

Consumption of industrial processed foods and risk of premenopausal breast cancer among Latin American women: the PRECAMA study

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ABSTRACT

Ultra-processed food intake has been linked to an increased risk of breast cancer in Western populations. No data are available in the Latin American population although the consumption of ultra-processed foods is increasing rapidly in this region.

We evaluated the association of ultra-processed food intake to breast cancer risk in a case–control study including 525 cases (women aged 20–45 years) and 525 matched population-based controls from Chile, Colombia, Costa Rica and Mexico. The degree of processing of foods was classified according to the NOVA classification. Overall, the major contributors to ultra-processed food intake were ready-to-eat/heat foods (18.2%), cakes and desserts (16.7%), carbonated and industrial fruit juice beverages (16.7%), breakfast cereals (12.9%), sausages and reconstituted meat products (12.1%), industrial bread (6.1%), dairy products and derivatives (7.6%) and package savoury snacks (6.1%). Ultra-processed food intake was positively associated with the risk of breast cancer in adjusted models (OR_{T3-T1}=1.93; 95% CI=1.11 to 3.35). Specifically, a higher risk was observed with oestrogen receptor positive breast cancer (OR_{T3-T1}=2.44, (95% CI=1.01 to 5.90, P-trend=0.049), while no significant association was observed with oestrogen receptor negative breast cancer (OR_{T3-T1}=1.87, 95% CI=0.43 to 8.13, P-trend=0.36).

Our findings suggest that the consumption of ultra-processed foods might increase the risk of breast cancer in young women in Latin America. Further studies should confirm these findings and disentangle specific mechanisms relating ultra-processed food intake and carcinogenic processes in the breast.

INTRODUCTION

Risk factors for breast cancer in premenopausal women are not well understood.¹ This may be related to different tumour biology in young women compared with older women.² In Latin America close to 27% of

What this paper adds

- ⇒ This study reports for the first time that the consumption of ultraprocessed foods is associated with an increased risk of breast cancer in young women from Latin American countries.
- ⇒ Our findings, if confirmed, will be instrumental to support public policies.

breast cancer occurs between 20 and 45 years and increasing trends have been observed in the last decades with a larger proportion of oestrogen receptor negative (ER-) and triple negative (TN) tumours than observed in Western countries.¹ This population is currently undergoing a rapid lifestyle and nutritional transition switching from a varied traditional diet (including corn tortillas, corn flour cakes, beans and other legumes, soup, homemade stew, vegetable, whole fruit) towards a more homogenous diet rich in industrial ultra-processed foods.^{3,4}

Ultra-processed foods refer to formulations of substances that have been extracted or refined from whole foods through processes like baking, frying, extruding, moulding, reshaping, hydrogenation and hydrolysis. The final food or beverage product created is durable, convenient and palatable, ready-to-heat or ready-to-eat as snacks or to replace freshly prepared food-based dishes and meals.⁵ Ultra-processed foods include mostly carbonated soft drinks, sweet and savoury snacks, breakfast cereals, packaged bread, confectionary, cakes and pastries, ice cream, cookies, fruit and vegetables juices, energy drinks, pre-prepared meat, margarine and

spreads sauces and ready meals.⁶ A recent report from the WHO/Pan American Health Organisation⁷ pointed out the importance of this nutritional change, highlighting a 43% increase in the sales of ultra-processed foods and drinks in Latin America between 2000 and 2013 while the increase in North America during the same period was only 3%. The fastest increases were observed in Mexico and Chile, while in Costa Rica and Colombia the increase was lower. In 2013, the annual retail sales of ultra-processed foods and drink products in kilograms per capita were 307 in the USA (the world highest), 210 in Mexico (fourth), 202 in Chile (seventh), 120 in Costa Rica and 87 in Colombia.⁷

Classification of foods by degree of food processing could improve our understanding of how dietary factors may relate to chronic diseases.⁸ The NOVA classification categorises food items in accordance to their level of processing into minimally processed, processed culinary ingredients, processed foods and ultra-processed food and drink products.⁵ Ultra-processed foods are often energy dense, rich in saturated and trans-fatty acids, added sugar and salt and low content in fibres and vitamins and have been associated with an increased risk of several chronic diseases, in particular obesity, metabolic syndrome and dyslipidaemia^{6–11} as well as cancer.¹¹ Possible mechanisms for these associations may include industrial trans-fatty acids, formed when fats and oils are partly hydrogenated during industrial processing, and that have been linked to the risk of breast and ovarian cancer.¹⁰ To date there is no study on the potential role of ultra-processed food intake on the risk of cancer in Latin America.

