Choline and betaine consumption lowers cancer risk: a meta-analysis of epidemiologic studies

Shanwen Sun1,*, Xiao Li2,*, Anjing Ren1,*, Mulong Du3,4, Haina Du5, Yongqian Shu1, Lingjun Zhu1 & Wei Wang6

A number of human and animal in vitro or in vivo studies have investigated the relationship between dietary choline and betaine and cancer risk, suggesting that choline and betaine consumption may be protective for cancer. There are also a few epidemiologic studies exploring this relationship, however, with inconsistent conclusions. The PubMed and Embase were searched, from their inception to March 2016, to identify relevant studies and we brought 11 articles into this meta-analysis eventually. The pooled relative risks (RRs) of cancer for the highest versus the lowest range were 0.82 (95% CI, 0.70 to 0.97) for choline consumption only, 0.86 (95%CI, 0.76 to 0.97) for betaine consumption only and 0.60 (95%CI, 0.40 to 0.90) for choline plus betaine consumption, respectively. Significant protective effect of dietary choline and betaine for cancer was observed when stratified by study design, location, cancer type, publication year, sex and quality score of study. An increment of 100 mg/day of choline plus betaine intake helped reduce cancer incidence by 11% (0.89, 95% CI, 0.87 to 0.92) through a dose-response analysis. To conclude, choline and betaine consumption lowers cancer incidence in this meta-analysis, but further studies are warranted to verify the results.

Cancer is a major cause of death on a global scale. According to the GLOBOCAN estimates, new cancer cases and deaths are approximately 14.1 million and 8.2 million in 2012 worldwide1. To slow and even reverse the global trend of increasing in cancer ultimately, preventive measures could provide the merely possible approach2.

In the process of cancer prevention, dietary factors have long been regarded as a quite crucial role, among which choline and betaine (choline's metabolites in the liver and kidney) are likely to be essential and protective nutrients. Choline can be obtained from diet or produced by denovo synthesis in tissues whereas betaine can only be got from diet. Choline has a wide variety of functions, such as maintaining the structural integrity of cells, affecting signaling and transport across membranes and serving as a basic component of the neurotransmitter acetylcholine. Moreover, both choline and betaine participate in one-carbon metabolism, as major methyl-group donors3. Evidence from some humans and animals studies has demonstrated that choline insufficiency alters the structure of DNA and histones, resulting in DNA strand breaks4,5.

So far, there have been a number of epidemiologic studies exploring whether dietary consumption of choline and betaine is associated with the risk of cancer but the results are conflicting. Also, to our knowledge, no article has been found to attempt to make a summary of the results. Hence, it will be of interest to evaluate whether the consumption of choline and betaine is one of the dietary factors that are related with cancer incidence, on the basis of present epidemiologic evidence. To pool the results of relevant reports on the association and evaluate the dose-response relationship between choline and betaine consumption and the risk of cancer as well, we conducted this quantitative meta-analysis.

1Department of Oncology, The First Affiliated Hospital of Nanjing Medical University, Nanjing, China. 2Department of Pathology, The First Affiliated Hospital of Nanjing Medical University, Nanjing, China. 3Department of Environmental Genomics, Jiangsu Key Laboratory of Cancer Biomarkers, Prevention and Treatment, Cancer Center, Nanjing Medical University, Nanjing, China. 4Department of Genetic Toxicology, the Key Laboratory of Modern Toxicology of Ministry of Education, School of Public Health, Nanjing Medical University, Nanjing, China. 5Department of Oncology, Nanjing Hospital of T.C.M, Nanjing, China. 6Department of Thoracic Surgery, The First Affiliated Hospital of Nanjing Medical University, Nanjing, China. *These authors contributed equally to this work. Correspondence and requests for materials should be addressed to L.Z. (email: zhulingjun@njmu.edu.cn) or W.W. (email: wangwei-doctor@163.com)
Results

Literature search. Figure 1 tells the detailed procedures of how we searched and selected relevant articles. In brief, we retrieved 861 articles from PubMed and 825 articles from Embase, 1,686 articles in total, of which 271 articles were excluded due to duplication. Nineteen articles were left for further evaluation of the full text after screening the titles or abstracts of the remaining articles. We further eliminated 8 articles owing to reasons as listed below: no odds ratio (OR)/relative risk (RR) or 95% confidence interval (CI) reported (n = 3); data on the same population (n = 3); conference abstract lacking enough details for quality assessment (n = 2). Finally, we included 11 articles6–16 in accordance with the inclusion criteria without additional articles from the reference review. Among these articles, one paper by Ibiebele et al.7 reported two outcomes of esophageal cancer on the basis of its two subtypes: esophageal adenocarcinoma (EAC) and esophageal squamous cell carcinoma (ESCC); the article by Zhang et al.10 reported data from a two-stage case-control; another entry by Cho et al.16 provided information on males and females separately. Thus, our meta-analysis included 14 comparisons.

