Ingested Nitrate and Breast Cancer in the Spanish Multicase–Control Study on Cancer (MCC-Spain)

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BACKGROUND: Ingested nitrate leads to endogenous formation of $N$-nitroso compounds that are breast carcinogens in animals, but human evidence is limited.

OBJECTIVE: We evaluated ingested nitrate as a risk factor for breast cancer (BC) in a multicase–control study.

METHODS: Hospital-based incident BC cases and population-based controls were recruited in eight Spanish regions in 2008–2013; participants provided residential and water consumption from 18 years of age and information on known BC risk factors. Long-term nitrate levels (1990–2010) were estimated and linked with residential histories and water consumption to calculate waterborne ingested nitrate (milligrams/day). Dietary ingested nitrate (milligrams/day) was calculated using food frequency questionnaires and published dietary nitrate contents. Interactions with endogenous nitrosation factors and other variables were evaluated. A total of 1,245 cases and 1,520 controls were included in the statistical analysis.

RESULTS: Among the study regions, average ± SD waterborne ingested nitrate ranged from 2.9 ± 1.9 to 13.5 ± 7.5 mg/day, and dietary ingested nitrate ranged from 88.5 ± 48.7 to 154 ± 87.8 mg/day. Waterborne ingested nitrate was not associated with BC overall, but among postmenopausal women, those with both high nitrate (>6 vs. <2.6 mg/day) and high red meat intake (≥20 vs. <20 g/day) were more likely to be cases than women with low nitrate and low red meat intake (adjusted odds ratio = 1.64; 95% confidence interval: 1.08, 2.49; overall interaction $p$-value = 0.17). No association was found with dietary nitrate.

CONCLUSIONS: Waterborne ingested nitrate was associated with BC only among postmenopausal women with high red meat consumption. Dietary nitrate was not associated with BC regardless of the animal or vegetable source or of menopausal status.

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Introduction

Breast cancer (BC) is the leading cause of cancer mortality and is the most common cancer among women worldwide. In Spain, 25,215 new cases are diagnosed annually (Ferlay et al., 2013), and incidence rates have increased over the last three decades (Pollan et al., 2009). Several risk factors for BC have been identified, including sex, age, nulliparity, short breastfeeding, menstrual and reproductive history, high body mass index (particularly in postmenopausal women), physical inactivity, high alcohol or energy intake, use of drugs with estrogogenic action, exposure to ionizing radiation, specific genetic factors, family history of BC, previous diagnosis of non-malignant breast diseases, and high mammographic density (Hankinson et al., 2004; Romieu et al., 2015; Stewart and Wild, 2014). Established risk factors explain ~50% of the incidence variation of this malignancy, and other environmental exposures may partly explain the remaining variation (Brody et al., 2007).

Nitrate is a frequent contaminant in drinking water worldwide; its presence is related to excessive fertilizer use or to sewage (Wakida and Lerner, 2005). Humans are exposed to nitrate through diet and through drinking water ingestion. The maximum nitrate level in drinking water [50 mg/L as nitrate ion (NO$_3^-$)] or 10 mg/L of nitrate-nitrogen (nitrate-N) (EU, 1998; WHO, 2008b) was established to prevent acute health effects in children (methemoglobinemia), but the effects of long-term exposure to lower levels, including cancer risk, are not well established (Ward et al., 2005).

Ingested nitrate is classified as a probable human carcinogen in conditions of endogenous nitrosation (IARC, 2010). This process involves the conversion of nitrate into nitrite and the synthesis of $N$-nitroso compounds (NOCs) in the gastrointestinal tract. The intake of antioxidant vitamins and the use of nonsteroidal antiinflammatory drugs (NSAIDs) inhibit endogenous nitrosation, whereas meat intake and inflammatory gastrointestinal conditions promote it (Ward et al., 2005). NOCs are potent carcinogens in exposed animals, and NOC-related DNA adducts are found in humans (Hendzel et al., 1998; WHO, 2008b). Humans are exposed to NOCs through diet (methyl esters of NOCs) and from the ingestion of drinking water (nitrite-N). NOCs are found in drinking water in concentrations ranging from 5 to 20 μg/L (Liu et al., 2002; WHO, 2008b). Water intake is an important source of NOCs in the diet, and the quantity of NOCs ingested can be estimated from drinking water intake (WHO, 2008b).