Given the potential for preventive measures, there is an urgent need to investigate the impact of ultra-processed foods on the incidence of cancer, and principally breast cancer, the most common cancer among women in Latin America.⁹

In this manuscript, we report the association of ultra-processed food intake and risk of breast cancer, overall and by hormonal receptors status, within the framework of the PRECAMA project, an ongoing multicentric population-based case–control study coordinated by the International Agency for Research on Cancer (IARC).

METHODS

The PRECAMA study

The PRECAMA study involves scientific teams in four Latin American countries (Chile, Colombia, Costa Rica and Mexico). The methodology has previously been reported.¹² Through IARC and a central laboratory (the Porter Lab) at the Fred Hutchinson Cancer Research Center (FHCRC) in Seattle, the research teams have harmonised and standardised the protocol approved by at least one Institutional Review Board within each country and by the IARC Ethics Committee.

Selection of cases and controls

Women with incident primary invasive breast cancer cases were recruited from major cancer hospitals in four Latin American countries before any therapies.¹² The inclusion criteria were: age 20–45 years; being resident for ≥ 3 years in the same city district; having an incident primary invasive breast cancer with positive biopsy and clinical staging according to TNM standards and having menstruated at least once in the past 12 months. To date 525 primary invasive breast cancer incident cases have been recruited among premenopausal women (40 in Chile, 129 in Colombia, 43 in Costa Rica and 313 in Mexico).

Controls (one per case) were selected from the population residing in the same city district as the case for at least 3 years, applying the same exclusion criteria as for the cases, and matched to cases on age (± 3 years), area of residence and healthcare institution membership. Recruitment of cases and controls started in 2012.

Exposure assessment

During the recruitment visit, trained nurses administered health and lifestyle questionnaires as well as validated and standardised food frequency questionnaires to assess diet. Anthropometry was measured according to standardised protocols and following Lohman's recommendations.¹³ Biological samples (fasting blood and urine) were also collected for biomarker analyses.

Dietary intake assessment

Dietary intake was assessed through a standardised semi-quantitative food frequency questionnaire with country-specific food list and a standard portion size for each food item. These questionnaires have been validated in women residing in Mexico City^{14 15} and in Costa Rica.¹⁶ In each country, the list of food items was carefully reviewed with input from local nutritionists, and additional foods were added if necessary, according to local nutrition surveys, in order to capture the local diet. Thus, the number of food and drink items in each questionnaire varied across countries, and included 195, 217, 168 and 191 items, respectively, for Chile, Colombia, Costa Rica and Mexico.

For each food item, a commonly used unit or portion size was indicated (specified serving size: a slice, a glass or a natural unit such as one apple), and participants were asked how often, on average, over the previous year, they consumed the specified amount of each food, choosing among 10 frequencies of consumption: >6 times/day, 4–5 times/day, 2–3 times/day, 1 time/day, 5–6 times/week, 2–4 times/week, 1 time/week, 2–3 times/month, <1 time/month or never. In addition, information was collected on intake of alcoholic and non-alcoholic drinks (sugar-sweetened beverages, tea, coffee and water), as well as on added sugar, added salt, fat used for cooking and intake of fat from meat. Food items were linked to their macronutrient and micronutrient profile and total caloric intake was calculated based on context-specific food composition tables.

Categorisation of dietary intake NOVA classification

Dietary data from the semi-quantitative food frequency questionnaire were used to develop the NOVA classification based on the work by Monteiro *et al.*⁵ Foods were categorised into four groups according to their degree of processing as: (1) unprocessed or minimally processed foods or NOVA 1 (eg, fresh fruits and vegetables, unprocessed meat, fish...); (2) processed culinary ingredients or NOVA 2 (eg, oils, butter, sugar...); (3) processed foods or NOVA 3 (eg, canned fish, cheeses, fresh bread...); and (4) ultra-processed foods or NOVA 4 (eg, soft drinks, sweet and savoury packaged snacks, processed meat, ready-to-eat/heat foods (burgers, hot dogs, instant soups,....)....). The classification of the various food items in the different NOVA categories was reviewed by local nutritionists in each country (considering the degree of processing of each food item in their local context) and classified in collaboration with Dr Khandpur (University of Sao Paulo, Brazil). In case of doubt between two NOVA categories (3.8% of the foods), the food item was allocated to both groups while using a percentage in each group (eg, baked custard 30% NOVA 1% and 70% NOVA 4) based on the dietary habits provided by local nutritionists.