Study characteristics and quality assessment. Characteristics extracted from the included 11 articles were displayed in Table 1. Of these articles published between 2007 and 2015, 6 were case-control studies and 5 were cohort studies. In aggregate, we documented 14,488 cases among 513,390 participants, of which the number varied from 738 to 159,957. Seven studies were carried out in the USA, three studies in China and only one study in Australia. The range of the age of all participants was from 18 to 98 years if not considering the article by Ying et al.8 in which there was no age restrictions. Four studies included both males and females, five studies included only females, one study included only males and one study provided information on males and females separately. In these studies, the role of choline and betaine was investigated in colorectal cancer, breast cancer, esophageal cancer, lung cancer, nasopharyngeal carcinoma, epithelial ovarian cancer and renal cell cancer. To estimate dietary choline and betaine intake, all studies used the Food Frequency Questionnaire (FFQ). Categories of choline and betaine consumption were divided into fourths or fifths. All studies provided adjusted risk estimates. Supplemental Tables 1 and 2 summarized quality scores of case-control studies and cohort studies, respectively. The quality scores of all studies ranged from 6 to 9. The median score of case-control studies was 6.5 and that of cohort studies was 9. Three case-control studies and all cohort studies were of high quality.

Choline consumption and cancer risk. All 14 comparisons from 11 articles included reported the relationship between dietary consumption of choline and cancer risk. Overall, for the highest versus the lowest choline consumption, the pooled RR was 0.82 (95% CI, 0.69 to 0.96; Fig. 2) with significant heterogeneity among the 14 comparisons (P < 0.001, I² = 80.2%). According to the Begg's funnel plot (Fig. 3) and Egger’s test (P = 0.030), the publication bias existed.

Betaine consumption and cancer risk. The 11 articles with 14 comparisons also reported the relationship between dietary consumption of betaine and cancer risk. For the highest versus the lowest betaine consumption, the pooled RR was 0.86 (95% CI, 0.76 to 0.97; Fig. 4) with significant heterogeneity among the 14 comparisons (P < 0.001, I² = 65.8%). Additionally, no evidence of publication bias was observed from the Beggs funnel plot (Fig. 5) and Egger’s test (P = 0.319).

Choline plus betaine consumption and cancer risk. Only 5 comparisons from 4 articles9–11,13 presented results of dietary consumption of choline and betaine combined and cancer risk. For the highest versus the lowest choline plus betaine consumption, the pooled RR was 0.60 (95% CI, 0.40 to 0.90; Fig. 6) with significant heterogeneity among the 5 comparisons (P < 0.001, I² = 82.0%). Publication bias was neither investigated graphically (Begg’s funnel plot) nor by testing (Egger’s test) because the small number of studies limits the usefulness of these methods.