Ingested nitrate has been associated with breast cancer risk in case–control studies from Europe (Borrell et al., 2002; Estévez et al., 2014), Spain (Isasi et al., 2007), the United States (Baker et al., 2005), and China (Zhang et al., 2011). These studies were mainly conducted among postmenopausal women, with a small number of cases and controls being included (Baker et al., 2005; Borrell et al., 2002; Estévez et al., 2014; Isasi et al., 2007; Zhang et al., 2011). The objective of the present study was to evaluate the association between ingested nitrate and breast cancer risk in a large multicase–control study conducted in Spain.
Ingested nitrate and breast cancer

For several animal species (Lijinsky et al. 1992). Some NOCs, such as N-methyl-N-nitrosourea (MNU), are used to induce BC in experimental animal studies, and young rats exposed to MNU were more susceptible to developing breast tumors (Tsubura et al. 2011). In cell-based studies, low doses of nitrite and nitrate were able to mimic estradiol and to activate estrogen receptors, suggesting a potential role of these anions in the etiology or progression of cancer (Veselik et al. 2008).

Despite the evidence in animals, few epidemiologic studies have evaluated the association between exposure to nitrate or to its derivatives and BC. Relevant studies were conducted in the United States (Brody et al. 2006; Weyer et al. 2001); these studies did not find associations between waterborne or dietary ingested nitrate and BC. A recent cohort study of postmenopausal women in the United States reported that BC was increased in the highest versus lowest quintile of water nitrate intake among women who also had folate ingestion of ≥ 400 μg/day, but the study did not find any association with dietary nitrate (Inoue-Choi et al. 2012).

The authors of previous studies attributed their null associations to limitations in the exposure assessment (i.e., a lack of data on water daily intake), to the coexistence of other NOCs, such as N-nitrosonitrosamines (N-Nitosourea), and to the lack of exposure to NOCs from specific dietary sources such as animal foods and processed meat. In summary, human evidence relating nitrate exposure and BC is limited and inconclusive. Studies evaluating different exposure windows, including individual water consumption information, endogenous nitrosation factors and other covariates, are required to enhance the available evidence.

In the present study, we aimed to evaluate nitrate ingested through drinking water and diet as a risk factor for BC in a population-based multicase–control study conducted in Spain (MCC-Spain).

Methods

Study Design and Population

This study is part of the MCC-Spain study, which aims to evaluate the influence of environmental exposures on common cancers in Spain (e.g., female breast or colorectal). The study population was recruited between 2008 and 2013 in eight Spanish provinces (see Table 1). Cases were identified shortly after diagnosis (average: 3.2 months, SD 4.2 months) through an active search by periodic visits to the collaborating hospital departments (i.e., gynecology, oncology, general surgery, radiotherapy, and pathology departments). Participant hospitals were the reference centers for oncologic diseases in each study area. Only incident cases who were diagnosed within the recruitment period, without malignant BC history, between 20 and 85 years of age, resided in the hospitals’ catchment areas for at least 6 months prior to recruitment, and were able to answer the epidemiological questionnaire (Castaño-Vinyals et al. 2015) were included. All cases had histological confirmation and included all malignant BC [International Classification of Diseases (10th Revision) (World Health Organization 2008a); ICD-10: C50] and frequent in situ breast cancers (ICD-10:D05.1, D05.7).

Table 1. Characteristics of the study population (1,245 cases* and 1,520 controls).

