# BMJ Open Coffee consumption and risk of prostate cancer: a systematic review and metaanalysis

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#### **ABSTRACT**

Objectives To conduct a systematic review with metaanalysis of cohort studies to evaluate the association of coffee consumption with the risk of prostate cancer. Data sources PubMed, Web of Science and Embase were searched for eligible studies up to September 2020. Study selection Cohort studies were included. Data extraction and synthesis Two researchers independently reviewed the studies and extracted the data. Data synthesis was performed via systematic review and meta-analysis of eligible cohort studies. Meta-analysis was performed with the "metan" and "glst" commands in Stata 14 0

Main outcomes and measures Prostate cancer was the main outcome. It was classified as localised prostate cancer which included localised or non-aggressive cancers; advanced prostate cancer which included advanced or aggressive cancers; or fatal prostate cancer which included fatal/lethal cancers or prostate cancerspecific deaths.

**Results** Sixteen prospective cohort studies were finally included, with 57 732 cases of prostate cancer and 1 081 586 total cohort members. Higher coffee consumption was significantly associated with a lower risk of prostate cancer. Compared with the lowest category of coffee consumption, the pooled relative risk (RR) was 0.91  $(95\% \text{ Cl } 0.84 \text{ to } 0.98), 1^2 = 53.2\%)$  for the highest category of coffee consumption. There was a significant linear trend for the association (p=0.006 for linear trend), with a pooled RR of 0.988 (95% CI 0.981 to 0.995) for each increment of one cup of coffee per day. For localised, advanced and fatal prostate cancer, the pooled RRs were 0.93 (95% CI 0.87 to 0.99), 0.88 (95% CI 0.71 to 1.09) and 0.84 (95% CI 0.66 to 1.08), respectively. No evidence of publication bias was indicated in this meta-analysis.

**Conclusions** This study suggests that a higher intake of coffee may be associated with a lower risk of prostate cancer.

### INTRODUCTION

Prostate cancer is the second most frequently diagnosed cancer and the sixth leading cause of cancer death in men. There were 1276000 new cancer cases and 359000 cancer deaths in 2018. It is estimated that nearly threequarters of prostate cancer cases occur in developed countries. Since the 1970s, the incidence of prostate cancer has also

### Strengths and limitations of this study

- Risk of selection and recall bias may be minimised due to the inclusion of prospective cohort studies.
- Large sample size ensures adequate statistical power to detect even a small effect of interest.
- Uncontrolled/residual confounding may distort the association between coffee consumption and prostate cancer.
- Misclassification of coffee consumption may occur due to the self-reported nature of the exposure.
- Significant heterogeneity among study results may come from various sources.

increased rapidly in some Asian countries such as China, Singapore and Japan, where the incidence has always been much lower than in some Western countries. 12 Therefore, primary prevention of prostate cancer is a significant public health problem worldwide.

Coffee is one of the most popular beverages. Since its popularity continues to increase worldwide, even a small effect on individual health may exert a substantial public health impact. Coffee is known to be a major source of dietary caffeine, cafestol and antioxidants in industrialised nations.<sup>3</sup> Its various constituents such as caffeine, caffeic acid and chlorogenic acid can potentially impact the development of cancer through multiple carcinogenesis pathways. 4 5 Inverse associations were observed between coffee consumption and the risk of cancer in sites such as the liver, colorectum and breast.<sup>6</sup> However, previous studies have reported inconsistent results on the association of coffee consumption with the risk of prostate cancer. Although earlier cohort studies did not detect an association, 7-15 more recent studies conducted in major Western countries such as the USA, Sweden and the UK reported that coffee consumption was associated with a lower risk of localised and advanced prostate cancer. 16-20 In Japan, a country with increasing popularity of coffee, a cohort study also found a



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significant inverse association between coffee consumption and the risk of prostate cancer.<sup>21</sup>

Previous meta-analyses of cohort studies up to 2015 reported a significant positive association for coffee consumption on total prostate cancer risk, with highly variable results in different subgroups. 22 23 Since then, five cohort studies have explored the association but still reported inconsistent results.<sup>24–28</sup> It was hypothesised that higher coffee consumption was associated with an increased risk of prostate cancer. Thus, the objective of this updated meta-analysis was to explore and evaluate the association of coffee intake with the risk of prostate cancer in adult men, and to direct the future primary prevention strategy on prostate cancer.