Dose-response analysis. Only 3 comparisons from 2 case-controls9,10 were eligible for exploring the dose-response relationship between dietary consumption of choline plus betaine and cancer risk. The summary
| First author year of publication (reference) | Cancer type | Country/ design | Cases/ Controls | Age(y)/sex | Follow-up (y) | Assessment of consumption/ food item number/ choline and betaine consumption calculation/nutrient database | Contrast (highest vs lowest) | Ajusted OR/RR(95%CI) (highest vs lowest) | Matched or adjusted variables |
|---------------------------------------------|-------------|----------------|----------------|------------|--------------|------------------------------------------------|---------------------------|--------------------------------|----------------------------------|
| Xu et al.11 | Breast cancer | USA, case-control (PB) | 1,508/1,556 | 20-98 F | NR | FFQ/100/ frequency × portion-size/USDA database | Quartile >247.2 vs < 122.7mg/d | 0.85(0.61–1.18) | 0.96 (0.73–1.25) | NR | Age and daily energy intake |
| Ihiebele et al. 2010 | Esophageal Cancer | Australia, case-control (PB) | 881/1,507 | 18-79 F | NR | FFQ/135/-/USDA database | Quartile 1171 vs 308mg/d | NR | EAC: 1.25(0.91 – 1.72) ESCC: 1.39(0.88–2.30) | NR | Age, gender, education, BMI 1 y previously, frequency of heartburn or acid reflux 10 y prior to diagnosis, lifetime alcohol intake, pack-years of smoking, lung cancer history, total energy intake |
| Ying et al.11 | Lung Cancer | USA, case-control (PB) | 2,821/2,923 | no age restrictions | M/F | FFQ/214/ frequency × portion-size/Chinese Food Consumption Table | Quartile (highest vs lowest) | 0.92(0.78–1.07) | 0.71 (0.61–0.84) | NR | Sex, race/ethnicity, age, pack-years, total calorie intake, family cancer history, smoking, current drinking, exposure to potential carcinogens, occupational exposure to mutlivitamin supplement, chronic rhinitis, physical activity, dietary folate intake and daily energy intake |
| Zeng et al.11 | Nasopharyngeal Carcinoma | China, case-control (PB) | 600/600 | 30-75 M/F | NR | FFQ/76/ frequency × portion-size/Chinese Food Composition Table | Quartile 285 vs 125mg/d | 0.42(0.29–0.61) | 0.50 (0.35–0.72) | 0.44 (0.30–0.64) | Age, BMI, occupation, marital status, educational level, household income, current smoking, current drinking, exposure to potential carcinogens, NSAID use, alcohol, liver disease, body mass index, number of children, number of previous pregnancies, family cancer history, total energy intake, and potential confounders |
| Zhang et al.11 | Breast cancer | China, stage-case-control (PB) | 1,438/1,438 stage 1: 438/438 stage 2: 369/369 Pooled: 807/807 | 25–70 F | NR | FFQ/81/ frequency × portion-size/Chinese Food Composition Table | stage 1: Quartile >207.46 vs < 111.44mg/d | stage 1: 0.55(0.39–0.76) | 0.39 (0.28–0.54) | stage 1: 0.43 (0.28–0.66) | Occupation, body mass index, age at menarche, live births and age at first live birth, mother/sister/ father with breast cancer, passive smoking, alcohol consumption, alcohol intake, physical activity, dietary folate intake and study stage |
| Lu et al.11 | Colorectal Cancer | China, case-control (PB) | 890/890 | 30-75 M/F | NR | FFQ/81/ frequency × portion-size/Chinese Food Composition Table | Quartile >176.40 vs < 99.60mg/d | 0.92 (0.65–1.29) | 0.91 (0.65–1.28) | Age, sex, education, income level, occupation, family history of cancer, smoking status, passive smoking, alcohol drinking, diet of physical activity, BMI, red meat, fish, beans and folate intake |

Continued
| First author year of publication (reference) | Cancer type | Country/ design | Cases/ Controls | Age(y)/sex | Follow-up(y) | Assessment of consumption/ food item number/ choline and betaine consumption calculation/nutrient database | Contour (highest vs lowest) | Adjusted OR/RR(95%CI) (highest vs lowest) | Matched or adjusted variables |
|-------------------------------------------|-------------|----------------|----------------|------------|--------------|------------------------------------------------|------------------|-----------------------------------------------|----------------------------------|
| Cho et al.15                             | Breast cancer | USA; cohort | 1,032/9,631 | 62–66 F    | 12          | FFQ/ about 130/140/ USDA database or other sources | Quintile 397 vs 567mg/d | 0.88 (0.72–1.07) | NR                                      | Smoking, height, parity and age at first birth, body mass index, age at menarche, family history of breast cancer, history of benign breast disease, oral contraceptive use, and intakes of alcohol, energy, and animal fat |
| Kostopoulos et al.16                     | Epithelial ovarian cancer | USA; cohort | 526/139,431 | 25–55 F    | 22          | FFQ/ about USDA database or other sources | Quintile NHS: 338.8 vs < 240.5 mg/d; NHSII: 367.3 vs < 267.8 mg/d | 0.98 (0.73–1.31) | NR                                      | Age at menarche and parity, duration of oral contraceptive use, tubal ligation, height, family history of breast or ovarian cancer, caffeine intake, hysterectomy, PMH, postmenopausal status, calorie intake, alcohol consumption and total folate |
| Lee et al.14                             | Colorectal Cancer | USA; cohort | 907/46,315 | 40–75 M    | 18          | FFQ/ about frequency × nutrient content/ USDA database | Quintile (highest vs lowest) | 0.97 (0.79–1.20) | NR                                      | Total energy intake, aspirin dose, pack-years of smoking before age 30, body mass index, family history of colorectal cancer, history of endoscopy, alcohol intake, and total folate |
| Cho et al.15                             | Breast cancer | USA; cohort | 3,900/70,594 | 30–55 F    | 20          | FFQ/ about 130/140/ USDA database or other sources | Quintile 396 vs 267mg/d | 1.10 (0.99–1.22) | NR                                      | Smoking status, height, parity and age at first birth, body mass index at age 18, weight change between age 18 and current, physical activity, age at menarche, family history of breast cancer, history of benign breast disease, use of post-menopausal hormones, and intakes of alcohol, energy, and folate |
| Cho et al.15                             | Renal cell cancer | USA; cohort | 225/176,985 | 40–75 M    | 22          | FFQ/ about 130/140/ USDA database or other sources | Quintile 398 vs 267mg/d; NHS: Quintile 471 vs 303.5mg/d | 0.96 (0.62–1.48) | NR                                      | Age, smoking status, body mass index, history of hypertension, history of diabetes, physical activity, fruit intake, vegetable intake, and alcohol intake in NHS and HPFS and parity in NHS |