| Characteristic | Cases n(%) | Controls n(%) | p-Value χ² test |
|---------------|-----------|--------------|----------------|
| Smoking       |           |              |                |
| No            | 670 (61.4) | 905 (59.6)   |                |
| Yes           | 569 (45.9) | 613 (40.4)   | 0.003          |
| Missing       | 6          | 2            |                |
| Alcohol intake* |         |              |                |
| No            | 384 (35.4) | 516 (38.5)   |                |
| Yes           | 700 (64.6) | 624 (61.5)   | 0.20           |
| Missing       | 161        | 180          |                |
| Energy intake (kcal/day) |       |              |                |
| ≤ 1,479       | 278 (26.6) | 447 (33.4)   |                |
| >1,479–1,894  | 362 (34.3) | 447 (33.3)   | 0.001          |
| >1,894        | 434 (40.0) | 446 (33.3)   | 0.001          |
| Missing       | 161        | 180          |                |
| Vitamin C intake (mg/day) |     |              |                |
| No            | 396 (36.5) | 447 (33.4)   |                |
| Yes           | 731 (69.2) | 447 (33.3)   | 0.003          |
| Missing       | 161        | 180          |                |
| Folate intake (μg/L) |      |              |                |
| No            | 325 (30.9) | 447 (33.4)   |                |
| Yes           | 374 (34.5) | 446 (33.3)   | 0.82           |
| Missing       | 161        | 180          |                |
| Vegetable intake* (g/day) |   |              |                |
| No            | 393 (36.3) | 447 (33.4)   |                |
| Yes           | 356 (32.9) | 447 (33.4)   | 0.42           |
| Missing       | 161        | 180          |                |
| Red meat intake (g/day) |   |              |                |
| No            | 311 (28.7) | 447 (33.4)   |                |
| Yes           | 346 (31.9) | 447 (33.3)   | 0.31           |
| Missing       | 161        | 180          |                |
| Processed meat intake (g/day) | |              |                |
| No            | 253 (27.0) | 447 (33.4)   |                |
| Yes           | 360 (33.2) | 447 (33.3)   | 0.01           |
| Missing       | 161        | 180          |                |
| Interview quality |       |              |                |
| Unsatisfactory | 3 (0.2)   | 1 (0.1)      |                |
| Questionable  | 92 (7.5)   | 87 (6.1)     |                |
| Reliable      | 562 (46.0) | 736 (51.8)   |                |
| High quality  | 564 (46.2) | 596 (42.0)   | 0.02           |
| Missing       | 24         | 100          |                |
| Histologic type |       |              |                |
| Ductal        | 961 (76.4) |                |                |
| Others        | 162 (13.0) |                |                |
| Undefined     | 132 (10.6) |                |                |
| Estrogen receptors |      |              |                |
| Positive      | 990 (79.5) |                |                |
| Negative      | 218 (17.5) |                |                |
| Undefined     | 37 (2.9)   |                |                |

*Breast cancer cases. Distribution only among postmenopausal women. Distribution among parous women. Alcohol intake from 30 to 40 years of age. Vegetable intake includes vegetables and fruits.
Population-based controls were frequency-matched to cases by age, sex, and region, ensuring at least one control of the same sex and 5-year interval age for each case. Eligible controls were randomly selected from administrative records of primary care health centers located within hospitals’ catchment areas. For each control needed, five potential participants of similar age, sex, and hospital catchment area were randomly selected from the lists of general practitioners. If contact with the first person on the list was not achieved (after at least five attempts made at different times of the day), or if he/she refused to participate, the next person on the list was approached. The study protocol was approved by the ethics review board from each participating center, and participants signed an informed consent before recruitment.

**Questionnaires and Response Rates**

A structured computerized questionnaire was administered by trained personnel in face-to-face interviews (http://www.mccspain.org). Collected data included: a) sociodemographic characteristics; b) lifetime residential history; c) type of water consumed in each residence (municipal/bottled/well/other); d) amount of water intake at home, including water per se, coffee, tea, and other water-based beverages; e) smoking habits; f) history of genital ulcers and use of nonsteroidal anti-inflammatory drugs (NSAIDs); g) gynecologic and reproductive history; h) use of oral contraceptives (OCs) or hormonal replacement therapy (HRT); and i) physical activity. Anthropometric measurements were self-reported (weight, height) or measured (waist and hip circumference) during face interviews (http://www.mccspain.org). Questionnaires and Response Rates administered by trained personnel in face-to-face interviews. Questionnaires and Response Rates were self-reported (weight, height) or measured (waist and hip circumference) during face interviews (http://www.mccspain.org).

**Nitrate Levels in Drinking Water**

We collected environmental data from municipalities covering ≥ 80% of person-years in each area. We sent a standardized questionnaire to local authorities and water companies to ascertain current and historical nitrate measurements in water from municipal distribution systems and water source characteristics (surface/groundwater proportion). We also calculated the average waterborne nitrate consumption in this area according to the postal code of the residence. Nitrate levels in well water were not available for other areas, and waterborne ingested nitrate was considered missing for years when well-water consumption was reported among women from those areas (range: 0.6–8.2% of controls in the longest residence). The annual nitrate estimates were averaged and multiplied by the average daily water intake at home (1.3 ± 0.7 L/day in cases, 1.2 ± 0.7 L/day in controls). Water intakes > 99th percentile (4 L/day), considered implausible, were treated as missing values in the analyses. We also calculated the average waterborne ingested nitrate in two alternative exposure periods: from 15 to 2 years before the interview (“recent” exposure), and from 18 to 30 years of age (“early adulthood” exposure).