This study focuses primarily on identifying and classifying ultra-processed foods (NOVA 4). Ultra-processed foods were classified into 14 subgroups including carbonated beverages and industrial fruit juices, flavoured yoghurt and dairy products, distilled alcoholic beverages, industrial breads, breakfast cereals, confectionery, package sweet snacks, packaged savoury snacks, cakes and desserts, sausage and reconstituted meat products, ready-to-eat/heat foods, industrial cheese and cheese substitutes, margarine and butter substitutes and other ultra-processed foods (including sauces, gravies, jam, mayonnaise, artificial sweeteners).

Pathology review and immunohistochemistry analyses

Prepared histology sections from tumour biopsy obtained prior to any treatment were reviewed for diagnosis, tumour grade, lymphovascular invasion and stromal and lymphocyte response. Testing was conducted by immunohistochemistry (IHC) for ER (SP1, LabVision, Fremont California), PgR (PgR 636, DAKO, Denmark), human epidermal growth factor receptor 2 (AO485, DAKO, Denmark), according to standardised and optimised protocols that included antigen retrieval when required. All analyses were conducted centrally at the Porter Lab at the FHCRC. Each centre applies standardised protocols for specimen collection.¹²

Laboratory analysis of trans-fatty acids

Industrial trans-fatty acids blood concentration was measured in a subsample (n=284) in order to evaluate the correlation with ultra-processed food intake. The methodology has been previously described.¹⁰ Samples from cases and controls were processed in the same batch, and laboratory staff was blinded to any participant

characteristics. The relative amount of each fatty acid, expressed as per cent of total fatty acids, was quantified by integrating the area under the peak and dividing the result by the total area. Coefficients of variation for fatty acids ranged from 1.81% for large peaks to 9.75% for the smallest peaks.

Statistical analyses

Descriptive statistics (median/p10–p90) and proportions were calculated overall and stratified by tumour subtypes considering oestrogen receptors positive (ER+), ER– and TN tumour classification. Matching variables included: age ± 3 years, country, place of residence and health system membership. Wilcoxon tests were used to compare main characteristics and ultra-processed food intake (kcal) between cases and controls.¹⁷ Conditional logistic regression was used to calculate OR and 95% CI for the association of ultra-processed foods intake and breast cancer overall and by receptor status adjusting for potential confounding factors. The quantity of energy (kcal) of ultra-processed foods intake was modelled as tertiles based on the distribution of the controls. Tests for trend were calculated based on the scores of the tertiles. Stratified analyses were conducted on ER + and ER–, and TN tumours and compared with controls groups. Our final multivariate models were adjusted for factors that changed our risk estimate by more than 10% including education level (proven to be a good marker of overall socio-economic status in Latin American countries classified as \leq primary school, secondary school, $>$ secondary school), age at first full-term pregnancy (nulliparous/ $<$ 20, 20– $<$ 25, $>$ 25 years), number of full-term pregnancy (continuous), breast feeding (yes/no), physical activity (in hours per day of moderate intensity physical activity) body mass index (BMI) (continuous) and total caloric intake (continuous) (model 1). Further adjustment was conducted by adding to model 1 the quantity of energy from other nova groups (NOVA 1, NOVA 2 and NOVA 3) (model 2) as independent variables. This adjustment was used to further control the potential confounding effect of caloric intake from other food items than ultra-processed foods. These analyses were repeated using the percentage of total energy intake from ultra-processed foods as sensitivity analyses. The consumption of ultra-processed foods subgroups was compared in cases and controls and logarithmic transformation was used to normalise skewed variables. Thereafter, the association between ultra-processed food subgroups and breast cancer was explored in multivariate conditional regression models using the Benjamini-Hochberg correction to control for multiple comparisons.¹⁸

Partial Spearman correlation between blood levels of total industrial trans-fatty acids resulting from industrial processing (in percentage of total fatty acids) and ultra-process food intake (in grams) was calculated among the 142 cases and 142 controls with available data (joining data from cases and controls to increase sample size) adjusting for case/control status and laboratory batches.