Table 1. Characteristics of studies on choline and betaine consumption and cancer risk. Abbreviations: EAC, esophageal adenocarcinoma; ESCC, esophageal squamous cell carcinoma; FFQ, food frequency questionnaire; HB, hospital-based; HPFS, The Health Professionals Follow-up Study; PB, population-based; MB, mixed based (including both hospital-based and population-based); NASID, Non-steroidal Anti-inflammatory Drugs; NHS, The Nurses’ Health Study; NHSII, The Nurses’ Health Study II; NR, no record; PMH, postmenopausal hormone; USDA, US Department of Agriculture.

RR for an increment of 100 mg/day of choline plus betaine consumption was 0.89 (95% CI, 0.87 to 0.92) without heterogeneity (P = 0.965, I² = 0%).
Stratified analysis. For the purpose of exploring the sources of heterogeneity among the primary results, we conducted stratified analyses for choline consumption only, betaine consumption only as well as choline consumption combined with betaine consumption.
plus betaine consumption, respectively (Table 2). Significant effect of protection for cancer was revealed in hospital-based case-control studies: 0.52 (95% CI, 0.30 to 0.90) for choline consumption only, 0.60 (95% CI, 0.49 to 0.75) for betaine consumption only and 0.43 (95% CI, 0.33 to 0.55) for choline plus betaine consumption. When stratifying by the location, the RRs were 0.45 (95% CI, 0.36 to 0.55) for choline consumption only, 0.61 (95% CI, 0.44 to 0.85) for betaine consumption only and 0.52 (95% CI, 0.34 to 0.80) for choline plus betaine consumption, all producing significant reduction in cancer risk in China. In the subgroup analysis of cancer type, it was found significantly protective effect for breast cancer only in dietary consumption of choline and betaine combined (RR 0.42, 95% CI, 0.30 to 0.59). The risk for developing cancer was significantly lower in female only (RR 0.78, 95% CI, 0.61 to 0.99) for choline consumption only as well as male and female (RR 0.78, 95% CI, 0.62 to 1.00) for betaine consumption only. Besides, the protective effect for cancer was also observed in these subgroups (publication year after 2010; quality score ≥ 7 stars).

Meta-regression analysis. As shown in Supplemental Table 3, the meta-regression indicated that location (P < 0.001) and publication year (P = 0.008) but not the study design were significant sources of heterogeneity in the relationship of choline consumption only and the risk of developing cancer. The estimated between-study variance (τ²) was decreased from 0.069 to 0.004 (location) and 0.054 (publication year). Moreover, the location alone could explain 94.20% and the publication year could alone explain 21.74% of the τ². Hence, for the relationship of the consumption of choline only and cancer risk, the location was the main source of the heterogeneity. As the same time, for the relationship of betaine consumption only and cancer risk, meta-regression indicated that study design (P = 0.045), location (P = 0.016) and publication year (P = 0.006) were all significant sources of heterogeneity. The τ² was reduced from 0.031 to 0.009 (study design), 0.018 (location) and 0 (publication year). Besides, the study design alone could explain 70.97%, the location alone could explain 41.94% and the publication year could alone explain 100% of the τ². Thus, the heterogeneity across the comparisons about the relationship of betaine consumption only and cancer risk was almost entirely from publication year. On account of only 5 comparisons from 4 articles included in the relationship between consumption of choline and betaine combined and cancer risk, we did not run the meta-regression.
Sensitivity analysis. We also analyzed the sensitivity, attempting to explain the heterogeneity, examine whether varying in the criteria of inclusion had influence over the overall results and confirm the robustness of our results by omitting 1 comparison at every turn and recalculating the pooled relative risks for the remaining. Inclusion of another 3 articles that reported the hazard ratio (HR) with the corresponding 95% CI yielded similar results: a RR of 0.86 (95% CI, 0.76 to 0.97) with substantial evidence of heterogeneity (P < 0.001, I² = 80.0%) for choline consumption only and a RR of 0.87 (95% CI, 0.77 to 0.97) with substantial evidence of heterogeneity (P = 0.001, I² = 61.9%) for betaine consumption only. Under the circumstance of omitting any single comparison, the overall results did not materially alter, with ranges of the pooled RRs from 0.79 (95%CI, 0.67 to 0.94) to 0.90 (95%CI, 0.79 to 1.00) for betaine consumption only (Supplemental Figure 3) and from 0.65 (95%CI, 0.42 to 0.92) to 0.68 (95%CI, 0.42 to 1.02) for choline consumption only (Supplemental Figure 1). For consumption of choline plus betaine (Supplemental Figure 3), the range was from 0.52 (95%CI, 0.34 to 0.80) to 0.65 (95%CI, 0.42 to 1.02) and exclusion of any of the 3 comparisons from articles by Zeng et al. and Zhang et al. influenced the overall risk estimates. The pooled RRs for the remainders were 0.65 (95%CI, 0.42 to 1.02), 0.65 (95%CI, 0.42 to 1.02) and 0.65 (95%CI, 0.42 to 1.01), respectively, with no heterogeneity changing.