**Estimation of Long-Term Nitrate Levels in Drinking Water**

We calculated annual average nitrate levels back to 1940 by water zone (defined as a geographical area supplied by water with a homogeneous source and quality) that usually corresponded to municipality. We calculated annual averages based on available nitrate measurements. For years without measurements, we assigned the average of the total measurements available in the water zone, as long as the water source remained constant. In cases where the water source changed, the ground water percentage was used as a weight to modulate the estimations, assuming that nitrate levels were higher at higher ground water proportions. In municipalities without any nitrate measurement (covering 0.5% of the total person-years), we imputed the levels of neighboring municipalities supplied with similar ground water proportion ± 10%.

**Individual Exposure Variables**

We linked nitrate levels in drinking water (measured and imputed) and residential histories by year and municipality covering the exposure window from 18 years of age to 2 years before the study interview (henceforth referred to as “adult life” or “long-term exposure”). To calculate waterborne ingested nitrate (milligrams/day), we assigned nitrate levels [milligrams/liter nitrate ion (NO₃⁻)] in drinking water by year according to the water type consumed. Nitrate levels in municipal water (residential levels) were assigned for tap-water consumption. Levels in the sampled bottled waters were averaged using the sales frequency of each brand as a weight. This weighted average (6.1 mg/L of NO₃⁻) was assigned when bottled water consumption was reported. Levels in well-water samples from León (range: 0.5–93 mg/L) were assigned to women reporting well-water consumption in this area according to the postal code of the residence. Nitrate levels in well water were not available for other areas, and waterborne ingested nitrate was considered missing for years when well-water consumption was reported among women from those areas (range: 0.6–8.2% of controls in the longest residence). The annual nitrate estimates were averaged and multiplied by the average daily water intake at home (1.3 ± 0.7 L/day in cases, 1.2 ± 0.7 L/day in controls). Water intakes > 99th percentile (4 L/day), considered implausible, were treated as missing values in the analyses. We also calculated the average waterborne ingested nitrate in two alternative exposure periods: from 15 to 2 years before the interview (“recent” exposure), and from 18 to 30 years of age (“early adulthood” exposure). In a subset of participants from Barcelona with information on water type changes within residences, 86% of subjects reporting bottled water consumption in the last residence actually switched from municipal to bottled water after the year 2000. Potential misclassification of the water type consumed (municipal/bottled), particularly in recent residences, was a concern. To address this issue, we calculated an alternative variable for waterborne ingested nitrate in adult life. We assumed that women reporting bottled water consumption and living during at least 10 years in the last residence (or in the previous one), actually consumed municipal water before the year 2000 and bottled water thereafter.
Statistical Analyses

The population analyzed (1,245 cases, 1,520 controls) included women with data on both waterborne ingested nitrate covering ≥ 70% of the main exposure period (from 18 years of age to 2 years before the interview) and on daily water intake. We estimated odds ratios (ORs) and 95% confidence intervals (CIs) of BC for categorized nitrate intake using unconditional logistic regression. Categories of exposure (quartiles or tertiles) were specifically defined for pre- and postmenopausal women according to the distribution in controls. Basic models were adjusted for age (continuous), study area, and education (three categories: ≤ primary, high school, and ≥ university). Several potential confounders were explored separately for pre- and postmenopausal women, including smoking (yes/no 5 years before recruitment), average leisure physical activity from 16 years of age until 2 years before the interview (measured in metabolic equivalents of task (METS)/hour/week), body mass index (BMI), family history of malignant BC in any blood relative (yes/no), NSAID use (yes/no), age at menarche, age at menopause (both, continuous variables in years and categorized variables). Menopause and age at menopause were defined according to the date of the last regular menstrual period. Age at first birth, nulliparity (yes/no), parity (number of births), total months of breastfeeding (categorized), OC and HRT use (never/ever), intake of alcohol (no/yes at 30 years of age), intake of energy and folate (tertiles), and endogenous nitrosation modulators (intake of vitamin C, vitamin E, red meat, and processed meat) were also explored as potential confounders. Only established BC risk factors (Stewart and Wild 2014), and variables that changed the risk estimates > 10% were included in the adjustment (age, study area, education, BMI, family history of BC, age at first birth, use of OC, energy intake and age at menopause for postmenopausal women). For each model covariate, missing data in categorical variables were classified as missing data in covariables and women with unsatisfactory or missing interview quality. Interview quality was assessed by the interviewers as unsatisfactory, questionable, reliable, or high quality based on the completeness of the information provided. All statistical analyses were performed using STATA version 12.0 (StataCorp LP).