Further sensitivity analyses were conducted including only ultra-processed food items with potential high content of industrial trans-fatty acids (no detailed food composition was available to identify individual content in trans-fatty acids isomers). Two-sided *p* values <0.05 were considered statistically significant. All analyses were conducted using SAS software V.9.4.

RESULTS

Characteristics of the population at inclusion

Compared with controls, a higher percentage of cases had no children (cases: 17.8%; controls: 9.1%) (table 1). Cases had a first full-term pregnancy at older ages (cases: 23 years; controls: 21 years), breast fed their children for shorter periods (breast feeding for at least 12 months: cases: 46.1%; controls: 65.5%), had more frequently a history of benign breast disease (cases: 36.4%; controls: 12.6%) and shorter duration of daily moderate intensity physical activity (cases: 2.0 hours; controls: 3.6 hours), and higher percentage completing secondary school (cases: 39.4%; controls: 25.9%) than controls. In addition, cases were taller (1.57 vs 1.56 m), leaner (64 vs 68 kg) and had lower BMI (26.1 vs 28.1 kg/m²) compared with the controls. Among tumours with available IHC (n=332), 72% were ER+, 69% were progesterone receptor positive, while 21% of the tumours were TN.

Energy intake

Total caloric intake was slightly higher among cases than controls (cases: 2853 kcal/day; controls: 2694 kcal/day) as well as the caloric intake from ultra-processed foods (NOVA 4) (cases: 729 kcal/day; controls: 641 kcal/day) (table 1). However, differences were not statistically significant.

Relative contribution of subgroups to ultra-processed food consumption in diet

The food subgroups contributing to ultra-processed foods in our population overall and by countries are presented in figures 1 and 2. Overall the major contributors were ready-to-eat/heat foods (18.2%), cakes and desserts (16.7%), carbonated and industrial fruit juice beverages (16.7%), breakfast cereals (12.9%), sausages and reconstituted meat products (12.1%), yoghurt and dairy-based drinks (7.6%), industrial bread (6.1%) and package savoury snacks (6.1%). Some differences were observed between countries. In Costa Rica and Colombia the highest contributors were carbonated and industrial fruit juice beverage and ready-to-eat/heat foods while in Chile and Mexico ready-to-eat/heat foods and cakes and desserts were the most frequently eaten ultra-processed foods (figure 2).

Higher energy intake among cases than controls were observed for industrial fruit juice beverages (*p*=0.0001), industrial bread (*p*=0.0001), breakfast cereals (0.0001), packaged sweet snacks (*p*=0.0001), packaged savoury

snacks (*p*=0.001) and margarine and butter substitute (0.006) (online supplemental table 1).

Ultra-processed food intake and breast cancer

In the multivariate conditional regression model (table 2, model 1), a positive association was observed between overall ultra-processed food intake and risk of breast cancer (OR_{T3-T1}=1.49, 95% CI=0.96 to 2.33, *P*-trend=0.07). Further adjustment for energy intake from other NOVA groups (NOVA 1–NOVA 3) strengthened this association (OR_{T3-T1}=1.93, 95% CI=1.11 to 3.35, *P*-trend=0.02, model 2). In stratified analyses by breast cancer subtypes, a positive association of ultra-processed foods intake and ER +breast cancer was observed (OR_{T3-T1}=1.70, (95% CI=0.82 to 3.52, *P*-trend=0.15 model 1 and OR_{T3-T1}=2.44, 95% CI=1.01 to 5.90, *P*-trend=0.049 model 2) while among ER– breast tumours, the overall ultra-processed food intake was not associated with the risk of breast cancer (OR_{T3-T1}=1.28, 95% CI=0.44 to 3.72, *P*-trend=0.64 model 1, OR_{T3-T1}=1.87, 95% CI=0.43 to 8.13, *P*-trend=0.36 model 2). Similar results were observed when percentage of total calories from ultra-processed foods was used as exposure variable (online supplemental table 2).