Discussion

Findings from our meta-analysis manifested that high choline and betaine consumption contributed to cancer prevention. Moreover, the dose-response analysis further demonstrated the protective effect from choline and betaine consumption towards cancer occurrence: an increase in choline plus betaine consumption of 100 mg/day significantly lowered the chance of developing cancer by 11%.

Table 2. Stratified analyses of dietary choline and betaine consumption and cancer risk.
Homocysteine depends on a one-carbon unit to produce methionine and then generate S-adenosylmethionine (SAM), the universal methyl donor, which contributes to methylation actions, such as histone and DNA methylation. Just like 5-methyl tetrahydrofolate, choline can offer the one-carbon unit when oxidized to betaine. Therefore, in case that this pathway of one-carbon metabolism gets disrupted, it will exert an influence over the process of DNA synthesis and repair, as well as the genetic expression regulation by means of methylation, and consequently promotes carcinogenesis. Moreover, choline metabolism and folate metabolism are closely connected, interacting at the point of the conversion from homocysteine to methionine. A few studies on animals and humans have shown that dietary choline and folate are supplementary to each other if either of them gets in a state of deprivation. In other words, diets of choline deficiency also result in the deficiency of tissue folate, which plays a vital role in DNA synthesis and one-carbon metabolism when converted into the aforementioned 5-methyl tetrahydrofolate. Furthermore, in animals studies, choline deficiency causes cancer via inducing dysfunction in mitochondria and excessive production of reactive oxygen species (ROS), which has long been considered to be one of mechanisms in promoting cancer. In addition, Ghoshal et al. have reported that in experimental animals, a 0.8% supplemented dietary consumption of choline aids in the action of cancer prevention in complete.

The significant protective effect of choline and betaine consumption against cancer was weaker in cohort studies in the subgroup analysis of study design. The discrepant results could be explained by the retrospective nature of case-control studies which were more inclined to have greater recall and selection biases and in which, cancer cases were more likely to change their eating habits and dietary patterns because of information of choline intake and cancer consumption was gathered in their post-cancer diagnosis life. Besides, we found results from the subgroup analysis of the location which studies conducted in suggest that people settling in China were greatly protected from the risk of developing cancer (a 55% compared with 48% risk reduction in for choline consumption only, a 39% compared with 14% risk reduction for betaine consumption only and a 48% compared with 40% risk reduction for choline plus betaine consumption) and we could not find the analogous results in studies conducted in non-China. These inconsistent findings may have been attributed to differences in the compositions of the diets, diverse eating habits, dissimilar susceptibility to cancer of Chinese and non-Chinese as well as, to some degree, differences stemming from levels of scientific researches in different regions. Additionally, in the separate analysis of cancer type, dietary choline and betaine separately did not show the protective effect for breast cancer while consumption of choline plus betaine could reduce the risk of cancer incidence by 58%. Such results may be due to not enough studies on the relationship between dietary choline and betaine consumption and cancer risk to be summarized to obtain more reliable results. Different results also appeared when stratified by publication year and quality score of the study. The former could be explained by differences in the measures of scientific researches in different eras, and as to the latter, the relationship may have been attenuated by poor methodological quality of studies.

To our knowledge, meta-analysis is a crucial method for revealing trends that might not be evident in a single study. One of strengths of this meta-analysis is that the number of total subjects (513,390) was substantial, which significantly increased the statistical power. Moreover, a significant dose-response relation was observed between choline plus betaine intake and cancer risk, further strengthening the relationship. We also conducted sensitivity analyses and the combined estimates were robust except the one for choline plus betaine consumption and cancer risk. The reason for this phenomenon may be interpreted by the very inconsistent ranges of exposure in the studies by Zeng et al., Zhang et al., Liu et al. and Kotsopoulos et al. Several limitations exist in our meta-analysis, though. First, in observational studies, the possibility that residual confounders may account for the protective effect of choline and betaine could not be ruled out. Nevertheless, all studies except the one by Xu et al. included in this meta-analysis have adjusted for a wide spectrum of potential confounders, like age, gender, total energy intake, folate consumption, smoking, alcohol drinking and so on. Second, the distribution of the highest and lowest choline and betaine consumption is not uniform among studies, which may conduces to the heterogeneity in the pooled analysis and conclusions limited. Third, this present meta-analysis contains a small number of observational studies and only 2 case-control studies containing 3 comparisons did not fail to meet the applied conditions of the dose-response meta-analysis. Therefore, more observational studies with standardized choline and betaine consumption collection strategies are needed to answer the two questions more completely. Fourth, a problem with dietary assessments in case-control studies is that dietary patterns and habits may have changed after the diagnosis of cancer because of symptoms related to the cancer. Fifth, many studies only provided the results without showing detailed calculation methods or the raw data (e.g., the number of case and non-case or person-year at different ranges of consumption) and we contacted the corresponding authors for the data, but unfortunately, we failed to get any reply. So the investigators of all the published studies are encouraged to share their raw data. Sixth, substantial heterogeneity was explored among the comparisons in our meta-analysis, which could be mostly explained by study design, location and publication year, and the meta-regression analysis. Finally, publication bias appeared when examining the relationship between consumption of choline only and cancer risk and the tendency of small studies with null results not being published could be the fairly reasonable explanation.