Results

General characteristics of the study population are shown in Table 1. Compared with controls, cases showed higher frequency of family history of BC; age at menopause > 50 years; age at first birth > 30 years; higher intake of energy, red meat, and processed meat; lower intake of vitamin C; and nulliparity (p-value < 0.05 in χ² test). Among the women analyzed, 24.6% (n = 679) were premenopausal and 75.4% (n = 2,086) were postmenopausal. Women with assigned nitrate levels in drinking water for < 70% of their residential history in adult life and those without information on daily water intake were excluded from the analyses. Compared with those who were excluded, the women who were analyzed showed a higher proportion of controls (55% vs. 47%) and

Figure 1. Ingested nitrate levels (milligrams/day) through drinking water from 18 years of age to 2 years before the interview (A) and diet (B) across study areas. Women with waterborne ingested levels > 44 mg/day (n = 6) or with dietary ingested levels > 476 mg/day (n = 7) were excluded from the graphics. Boxes extend from the 25th to the 75th percentile. Horizontal bars represent the median, whiskers indicate the 10th and 90th percentiles, and outliers are represented as points.
postmenopausal women (75.4% vs. 63.9%); a lower proportion of university education and nulliparity; were older; and had a lower intake of vitamins C and E; however, their levels of waterborne ingested nitrate were similar (see Table S1).

Figure 1 shows the average ingested nitrate levels in adult life for cases and controls. Across the investigated areas, levels of waterborne ingested nitrate (mean ± SD) ranged from 2.9 ± 1.9 to 13.5 ± 7.5 mg/day (Figure 1A) and were higher among post- versus premenopausal women (6.74 ± 7.1 vs. 5.12 ± 5.6 mg/day; p-value < 0.001 for Mann–Whitney U test). Ingested levels during alternative exposure periods (from 15 to 2 years before the study interview and from 18 to 30 years of age) were similar to the levels presented in Figure 1A (results not shown). Across the investigated areas, dietary ingested nitrate levels (mean ± SD) ranged from 88.5 ± 48.7 to 154 ± 87.8 mg/day (Figure 1B) and were higher among post- versus premenopausal women (129.0 ± 86.2 vs. 109.7 ± 62.1 mg/day; p-value < 0.001 for t-test). On average, 6.0% ± 7.0 of the total dietary nitrate was derived from animal sources, 84.7% ± 12.1 from vegetables, and the remaining portion from other food products (e.g., grains). Ingested nitrate from animal sources (mean ± SD: 5.5 ± 2.9 mg/day) was greater among pre- versus postmenopausal women (5.9 ± 2.7 vs. 5.2 ± 3.0 mg/day; p-value < 0.0001 for t-test), but ingested nitrate from vegetable sources (mean ± SD: 110 ± 79.6 mg/day) was lower among pre- versus postmenopausal women (96.5 ± 60.4 vs. 115.0 ± 84.5 mg/day; p-value < 0.0001 for t-test) (results not shown).

Table 2 shows the association between waterborne ingested nitrate and BC. Among postmenopausal women, the fully adjusted OR (95% CI) was 1.29 (0.92, 1.81) for > 8.8 mg/day compared with the lowest intake levels (< 2.3 mg/day). After excluding postmenopausal women with missing or unreliable interview quality (n = 118), the OR (95% CI) was 1.32 (0.93, 1.86) for > 8.8 mg/day versus < 2.3 mg/day. Among premenopausal women, the fully adjusted OR (95% CI) was 1.14 (0.67, 1.94) for > 6.3 mg/day compared with the lowest intake levels (< 1.8 mg/day), and the results were similar after excluding premenopausal women with unreliable interviews (n = 10). The results were also similar when waterborne exposure from 18 years of age to 2 years before the study interview was defined assuming bottled water use after 2000, for exposures from 15 to 2 years before the study interview, and from 18–30 years of age (see Table S2). Exposure–response curves among study areas did not show associations except at the highest levels, where estimates were extremely imprecise (see Figure S1).

Table 3 shows the associations between waterborne ingested nitrate and BC for postmenopausal women across categories of relevant covariables. BC was inversely associated with high vitamin C + E intake (> 181 mg/day) versus low vitamin C + E intake among women with low waterborne nitrate intake (< 2.6 mg/day) (OR = 0.60; 95% CI: 0.39, 0.92), and the overall interaction p-value was 0.08. However, there was no evidence that vitamin C + E intake modified the odds of BC among those in the second or third tertile of waterborne nitrate. This inverse association was not observed when vitamins C and E were analyzed separately.

