; $\mathsf{HT}\,{=}\,\mathsf{hormone}$ therapy.

^aAdjusted for age, duration of HT use, diabetes, hypertension, age at menarche, age at menopause, BMI (continuous), waist-to-hip ratio, smoking status, pack years of smoking, total energy and alcohol use.

DISCUSSION

Coffee consumption was most strongly associated with a lower risk of Type I EC among obese postmenopausal women, and these associations were generally stronger and statistically significant for caffeinated relative to decaffeinated coffee intake. There were no statistically significant associations of coffee consumption with Type I EC among non-obese women or for Type II EC. Tea, cola and chocolate intake were not associated with risk of Type I or Type II EC.

A recently updated meta-analysis of 6 cohort and 10 casecontrol studies (Je and Giovannucci, 2011) reported a pooled RR of 0.71 (95% CI: 0.62–0.81) for the risk of EC for the highest *vs* lowest categories of coffee intake, with the strongest inverse association observed in Japanese studies (RR = 0.40; 95% CI: 0.25–0.63), intermediate for North American studies (RR = 0.69; 95% CI: 0.60–0.79) and weakest but still evident for European studies (RR = 0.79; 95% CI: 0.63–0.99). Consistent with our results, four recent studies found an inverse association of coffee with EC, particularly among women with BMI \geq 30 kg m⁻² (Friberg *et al*, 2009; Giri *et al*, 2011; Gunter *et al*, 2011; Je *et al*, 2011). For the first time, we extend this association specifically to Type I EC and to coffee but not other common sources of methylxanthines, which were not addressed by these prior studies.

The exact mechanisms involved in any putative beneficial effect of coffee on EC remain largely unknown. Coffee is a major source of caffeine, and this methylxanthine may increase levels of circulating sex-hormone-binding globulin, thus reducing the concentrations of bioavailable sex-steroid hormones, in particular free oestradiol, and consequently modifying the hormonal milieu leading to downregulation of endometrial hyperproliferation (Ferrini and Barrett-Connor, 1996; Nagata *et al*, 1998). However, coffee, irrespective of caffeine content, also contains additional compounds with antioxidant activities. These compounds vary widely depending on the type of coffee, roasting and preparation, and many have been found to inhibit the proliferation of tumour cells *in vitro* (Vivani, 1993; Cavin *et al*, 2002).

An intriguing hypothesis suggests that coffee may be an insulin sensitiser (Wu *et al*, 2005; Huxley *et al*, 2009; Loopstra-Masters *et al*, 2011). Coffee (both caffeinated and decaffeinated) and caffeine intake were inversely associated with levels of circulating C-peptide, a marker of insulin secretion and resistance, and this association was much stronger in overweight and obese women (Wu *et al*, 2005).

An inverse association with coffee was not observed for Type II EC, although our analysis was limited by the relatively small number of Type II cases and by the absence of central pathology review. Type I and Type II EC may have different aetiologic pathways and distinct risk factors (Uccella *et al*, 2011). From a molecular point of view, Type II EC is often associated with p53 mutations, which commonly lead to DNA derangements, chromosomal instability and a more aggressive clinical behaviour (Doll *et al*, 2008). Conversely, alterations of p53 have been reported

in only a small proportion of Type I tumours and, when they occur, they are usually a late event (Doll *et al*, 2008). Apoptosis of rapidly growing cells induced by caffeine *in vitro* is dependent on the presence of a functional p53 product, so when p53 is mutated cellular growth is not inhibited by caffeine (He *et al*, 2003).

In conclusion, our results suggest that coffee consumption, perhaps in part related to caffeine, may be relevant for chemoprevention of Type I EC, particularly among obese women.

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