Cancer is a large burden worldwide. The relationship between dietary choline and betaine consumption and the risk of cancer remains controversial. The results of this meta-analysis indicate that a 100 mg/d increment in consumption of choline plus betaine decreases cancer occurrence by 11%, which means that the above mentioned approximately 14.1 million new cancer cases in 2012 worldwide could fall to about 12.5 million. Besides, choline must be consumed through the diet to play its role of maintaining health of humans, though choline can also be synthetised by the liver. Dietary sources of choline are mainly eggs, beef, pork, liver, soybean, and wheat germ, while we can obtain betaine mainly from wheat bran, wheat germ, and spinach. However, if the recommended dietary adequate intake (550 mg/day for men and 425 mg/day for women) for choline is taken as a standard, there are only a minority of people meeting it. Some studies have reported that 20–25% of Americans are not up to that...
‘standard’. For example, in the Framingham Heart Study33, the Atherosclerosis Risk In Communities study34,35 and the Nurse’s Health Study46, the amount of dietary choline consumption is <203 mg/d, <217 mg/d and <293 mg/d, respectively. Choline and betaine intake may be increased by well-described dietary changes such as appropriate increases in consumption of the aforementioned foods.

To conclude, our meta-analysis manifests that choline and betaine consumption possesses the ability to lower cancer risk. However, these results should be considered with caution on account of the considerable heterogeneity, the potential biases and confounding factors. Further studies well designed as large prospective studies and placebo-controlled intervention trials on choline and betaine supplementation are warranted to ascertain the results and establish the potential dose-response relationship. For better and comprehensive understanding of the relationship between dietary consumption of choline and betaine and cancer risk, future studies are expected to use standardized collection strategy for choline and betaine consumption. Furthermore, it also requires further studies to elucidate the underlying mechanisms responsible for the relationship.

Methods

Data sources and searches. We retrieved epidemiologic studies examining the relationship between dietary choline and betaine intake and cancer risk from two databases (PubMed and EMBASE), from their inception to March 2016, using the following key words: (dietary OR consumption OR intake OR food) AND (choline OR betaine) AND (cancer OR tumor OR carcinoma OR neoplasm). The search was limited to no language restriction. In addition, the references of identified publications were scrutinised for further potentially relevant articles. This present meta-analysis was designed, performed and reported on the basis of the epidemiology guidelines for meta-analysis of observational studies37.

Study selection criteria. Published articles were included if they were in accordance with the inclusive criteria: 1) the study design was case-control or cohort; 2) the exposure of interest was dietary consumption of choline or betaine or choline plus betaine; 3) the outcome was cancer occurrence; 4) reported the relative risk (RR) or odds ratio (OR) with its 95%CI for the relationship between the consumption of choline or betaine or choline plus betaine and cancer risk adjusted at least for age. In case that more than one published paper reported data based on the same population, we selected the most recent or most informative one. Two of the authors independently identified and reviewed each relevant article, and discrepancies in study eligibility were discussed until an agreement of opinions was reached.

Data extraction and quality assessment. The relevant information was extracted by 2 of the authors independently. The extracted information for each study was as follows: the last name of the first author, year of publication, type of cancer, design of the study, the location of the study conducted in, age, gender, sample sizes, the assessment and comparison method for the consumption of choline and betaine, RRs or ORs and the corresponding 95% CIs (the highest versus the lowest consumption of choline only or betaine only or choline plus betaine) from the most fully adjusted model and adjustment for confounders in a multivariate analysis.

In order to evaluate the quality of the study, a 9-star system based on the Newcastle-Ottawa Scale38 which contains 3 aspects (the selection of study populations, the comparability of the populations and ascertainment of exposure or the outcomes of interest for case-control or cohort studies) was adopted to judge on the study. Additionally, if a study with data analysis using the residual method39 could achieve an additional star. In other words, the maximum score was 10 stars and when a study was assessed with ≥7 stars, it would be regarded as a high-quality study.