We explored further the role of some specific subgroups of ultra-processed foods. The most strongly associated with the risk of breast cancer were: industrial fruit juice beverages (OR_{T3-T1}=2.97, 95% CI=2.01 to 4.38, *P*-trend <0.0001 model 2); industrial breads (OR_{T3-T1}=1.98, 95% CI=1.33 to 2.94, *P*-trend=0.001 model 2), breakfast cereals (OR_{T3-T1}=1.73, 95% CI=1.17 to 2.56, *P*-trend=0.007 model 2), packaged sweet snacks (OR_{T3-T1}=1.75, 95% CI=1.22 to 2.50, *P*-trend=0.003 model 2), cakes and desserts (OR_{T3-T1}=1.83, 95% CI=1.20 to 2.79, *P*-trend=0.005 model 2) and ready-to-eat/heat foods (OR_{T3-T1}=1.97, 95% CI=1.30 to 2.99, *P*-trend=0.002 model 2) (online supplemental table 3). Similar results were observed among ER +breast cancer. Among ER– breast cancer and TN breast cancer strong associations were observed with ready-to-eat/heat foods (OR_{T3-T1}=5.53, 95% CI=1.71 to 18.0, *P*-trend=0.003 and OR_{T3-T1}=4.98, 95% CI=1.47 to 16.8, *P*-trend=0.01, respectively). Our results remained significant after correction for multiple comparisons.

Based on the proportion of calories intake from ultra-processed foods and assuming a linear relationship, we calculated that an increase of 20% in calories from ultra-processed foods was related to a 46% increase in the risk of breast cancer. Similar results were observed among ER + breast cancer, while no association was observed with ER– tumours risk.

Blood concentrations of total industrial trans-fatty acids resulting from industrial processing showed an increasing gradient ranging from 0.38% (Costa Rica) to 0.60% of total fatty acids (Mexico) and were marginally correlated with overall ultra-processed food intake (*r*=0.11 *p*<0.08). This correlation increased to *r*=0.21 (*p*=0.0005) when we restricted the ultra-processed items including yoghurts and dairy drinks, industrial bread, confectionary, cakes

Table 1 Main characteristics of the population

Variable	N	Controls	Cases	P value*
		N=525	N=525	
		Median (p10–p90) or N (%)	Median (p10–p90) or N (%)	
Age (years)	1050	40 (31–45)	40 (31–45)	0.06
Age at menarche (years)	1050	12 (10–15)	12 (11–15)	0.93
Pregnancy ever	911	478 (91)	433 (84)	<0.0001
Age at first full-term pregnancy†	908			<0.0001
<20		180 (37.7)	123 (28.5)	
20–25		174 (36.5)	128 (29.7)	
≥25		123 (25.8)	180 (41.8)	
Age at first full-term pregnancy†(years)	908	21 (16–29)	23 (17–33)	<0.0001
Age at last full-term pregnancy† (years)	908	28 (22–36)	30 (22–36)	0.05
Number of children	1036			<0.0001
0		47 (9.1)	92 (17.8)	
1		112 (21.6)	157 (30.4)	
≥2		360 (69.3)	268 (51.8)	
Ever breast feed, for parous women	911	443 (92.7)	383 (88.5)	<0.0001
Cumulated duration of breast feeding† ≥12 months,	911	313 (65.5)	200 (46.1)	<0.0001
History of benign breast disease	1050	66 (12.6)	191 (36.4)	<0.0001
Family history of breast cancer	1050	23 (4.4)	33 (6.3)	0.18
Daily alcohol intake‡ (g/day)	898	0.92 (0.05–4.95)	0.83 (0.04–5.53)	0.15
Daily physical activity§ (hours/day)	1050	3.6 (0.93–8.9)	2.0 (0.36–8.4)	<0.0001
Education level	1050			<0.0001
≤primary school (low)		103 (19.6)	65 (12.4)	
secondary school (medium)		286 (54.5)	253 (48.2)	
>secondary school (high)		136 (25.9)	207 (39.4)	
Ever smoker (yes)	530	285 (54.3)	245 (46.7)	0.009
<i>Anthropometry</i>				
Height (m)	1047	1.56 (1.50–1.65)	1.57 (1.49–1.66)	0.003
Weight (kg)	1049	68 (55–89)	64 (52–80)	<0.0001
BMI (kg/m ²)	1047	28.1 (22.3–36.2)	26.1 (20.9–32.6)	<0.0001
<i>Energy intake and UPF¶</i>				
Energy (kcal)	1050	2694 (1642–4732)	2853 (1790–4672)	0.09
Energy (kcal) for NOVA 1	1050	1314 (736–2427)	1372 (801–2472)	0.25
Energy (kcal) for NOVA 2	1050	102 (35–289)	102 (27–247)	0.16
Energy (kcal) for NOVA 3	1050	463 (208–893)	544 (236–997)	0.09
Energy (kcal) for NOVA 4	1050	641 (273–1599)	729 (322–1449)	0.23
<i>Biomarkers measured in blood</i>				
Plasma industrial trans-fatty acids (%)	284	0.27 (0.16–0.49)	0.29 (0.16–0.57)	0.05
<i>Tumour characteristics</i>				
Hormone receptor status**	332‡‡			
Oestrogen receptor positive		–	240 (72)	
Progesterone receptor positive		–	230 (69)	
HER2 positive		–	56 (15)	