Across the investigated areas, dietary nitrate from animal sources (mean ± SD: 110 ± 79.6 mg/day) was lower than among women with high waterborne nitrate (> 2.6 mg/day) and low red meat intake (OR = 1.64; 95% CI: 1.08, 2.49). BC was not increased in women with high waterborne nitrate and low red meat intake (OR = 1.08; 95% CI: 0.72, 1.47), but the overall interaction between nitrate and red meat intake was not significant (LRT p-value = 0.17). The results for processed meat intake followed a similar pattern. BC was also more common among women with the highest waterborne nitrate (> 6 mg/day) and the highest red meat intake (> 20 g/day) than among women with low waterborne nitrate (< 2.6 mg/day) and low red meat intake (OR = 1.64; 95% CI: 1.08, 2.49). BC was more common among women with the highest waterborne nitrate intake (> 8.8 mg/day) versus < 2.3 mg/day) and the highest red meat intake (> 20 g/day) than among women with low waterborne nitrate (< 2.6 mg/day) and low red meat intake (OR = 1.32; 95% CI: 0.93, 1.86) for > 8.8 mg/day compared with the lowest intake levels (< 2.3 mg/day). After excluding women with low waterborne nitrate (< 2.6 mg/day) and low red meat intake (OR = 1.64; 95% CI: 1.08, 2.49). BC was not increased in women with high waterborne nitrate and low red meat intake (OR = 1.08; 95% CI: 0.72, 1.47), but the overall interaction between nitrate and red meat intake was not significant (LRT p-value = 0.17).

Stratified analyses among pre- and postmenopausal women resulted in less-precise estimates of associations owing to smaller numbers of observations. Most of the ORs observed across strata were close to 1, and overall interactions were not significant (LRT p-values > 0.10) (see Table S3).

Among all BC cases, 951 (76.4%) were ductal (ICD-10 C50), 162 (13.0%) were other malignant and in situ cancers (ICD-10 D05.1, D05.7), and 132 (10.6%) were undefined. Regarding ER status, 990 (79.5%) were positive, 218 (17.5%) were negative, and 37 (2.9%) had missing ER status. Stratified analyses among pre- and postmenopausal women combined showed similar ORs for ductal and other/undefined tumors as well as for ER-positive and ER-negative cancers (see Table S4).

Overall, BC was not associated with dietary nitrate from animal or vegetable sources (Table 4). The ORs reported were similar after adjusting for endogenous nitration factor (intake of vitamin C, vitamin E, and red and processed meat) and other covariables listed in Table 1 (data not shown) and after excluding women with low interview quality (n = 128 among pre- and postmenopausal women). Similar results were observed in separate analyses for pre- and postmenopausal women (see Table S5).

Discussion

Average waterborne ingested nitrate levels from 18 years of age to 2 years before the interview was 6.2 ± 6.2 mg/day among controls and 6.6 ± 7.4 mg/day among cases. These levels were not associated with BC overall. However, in postmenopausal women, BC was significantly increased (p < 0.05) in women in the highest tertile of waterborne nitrate and with high red meat intake compared with women
in the lowest tertile of waterborne nitrate and with low red meat intake. Dietary ingested nitrate (mean ± SD: 125.7 ± 80.3 mg/day in controls and 123.2 ± 82.3 mg/day in cases) was not associated with BC among pre- or postmenopausal women regardless of the vegetable or animal source.

To our knowledge, this is the first case-control study on ingested nitrate and BC in a European population. Most previous studies of waterborne nitrate exposure and BC have reported null associations (Brody et al. 2006; Weyer et al. 2001). A recent cohort study conducted in postmenopausal women from the state of Iowa (n = 2,875 BC cases in total), suggested an association between BC and waterborne nitrate intake in interaction with folate intake (Inoue-Choi et al. 2012). Individual data on daily water intake were not available in that study, but estimated waterborne nitrate intake levels were higher than the levels in our study (median: 20 mg/day vs. 3.8 mg/day, respectively). We did not confirm an interaction with folate, most likely because of the differences in nitrate and folate intake levels, as well as other differences including the cancer subtypes evaluated.

Analyses stratified by endogenous nitrosation factors (intake of vitamin C, vitamin E, and red and processed meat) and by other variables (listed in Table 1) did not show significant differences across categories; the CIs were overlapped and included the null value. BC occurred more frequently among postmenopausal women with the highest waterborne nitrate and red meat intake than among postmenopausal women with low waterborne nitrate intake and low red meat intake, and the overall interaction p-value was > 0.10. However, this joint effect is plausible because red meat contains amines, amides, and heme iron that may increase endogenous formation of NOCs (Bingham et al. 2002). The combined intake of vitamins C and E seemed to exert a protective effect that was limited to postmenopausal women in the lowest tertile of waterborne nitrate intake. These findings require confirmation in future studies because multiple stratifications were conducted, and chance cannot be ruled out.