Statistical analysis. Adopting random effect models, which takes into account both within-study and between-study variations40, we estimated the pooled RRs with 95% CIs by summarizing the risk estimates of each study. Heterogeneity among studies was evaluated using the Cochran’s Q test and $I^2$ (inconsistency index) statistic41. A value of $I^2$ greater than 50% indicates severe heterogeneity and the value less than 25% suggesting no significant heterogeneity42. Based on the location, study design, cancer type, publication year, sex and quality score of study, we carried out stratified analyses to explore possible heterogeneity and test the robustness of the relationship. In the meta-regression model, we examined the variables of study design, location, year of publication, type of cancer, design of the study, the location of the study conducted in, age, gender, sample sizes, the assessment and comparison method for the consumption of choline and betaine, RRs or ORs and the corresponding 95% CIs (the highest versus the lowest consumption of choline only or betaine only or choline plus betaine) from the most fully adjusted model and adjustment for confounders in a multivariate analysis.

For the dose-response analysis, the method of generalised least square for trend estimation put forward by Greenland and Longnecker43 and Orsini et al.44 was adopted to estimate the trend from the correlated log RR estimates across categories of choline plus betaine intake. It is necessary for the method that the studies provided data of distribution of cases and non-cases or person-years and the RR or OR with its 95% CI known for at least 3 quantitative exposure categories as well as the median or mean values of choline plus betaine in each category. When studies merely provided the consumption data by ranges, we considered the average of the lower and upper bound as estimation of the midpoint of each category. Supposing that the two ends of the categories were both open ended, we regarded the length of the open interval to be the same as that of the closest interval for the highest category and set the lower boundary to zero for the lowest.

We also adopted Begg’s funnel plots and Egger’s linear regression test45,46 to evaluate whether publication bias existed. All the statistical analyses involved were performed with STATA software (version 13.1; StataCorp, College Station, Texas, USA). $P$ values were two-sided and $P < 0.05$ was considered statistically significant.