#### **METHODS**

This systematic review was conducted and reported in adherence to the PRISMA and MOOSE guidelines<sup>29</sup>; the corresponding checklists are shown in online supplemental table 1 and online supplemental file 1. Two researchers (YZ and ZT) independently conducted the literature search, study selection, data extraction and study quality assessment. Any discrepancies were resolved by discussion, but whenever consensus could not be reached between the two reviewers, a third reviewer (KW) acted as arbitrator.

#### Patient and public involvement

This is a meta-analysis based on study-level data and no individual-level data were involved in the study or in defining the research question or outcome measures.

#### **Inclusion criteria**

The eligibility criteria of the studies were as follows: (1) The study should use a longitudinal cohort design or case-control design nested within a cohort study. (2) The study should present information on coffee consumption as the exposure of interest. Coffee consumption was ascertained by self-reported dietary records or food diaries on the intake levels (highest intake category vs lowest intake category) or frequency measures (eg, per unit/cups/ mL per day/week). Since the intake levels were classified and defined differently in each study, the absolute coffee consumption in the highest and lowest intake categories varied across the included studies. (3) The study should report prostate cancer as the outcome of interest. Prostate cancer was defined by clinical diagnosis, physician diagnosis, medical records, self-reports or data linkage to a registry system such as a cancer registry. Based on definitions in each original study, the prostate cancer categories were classified as follows: (a) localised prostate cancer, which included localised or non-aggressive cancers; (b) advanced prostate cancer, which included advanced or aggressive cancers; and (c) fatal prostate cancer, which included fatal/lethal cancers or prostate cancer-specific deaths. (4) The study should provide relative risk (RR), HR, risk ratio, rate ratio or OR estimates with confidence

intervals (CIs) or standard errors for the association of coffee consumption with the risk of prostate cancer. If multiple estimates were provided, priority was given to the multivariable-adjusted risk estimates. If more than one study was conducted in the same population, the earlier reports or reports with less applicable information were excluded.

#### Literature search

A literature search was performed using PubMed, Web of Science and Embase up to September 2020 with the following keywords: coffee and prostate and (cancer or carcinoma or neoplasm or tumour). The full search strategy is shown in online supplemental file 2 The reference lists of relevant publications were also manually searched for identification of additional eligible studies. No language limitation was imposed.

When data or information in the publication were insufficient, we attempted to contact the corresponding authors of the original study to request the relevant data. Then two authors (Russnes and Nilsson) provided us with the relevant information about the person-years of follow-up for specific categories of coffee intake to facilitate the dose-response analyses. <sup>15 20</sup> Of note, we finally did not include the study by Russnes et al<sup>20</sup> in the current meta-analysis because the study population is the same as another included cohort study by Wilson et al<sup>18</sup> which reported more applicable information.

#### **Data extraction**

We extracted the following information from each eligible study: first author's name, year of publication, study country, follow-up time, number of participants in the cohort, number of prostate cancer cases, assessment of coffee consumption, primary study outcome, definitions and categories of coffee consumption, RRs and 95% CIs for all prostate cancer outcomes associated with coffee consumption, and the potential confounders considered or adjusted in the analysis.

### Study quality assessment

The 9-star Newcastle-Ottawa Scale tool was used to assess study quality.<sup>30</sup> The quality of each cohort study was judged on three broad categories—namely, selection of the study population, comparability of groups and ascertainment of either the exposure or outcome of interest.