The associations between waterborne ingested nitrate and BC were slightly higher in postmenopausal women than in premenopausal women. However, insufficient statistical power owing to small sample size may partly explain the null results among premenopausal women. We did not find an interaction between menopausal status and nitrate intake (p-value = 0.63) (data not shown), but we evaluated these groups separately because differences have been observed with other risk factors, such as body mass index, according to menopausal status (Cheraghi et al. 2012). These differences may be attributed to endogenous hormonal production and to other factors that are not well established. BC is a heterogeneous disease with potentially different etiologies in pre- and postmenopausal women; therefore, the evaluation of risk factors among these subgroups may have relevant public health implications.

### Table 3. Interaction of waterborne ingested nitrate from age 18 to 2 years before study interview with relevant dietary covariates and breast cancer associations among postmenopausal women.2

| Waterborne ingested nitrate | Cases (n) | Controls (n) | OR(95% CI) | ORb(95% CI) | Interaction p-valuec |
|----------------------------|-----------|-------------|------------|-------------|----------------------|
| < 2.6 mg/day               | 140       | 162         | Reference  | 104         | 161                  | 0.72 (0.51, 1.05)  |
| ≥ 2.6–6.0 mg/day           | 118       | 169         | 0.88 (0.62, 1.24) | 153         | 171                  | 1.08 (0.77, 1.52)  |
| > 6.0 mg/day               | 146       | 185         | 1.19 (0.80, 1.77) | 136         | 183                  | 1.06 (0.72, 1.58)  |
| < 10 mg/day                | 131       | 170         | Reference  | 113         | 153                  | 0.84 (0.58, 1.22)  |
| ≥ 2.6–6.0 mg/day           | 132       | 160         | 1.18 (0.83, 1.67) | 139         | 180                  | 0.94 (0.65, 1.34)  |
| > 6.0 mg/day               | 123       | 186         | 1.16 (0.77, 1.74) | 159         | 182                  | 1.25 (0.82, 1.89)  |
| < 10 mg/day                | 131       | 170         | Reference  | 113         | 153                  | 0.84 (0.58, 1.22)  |
| ≥ 2.6–6.0 mg/day           | 132       | 160         | 1.18 (0.83, 1.67) | 139         | 180                  | 0.94 (0.65, 1.34)  |
| > 6.0 mg/day               | 123       | 186         | 1.16 (0.77, 1.74) | 159         | 182                  | 1.25 (0.82, 1.89)  |

### Table 4. Odds ratio (OR) of breast cancer associated with dietary ingested nitrate (mg/day) from different sources (n = 2,424).2

| Ingested nitrate from | Cases (n) | Controls (n) | OR(95% CI) | ORb(95% CI) |
|-----------------------|-----------|-------------|------------|-------------|
| Animal sources        |           |             |            |             |
| < 4.0 mg/day          | 307       | 447         | Reference  | Reference   |
| ≥ 4.0–6.0 mg/day      | 364       | 447         | 1.12 (0.92, 1.38) | 1.02 (0.82, 1.26) |
| > 6.0 mg/day          | 413       | 446         | 1.19 (0.97, 1.47) | 1.04 (0.83, 1.31) |
| p for trend           |            |             | 0.09       | 0.72        |
| Vegetables sources    |           |             |            |             |
| < 78 mg/day           | 385       | 447         | Reference  | Reference   |
| ≥ 78–122 mg/day       | 355       | 447         | 0.92 (0.75, 1.12) | 0.90 (0.74, 1.11) |
| > 122 mg/day          | 344       | 446         | 0.90 (0.74, 1.11) | 0.86 (0.69, 1.06) |
| p for trend           |            |             | 0.33       | 0.15        |
| Total diet            |           |             |            |             |
| < 90 mg/day           | 387       | 447         | Reference  | Reference   |
| ≥ 90–138 mg/day       | 349       | 447         | 0.90 (0.74, 1.10) | 0.86 (0.70, 1.06) |
| > 138 mg/day          | 348       | 446         | 0.90 (0.73, 1.10) | 0.83 (0.67, 1.03) |
| p for trend           |            |             | 0.30       | 0.09        |
| Total diet + waterborne|          |             |            |             |
| < 96 mg/day           | 386       | 447         | Reference  | Reference   |
| ≥ 96–144 mg/day       | 341       | 447         | 0.89 (0.73, 1.09) | 0.84 (0.69, 1.04) |
| > 144 mg/day          | 357       | 446         | 0.94 (0.76, 1.15) | 0.87 (0.70, 1.08) |
| p for trend           |            |             | 0.51       | 0.19        |

Trend p-values derived from a likelihood ratio test that compared a model including the categorical nitrate intake variable as an ordinal variable (0, 1, 2) with a model that excluded this variable.