Continued

Table 1 Continued

Variable	N	Controls	Cases	P value*
		N=525	N=525	
		Median (p10–p90) or N (%)	Median (p10–p90) or N (%)	
Triple negative: ER–/PR–/HER2–		–	70 (21)	
Of which Basal-like††		–	61 (18)	

*P value Wilcoxon test from logistic regression conditional on matching factors.

†Among parous women only.

‡Among consumers.

§Daily moderate intensity physical activity.

¶NOVA group 1: Unprocessed or minimally processed foods; NOVA group 2: Processed culinary ingredients, NOVA group 3: Processed foods; NOVA group 4: Ultra-processed food and drink products.

**Immunohistochemistry is so far available for 332 cases. Percentages given for tumour characteristics are based on these 332 cases. Receptors status presented irrespective of other receptors.

††Basal-like: (TN and EGFR +and/or CK5/6+).

BMI, body mass index; CK5/6, cytokeratin 5/6; EGFR, epidermal growth factor receptor; ER, oestrogen receptor; HER2, human epidermal growth factor receptor 2; PR, progesterone receptor; TN, triple negative; UPF, ultra-processed foods.

and deserts, ready-to-eat/heat foods and margarine and butter substitutes. (online supplemental table 4)

DISCUSSION

In our population of young Latin American women, ultra-processed food intake was associated with an increased risk of breast cancer. An increase of 20% in calories from ultra-processed foods was related to a 46% increase in the risk of breast cancer. Similar results were observed among ER + breast cancer, while no association was observed with ER– tumours risk. The food groups most strongly associated to the risk of breast cancer were industrial fruit juices, industrial bread, packaged sweet snacks and ready-to-eat/heat foods.

This is the first study on the association of ultra-processed foods intake and breast cancer in young women from Latin American countries where a rapid nutritional transition is taking place.^{3 7} Ultra-processed foods are usually associated with high energy density, high intake

of sugar, low levels of micronutrients, greater content of unhealthy fats and lower content of protein and fibre.¹⁹ Previous studies have shown an association with increased risk of chronic diseases in particular obesity,^{8 20 21} hypertension,²² metabolic syndrome and dyslipidaemia^{8 23} and cardiovascular and cerebrovascular diseases.²⁴

Recently a large prospective study has reported a significant association of ultra-processed food intake and cancer incidence in the French population.¹¹ In this study, the contribution of ultra-processed foods to the overall diet in women was 18.7% while in our population, the contribution was 23.7% among the controls and 25.5% among the cases. An association of ultra-processed food intake was observed on overall cancers as well as postmenopausal breast cancer but no effect was observed among premenopausal breast cancer. This may be due to the small number of premenopausal breast cancers (n=264) in this study.¹¹

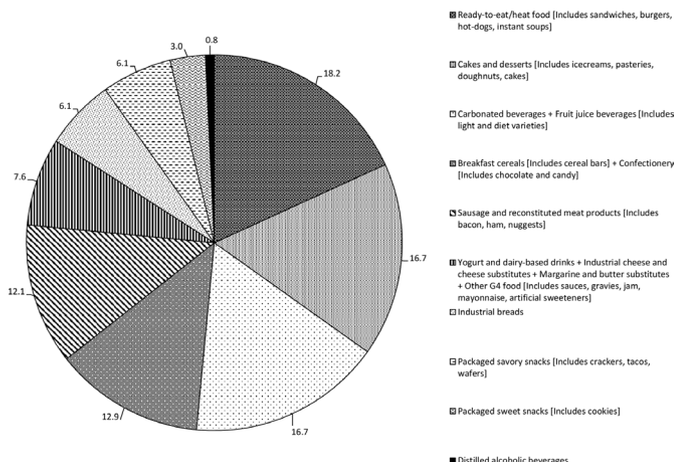


Figure 1 Food groups contributing to ultra-processed foods in our population overall.