#### Statistical analysis

In the meta-analysis, the RR estimate was used to measure the association between coffee consumption and the risk of prostate cancer in this meta-analysis. We pooled the study-specific RR estimates for the highest versus the lowest category of coffee consumption. A fixed effects model was used to pool the study-specific estimates; whenever significant heterogeneity was detected, the random effect model was used to address the heterogeneity across studies.<sup>31</sup> Subgroup analyses were conducted stratified by study location, prostate cancer stage and potential confounder adjustments including a history of prostate-specific antigen (PSA) testing, a family history of prostate cancer, total energy intake, cigarette smoking, alcohol consumption, physical activity, body mass index (BMI) or history of diabetes. Since PSA testing was generally introduced after 1986,<sup>32</sup> studies with follow-up periods that ended before 1986 were classified in the PSA-adjusted group. To explore the influence of each study on the pooled results, sensitivity analyses were also performed by excluding one study at a time and then repeating the meta-analysed approach.

We further examined the potential dose-response relationship between coffee consumption and the risk of prostate cancer. When the mean coffee intakes in each category were not reported, the midpoint values in each category were used instead; when the upper boundary of the highest intake category was not presented, we calculated the midpoint value assuming that the highest category had the same magnitude of intake as the preceding category.<sup>33 34</sup> The pooled RR for each increment of one cup of coffee per day was estimated using the method proposed by Orsini and Greenland.<sup>35</sup> We examined a potential non-linear relation between coffee consumption and prostate cancer risk by modelling coffee consumption using restricted cubic splines for non-linear trends with 4 knots at fixed percentiles (5%, 35%, 65% and 95%) of the distribution.<sup>36</sup> Non-linearity of the association was explored by testing the null hypothesis that the coefficients of the second and third splines were equal to zero.

We assessed the heterogeneity by using the Q and the  $I^2$  statistic. A p value <0.10 or an  $I^2$  >50% suggest that statistical heterogeneity may exist.<sup>37</sup> Small study effects such as publication bias were evaluated by funnel plots, as well as Begg's test and Egger's test. 38 39 Meta-analysis was conducted using the "metan" and "glst" commands in Stata version 14.0 (StataCorp, College Station, Texas, USA). Two-sided p values <0.05 were considered statistically significant in the meta-analysis.

#### **RESULTS**

#### Literature search

We identified 497 records after searching the three databases. After 217 duplicate records were removed, 280 records remained for screening of titles and abstracts, and after screening the titles and abstracts, 254 irrelevant records were excluded. Following a further fulltext review of the 26 remaining studies, 10 studies were excluded because of no useful risk estimates or 95% CIs; two studies were excluded as newer data or more informative data were available. Fourteen studies were obtained from full-text screening and a further two studies were identified by checking the reference lists of retrieved articles. Thus, 16 studies were included in the final analysis, 11 13-19 21 24-28 of which 15 reported on the risk of prostate cancer associated with the highest versus the lowest coffee consumption 9-11 13-19 21 24-27; 13 studies reported the risk associated with an increase of one cup

of coffee per day or provided sufficient data to estimate the dose–response risk 9 13–19 21 24 25 27 28 (figure 1).

#### Study characteristics and quality assessment

The characteristics of the eligible cohort studies are shown in online supplemental file 3. The included studies were conducted in North America (n=7), Europe (n=7) and Japan (n=2). There was a total of 1081586 men in the 16 cohort studies, of whom 57732 developed prostate cancer. To measure coffee consumption, 11 studies used food-frequency questionnaires and five used a selfadministered dietary questionnaire. Most studies considered or adjusted for the most potential confounders in the analysis, such as age at baseline, family history of prostate cancer, race, cigarette smoking, alcohol drinking, total energy intake, BMI and physical activity. The results of study quality assessment are presented in online supplemental file 4. The total scores for each cohort study ranged from 6 to 9. Fourteen studies awarded a total score of ≥7, which were considered as relatively highquality studies with a low risk of bias.