2Only women with complete information on dietary covariates (n = 1,928) or smoking (n = 2,080) were analyzed.

3Adjusted for study area, age, education, body mass index, family history of breast cancer, age at menopause, age at first birth, oral contraceptives use, and energy intake. p-Value for overall interaction was calculated by comparing the multivariate models with and without the interaction term using a likelihood ratio test.
The evaluation of BC’s association with nitrate and other environmental pollutants in different exposure periods is required because there is evidence suggesting that early exposure (e.g., before the first full-term pregnancy) might be the most relevant for inducing breast carcinogenesis (Brody et al. 2007). Although we evaluated three different exposure periods, we did not observe differences in the associations, most likely because of high correlations between exposure levels at different periods. In addition, we did not evaluate early-life exposure owing to a lack of data. This evaluation is warranted in future studies, particularly in settings with more available historical nitrate measurements in drinking water.

In the present study, dietary ingested nitrate levels were lower than levels observed in previous studies on this topic (Inoue-Choi et al. 2012), which may partly explain the lack of statistically significant associations. Our results suggested an inverse association between BC and ingested nitrate from vegetable sources. Vegetables contain endogenous nitrosation modulators (e.g., vitamins C and E), which may explain these results. Previous studies (Hord et al. 2009) have suggested beneficial health effects of nitrate from vegetable sources, which might also explain these results. Further research is needed to confirm these effects and to understand the underlying mechanisms.

Potential exposure misclassification is an area of concern in our study because most of the long-term nitrate levels in drinking water were imputed, particularly before 1980, and we did not account for water intake outside the home. However, the reported amount of water consumed at work (mean ± SD: 0.2 ± 0.3 L/day) and other places (0.01 ± 0.05 L/day) was smaller than that consumed at home (1.2 ± 0.7 L/day), and minor bias was expected. We conducted sensitivity analyses excluding women with the lowest interview quality, and slightly higher ORs were found, particularly among postmenopausal women. Changes in the type of water consumed, particularly in recent residences, may have led to exposure misclassification. To address this possibility, we calculated an alternative variable of waterborne ingested nitrate, which was described in “Methods.” In the analysis of this alternative variable, few women (n = 4 postmenopausal women and n = 6 premenopausal women) changed exposure categories, so the associations observed (see Table S2) were similar to the main results. Potential confounding by other environmental contaminants with estrogenic action and correlated to nitrate in drinking water may have occurred, although available data on selected pesticides (e.g., simazine, atrazine, terbuthylazine) showed levels below or around the quantification limit. Additionally, waterborne nitrate exposure was not evaluated because the available measurement showed unquantifiable or extremely low levels of nitrate.

Dietary nitrate estimations may also be prone to error because nitrate content was not available for some food items, and other relevant data, including vegetable storage and processing (i.e., washing, peeling and cooking), were not collected. Dietary nitrate intake was not evaluated; however, the lack of this information is not a major limitation because the main exposure route is through the endogenous reduction of nitrate (IARC 2010). Finally, because dietary intake was collected with an FFQ, ingested nitrate misclassification because of recall bias may be a concern.

The matched case–control design by area of residence may lead to overmatching in environmental studies, which may have occurred in this study. However, overmatching would not affect the validity of our results (Agudo and Gonzalez 1999). Controls had a higher education level than did the general population, which may hamper the external validity of the results. The heterogeneity of effects between some of the study areas may also be a limitation for the combined analyses.

A major strength of this study was the availability of detailed individual information on residential history, water consumption habits, and relevant covariates. Despite the limitations, the environmental data collected enabled us to evaluate BC associations for a long-term exposure window (from 18 years of age to 2 years before the study interview), in recent years, and in early adulthood. The information provided by the FFQ allowed us to evaluate nitrate ingestion from different dietary sources and to evaluate several potential confounders and effect modifiers that were not previously evaluated, including endogenous nitrosation modulators.

**Conclusion**

Waterborne nitrate ingestion at the exposure levels observed was not associated with BC overall. However, BC was more common among postmenopausal women with the highest waterborne nitrate and red meat intake than among women with low waterborne nitrate and low red meat intake. Dietary nitrate was not associated with BC regardless of the exposure source or of menopausal status.

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