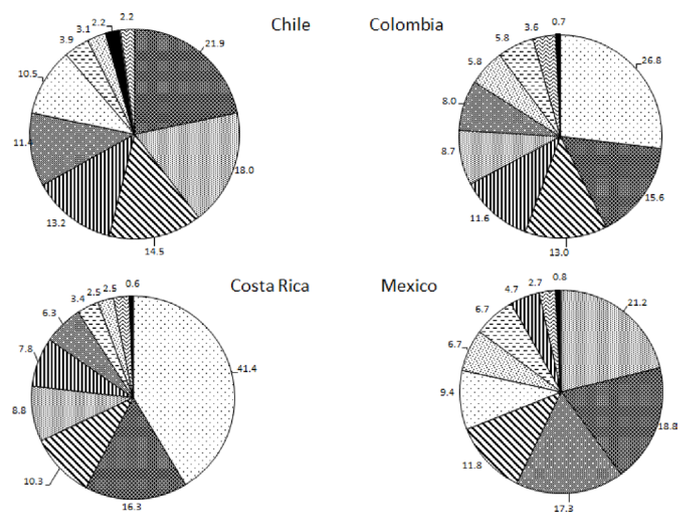


Figure 2 Food groups contributing to ultra-processed foods in our population by countries (Chile, Colombia, Costa Rica and Mexico).

Table 2 ORs and 95% CI for associations between ultra-processed food intake and risk of breast cancer, overall and by receptor status

Cases/controls	Model 1*	Model 2†	
	OR (95% CI)	OR (95% CI)	
All breast cancer‡			
Tertile 1	142/175	1.00 (ref.)	1.00 (ref.)
Tertile 2	184/175	1.27 (0.88 to 1.83)	1.38 (0.95 to 2.03)
Tertile 3	199/175	1.49 (0.96 to 2.33)	1.93 (1.11 to 3.35)
P-trend		0.07	0.02
By receptors status			
ER positive			
Tertile 1	58/80	1.00 (ref.)	1.00 (ref.)
Tertile 2	92/80	1.37 (0.79 to 2.36)	1.45 (0.81 to 2.58)
Tertile 3	90/80	1.70 (0.82 to 3.52)	2.44 (1.01 to 5.90)
P-trend		0.15	0.05
ER negative			
Tertile 1	24/31	1.00 (ref.)	1.00 (ref.)
Tertile 2	32/31	1.17 (0.44 to 3.10)	1.55 (0.55 to 4.37)
Tertile 3	36/30	1.28 (0.44 to 3.72)	1.87 (0.43 to 8.13)
P-trend		0.64	0.36

*Model 1: ORs were estimated by logistic regression conditioned on age (± 3 years), city district of residence and health insurance institution and adjusted for education, (\leq primary/secondary/ $>$ secondary), moderate intensity physical activity (continuous), number of full-term pregnancies (continuous), age at first full-term pregnancy (nulliparous/ < 20 ; 20–25; ≥ 25), breast feeding ever (yes/no), BMI (continuous) and total energy intake (continuous).

†Model 2: Additionally adjusted for energy intake from the other NOVA groups (NOVA 1, NOVA 2, NOVA 3 added simultaneously in the model).

‡Cut-off points for tertiles are respectively: Tertile 1= ≤ 495.5 ; tertile 2= $495.5-896.6$; tertile 3= > 896.6 kcal/day.

BMI, body mass index; ER, oestrogen receptor .

Several potential mechanisms might explain our findings. Ultra-processed foods have been associated with a higher glycaemic response and lower satiety effect.⁸ A recent meta-analysis of prospective studies reported a significant positive association between glycaemic index and glycaemic load and breast cancer.²⁵ Supporting this hypothesis, high intake of carbohydrate has been linked to a higher risk of breast cancer among Mexican women.²⁶ The strong association with industrial fruit juice and packaged sweet snacks in our study supports this hypothesis. Other dietary factors may be involved as a combination of low fat and high fibre intake has been associated to a significant reduction of breast cancer in premenopausal women.²⁷

Ultra-processed products are also rich in fat and industrially transformed fat, in particular trans-fatty acids formed from fats and oil by partial hydrogenation during industrial processing techniques. A large cohort study conducted in European countries (European Prospective Investigation into Cancer and Nutrition (EPIC)) has shown an increase in breast cancer risk in association with blood industrial trans-fatty acids in particular for ER- breast cancer.¹⁰ These fatty acids are found in fast food, and EPIC levels measured in blood have been used as biomarkers of dietary intake

of industrial trans fats.²⁸ In the present study, ultra-processed food intake was significantly correlated to the blood levels of total industrial trans-fatty acids. This may suggest that ultra-processed foods, through their content in trans fat, could affect the risk of breast cancer in premenopausal women.