References

1. Torre, L. A. et al. Global cancer statistics, 2012. CA Cancer J Clin 65, 87–108 (2015).
2. Thun, M. J., DeLancey, J. O., Center, M. M., Jemal, A. & Ward, E. M. The global burden of cancer: priorities for prevention. Carcinogenesis 31, 100–110 (2010).
3. Zeisel, S. H. & Blusztajn, J. K. Choline and human nutrition. Am. Rev. Nutr. 14, 269–296 (1994).
4. Da, C. K., Niculescu, M. D., Craciunescu, C. N., Fischer, L. M. & Zeisel, S. H. Choline deficiency increases lymphocyte apoptosis and DNA damage in humans. Am. J. Clin. Nutr. 84, 88–94 (2006).
5. James, S. J. & Yin, L. Diet-induced DNA damage and altered nucleotide metabolism in lymphocytes from methyl-donor-deficient rats. Carcinogenesis 10, 1209–1214 (1989).
6. Xu, X. et al. High intakes of choline and betaine reduce breast cancer mortality in a population-based study. The FASEB Journal 23, 4022–4028 (2009).
7. Ibiebele, T. L. et al. High Intake of Folate from Food Sources Is Associated with Reduced Risk of Esophageal Cancer in an Australian Population. J. Nutr. 141, 274–283 (2011).
8. Ying, J. et al. Associations between dietary intake of choline and betaine and lung cancer risk. Plos One 8, e54561 (2013).
9. Zeng, F. et al. Choline and betaine intakes are associated with reduced risk of nasopharyngeal carcinoma in adults: a case–control study. Br J Cancer 110, 808–816 (2013).
10. Zhang, C. et al. Choline and betaine intake is inversely associated with breast cancer risk: A two-stage case-control study in China. Cancer Sci 104, 250–258 (2013).
11. Lu, M. et al. Choline and Betaine Intake and Colorectal Cancer Risk in Chinese Population: A Case-Control Study. Plos One 10, e118661 (2015).
12. Cho, E., Holmes, M., Hankinson, S. E. & Willett, W. C. Nutrients Involved in One-Carbon Metabolism and Risk of Breast Cancer among Premenopausal Women. Cancer Epidemiol Biomar. 16, 2787–2790 (2007).
13. Kotsopoulos, J., Hankinson, S. E. & Tworoger, S. S. Dietary betaine and choline intake are not associated with risk of epithelial ovarian cancer. Eur J Clin. Nutr. 64, 111–114 (2010).
14. Lee, J. E. et al. Choline and Betaine Intake and the Risk of Colorectal Cancer in Men. Cancer Epidemiol Biomar. 19, 884–887 (2010).
15. Cho, E., Holmes, M. D., Hankinson, S. E. & Willett, W. C. Choline and betaine intake and risk of breast cancer among postmenopausal women. Br J Cancer 102, 489–494 (2010).
16. Cho, E., Giovannucci, E. L. & Joh, H. Nutrients related to one-carbon metabolism and risk of renal cell cancer. Cancer Cause Control 24, 373–382 (2013).
17. Richman, E. L. et al. Choline intake and risk of lethal prostate cancer: incidence and survival. Am. J. Clin. Nutr. 96, 855–863 (2012).
18. Liu, J. J. et al. One-carbon metabolism factors and endometrial cancer risk. Br J Cancer 108, 183–187 (2013).
19. Huang, J. Y. et al. Dietary Intake of One-Carbon Metabolism-Related Nutrients and Pancreatic Cancer Risk: The Singapore Chinese Health Study. Cancer Epidemiol Biomarkers Prev. 25, 417–424 (2016).
20. Mentch, S. J. & Locasale, J. W. One-carbon metabolism and epigenetics: understanding the specificity. Ann N Y Acad Sci 1363, 91–98 (2016).
21. Horne, D. W., Cook, R. J. & Wagner, C. Effect of dietary methyl group deficiency on folate metabolism in rats. J. Nutr. 119, 618–621 (1989).
22. Selhub, J., Seyoum, E., Pomfret, E. A. & Zeisel, S. H. Effects of choline deficiency and methotrexate treatment upon liver folate content and distribution. Cancer Res 51, 16–21 (1991).
23. Kim, Y. I. et al. Severe folate deficiency causes secondary depletion of choline and phosphocholine in rat liver. J. Nutr. 124, 2197–2203 (1994).
24. Guo, W. X. et al. Mitochondrial dysfunction in choline deficiency-induced apoptosis in cultured rat hepatocytes. Free Radic Biol Med 39, 641–650 (2005).
25. Banni, S. et al. Free radicals and lipid peroxidation in liver of rats kept on a diet devoid of choline. Free Radic Res Commun 7, 233–240 (1989).
26. Vafa, O. et al. c-Myc can induce DNA damage, increase reactive oxygen species, and mitigate p53 function: a mechanism for oncogene-induced genetic instability. Mol. Cell 9, 1031–1044 (2002).
27. Gao, P. et al. HIF-dependent antitumorigenic effect of antioxidants in vivo. Cancer Cell 12, 230–238 (2007).
28. Ghoshal, A. K. & Farber, E. The induction of liver cancer by dietary deficiency of choline and methionine without added carcinogens. Carcinogenesis 5, 1367–1370 (1984).
29. Zeisel, S. H. Choline: critical role during fetal development and dietary requirements in adults. Am. Rev. Nutr. 26, 229–250 (2006).
30. Zeisel, S. H., Mar, M. H., Howe, J. C. & Holden, J. M. Concentrations of choline-containing compounds and betaine in common foods. J. Nutr. 133, 1302–1307 (2003).
31. Sakamoto, A., Nishimura, Y., Ono, H. & Sakura, N. Betaine and homocysteine concentrations in foods. Pediatr. Int. 44, 409–413 (2002).
32. Institute of Medicine (US) Standing Committee on the Scientific Evaluation of Dietary Reference Intakes and its Panel on Folate, Other B Vitamins, and Choline. Dietary Reference Intakes for Thiamin, Riboflavin, Niacin, Vitamin B6, Folate, Vitamin B12, Pantothenic Acid, Biotin, and Choline. 12 (National Academies Press (US), Washington (DC) 1998).
Acknowledgements
This study was partially supported by the National Natural Science Foundation of China (NSFC 81472634) and
Jiangsu Province Clinical Science and Technology Projects (Clinical Research Center, BL2012008).

Author Contributions
L.Z. and W.W. conceived and designed the study; S.S., X.L. and A.R. performed the experiments; M.D., H.D.
and Y.S. analyzed the data; S.S., A.R. and X.L. contributed analysis tools; S.S., A.R. and H.D. critically revised the
manuscript; S.S. wrote the paper. All authors reviewed the manuscript.

Additional Information
Supplementary information accompanies this paper at http://www.nature.com/srep
Competing financial interests: The authors declare no competing financial interests.

How to cite this article: Sun, S. et al. Choline and betaine consumption lowers cancer risk: a meta-analysis of
epidemiologic studies. Sci. Rep. 6, 35547; doi: 10.1038/srep35547 (2016).

This work is licensed under a Creative Commons Attribution 4.0 International License. The images
or other third party material in this article are included in the article’s Creative Commons license,
unless indicated otherwise in the credit line; if the material is not included under the Creative Commons license,
users will need to obtain permission from the license holder to reproduce the material. To view a copy of this
license, visit http://creativecommons.org/licenses/by/4.0/

© The Author(s) 2016