#### Overall analyses and dose-response analyses

The reported RRs for the original cohort studies ranged from 0.47 (95% CI 0.25 to 0.87) in the study by Pounis et al to 1.42 (95% CI 0.77 to 2.61) in the study by Ellison et al (figure 2). Compared with the lowest coffee intake category, there was a 9% reduction in the risk of prostate cancer for the highest category (RR=0.91; 95% CI 0.84 to 0.98). Statistically significant heterogeneity was detected across the studies (p=0.008, I<sup>2</sup>=53.2%). In dose-response analyses, we found evidence of a linear inverse association between coffee consumption and prostate cancer risk (p=0.006 for linear trend) (figure 3). The pooled RR of prostate cancer was 0.988 (95% CI 0.981 to 0.995) for an increase of one cup of coffee per day. No evidence of a non-linear relationship was observed between coffee consumption and risk of prostate cancer (p=0.193 for non-linearity). Moreover, there was no indication of small study effects such as publication bias from the results of the Egger's test (p=0.409), Begg's test (p=0.843) as well as the funnel plot. The funnel plot and Egger's publication bias plot are shown in online supplemental file 5.

#### **Subgroup and sensitivity analyses**

As shown in table 1, compared with the lowest coffee intake category there was a 7% reduction in risk for the highest intake category (RR=0.93; 95% CI 0.87 to 0.99) for localised prostate cancer. For advanced and fatal prostate cancer, the corresponding pooled RRs were 0.88 (95% CI 0.71 to 1.09) and 0.84 (95% CI 0.66 to 1.08) (figure 4). When stratified by study location, the pooled RRs were 0.96 (95% CI 0.90 to 1.03), 0.85 (95% CI 0.74 to 0.98) and 0.85 (95% CI 0.48 to 1.51) for studies conducted in North America (six in the USA and one in Canada), European countries and Japan. Furthermore, significant inverse associations were observed in all of the confounder adjusted subgroups.

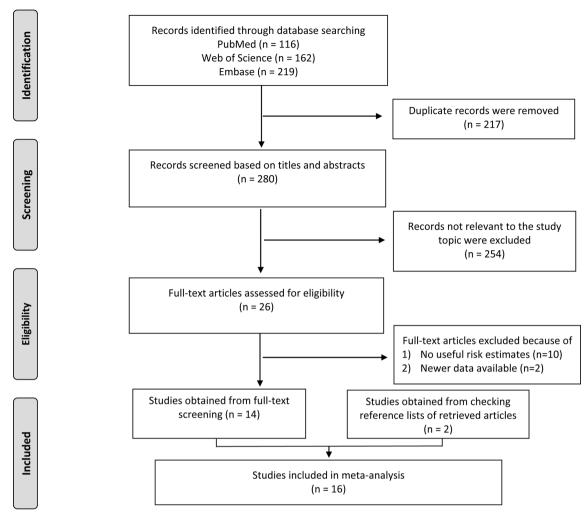


Figure 1 Flow diagram of study selection in the meta-analysis.

In sensitivity analyses, we sequentially excluded one study at a time and recalculated the pooled RRs of the remaining studies. The pooled RRs did not change substantially, ranging from 0.89 (95% CI 0.82 to 0.97) to 0.93 (95% CI 0.86 to 1.00) after omission of the studies by Hashibe *et al* and Terdal *et al*, respectively.

# DISCUSSION

#### **Summary of the findings**

In this meta-analysis, higher coffee consumption was significantly associated with a reduced risk of prostate cancer in men. In the dose–response analysis, a reduction in the risk of prostate cancer of nearly 1% was observed for each increment of one cup of coffee per day. The combined estimate for prostate cancer was robust across subgroup and sensitivity analyses.

#### **Comparison with other studies**

The previous meta-analysis detected a statistically significant positive association between coffee consumption and prostate cancer risk (RR=1.16; 95% CI 1.01 to 1.33). However, this observed effect was confined to case—control studies (RR=1.21; 95% CI 1.03 to 1.43), with no significant

association in cohort studies (RR=1.06; 95% CI 0.83 to 1.35) when stratified by study design.<sup>22</sup> Considering the case-control design, patients with prostate cancer might differentially recall their past coffee consumption habits compared with healthy controls, which might generally lead to biased estimates. This potential recall bias could generate a spurious positive association between coffee consumption and prostate cancer risk. Additionally, selection bias, which can occur in case-control studies, may distort the association between coffee consumption and prostate cancer risk. In another meta-analysis of cohort studies with 539577 participants and 34105 prostate cancer cases, the pooled RR for the highest category of coffee intake was 0.90 (95% CI 0.85 to 0.95) for total prostate cancer compared with the lowest intake category. In this updated meta-analysis of 1081586 cohort members and 57732 incident cases, the overall result was similar to the previous one. However, for subgroups of localised, advanced and fatal prostate cancers, the strength of associations tended to be weaker compared with the previous study.