Little research has examined whether ultra-processed foods have effects on health independent of their nutrient content. Several studies focusing on obesity have observed that the association between ultra-processed food intake and obesity persists after accounting for nutrient intake (eg, saturated fat, trans fat, added sugar and fibre intake, sodium) suggesting that ultra-processed foods may promote additional adverse health outcomes independent of nutrient content.⁸ The wide range of additives that are added to ultra-processed foods as well as other products such as acrylamide produced during heat treatment or contaminants migrating from plastic containers such as bisphenol A and phthalate which have endocrine disruptors properties could also play a role but no firm data are currently available.^{29 30}

Our study is the largest ongoing multicentric population-based case-control study of premenopausal breast cancer in Latin America. Strengths include the

study of young women with large variety and wide range of exposure in dietary intake, the use of standardised methodology to collected dietary data across centres using validated semi-quantitative food frequency questionnaires and standardised methods for food items classification regarding processing classification with nutritional experts, the use of standardised tools and questionnaires across centres to collect information on various current and past lifestyle factors and the centralisation of IHC analyses, thus limiting the inter-laboratory variability. Still some limitations need to be acknowledged. We used a case-control design in order to assemble primary incident breast cancer cases in young women and recall of exposure may be an issue. However, participants were not aware of the hypothesis of the study and had limited knowledge on dietary factor associated to breast cancer. While food frequency questionnaires provide less detailed information on food processing than data from 24 hours recalls, our questionnaire included a list of specific ready-to-eat/heat foods and type of food consumed and mainly aimed at assessing usual dietary intakes in the few years before breast cancer diagnosis (which cannot be obtained from 24 hours dietary recalls). It should be acknowledged that the few years prior to breast cancer diagnosis might not represent the exposure relevant for breast cancer development; however, several studies have shown that adult diet tends to track overtime.³¹ For some products, the food processing techniques might have changed. Thus, some misclassification could be expected. This would most likely be a random misclassification that would tend to lower the observed associations. Finally, our models were adjusted for potential confounding factors, total caloric intake and dietary intake from other NOVA groups, though potential residual confounding is still possible given that ultra-processed food intake is usually associated with unhealthy lifestyle. In addition, the modest sample size for ER- and TN tumours limits the interpretation of the results for these breast cancer subtypes.

Our results support an adverse effect of ultra-processed food intake on the risk of breast cancer in young Latin American women. Further studies are needed to confirm our results and disentangle mechanistic pathways. Given the already proven chronic adverse health effects of ultra-processed foods and the modifiable nature of the diet, a decrease in ultra-processed foods intake should be encouraged particularly in Latin America where an increasing trend of ultra-processed food intake is observed.

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Contributors IR, SR, GTM, GIS, CP, ACRO, MLG and PLP designed the study. GTM, AA-L, IA-C, GIS, MEM, CP, ACRO and MLG and members of the PRECAMA team (JT, EN, RJ, YRC, AA, CAO, WHA, GB, ACRA, CE, FH, IDY, AH, BC, PG, RO, DG, VL, LM and JLS) collected the data and sample materials. PLP designed the specific protocols for IHC and genetic analyses. PLP and MG, together with members of the PRECAMA team (ED, JG, TD, KW and HL), conducted molecular analyses. IR, IH, AK and NK developed the dietary data basis and NOVA classification together with members of the PRECAMA team (MLG, GTM, GIS and CP). VC developed the laboratory assay for fatty acids measurements, and VC and EKA conducted the analyses. CB assembled the database and analysed the data. IR, SR, NK, IH, GTM, GIS, CP, ACRO, MLG, MH, VC and MLG interpreted the data. All authors participated in writing the manuscript. All authors read and approved the final manuscript. SR is the guarantor for the study

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Data availability statement Data are available upon reasonable request. PRECAMA data and biospecimens are available for investigators who seek to answer important questions on health and disease in the context of research projects that are consistent with the legal and ethical standard practices of IARC/WHO and the PRECAMA Centres. The primary responsibility for accessing the data belongs to the PRECAMA centres that provided them. The use of anonymised data from the PRECAMA study can be requested by contacting the corresponding author. The request will then be passed to members of the PRECAMA Steering Committee for deliberation.

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