### Possible biological mechanisms

It is biologically plausible that coffee may reduce the risk of prostate cancer in men. Coffee improves

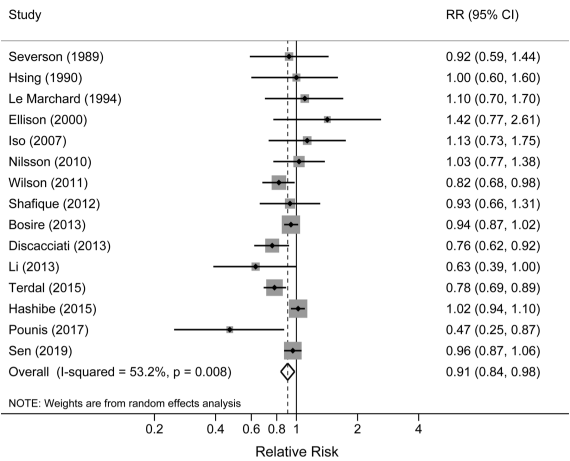


Figure 2 Forest plot for the association between coffee consumption and prostate cancer risk.

glucose metabolism, decreases concentrations of plasma insulin and insulin-like growth factor-1, has anti-inflammatory and antioxidant effects, and affects sex hormone levels, all of which may play roles in the initiation, development and progression of prostate cancer.<sup>3</sup> 18 20 40 Coffee is also a major source of chlorogenic acids; intake of quinides, the degradation products of chlorogenic acids, has been observed to increase insulin sensitivity and lower blood glucose levels.<sup>3</sup> Moreover, coffee intake may be associated

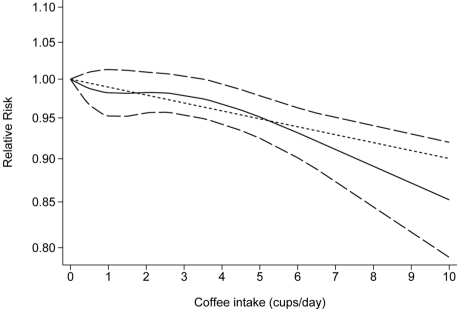


Figure 3 Dose–response relationship of coffee consumption with prostate cancer risk.

**Table 1** Summary risk estimates and corresponding 95% CIs for prostate cancer associated with the highest versus the lowest coffee consumption

lowest coffee consumption	N. 6		250/ 84	12 (2.1)	
	No of studies	Summary RR	95% <b>CI</b>	l² (%)	P value*
Overall	15	0.91	(0.84 to 0.98)	53.2	0.008
Prostate cancer category †					
Localised	6	0.93	(0.87 to 0.99)	37.0	0.159
Advanced	8	0.88	(0.71 to 1.09)	52.7	0.039
Fatal	6	0.84	(0.66 to 1.08)	46.3	0.097
Study location					
North America	7	0.96	(0.90 to 1.03)	17.7	0.295
Europe	6	0.85	(0.74 to 0.98)	63.3	0.018
Japan	2	0.85	(0.48 to 1.51)	68.5	0.075
NOS score					
6	2	1.20	(0.84 to 1.72)	0	0.507
7	3	1.02	(0.78 to 1.32)	0	0.810
8 or 9	10	0.88	(0.81 to 0.96)	66.2	0.002
Adjustment for confounders					
PSA testing ‡					
Yes	6	0.86	(0.77 to 0.96)	31.8	0.197
No	9	0.94	(0.84 to 1.06)	60.5	0.009
Family history of prostate cancer					
Yes	4	0.83	(0.72 to 0.96)	57.8	0.068
No	11	0.95	(0.85 to 1.05)	50.8	0.026
Total energy intake					
Yes	6	0.85	(0.76 to 0.96)	61.1	0.025
No	9	0.97	(0.85 to 1.09)	47.7	0.053
Smoking status					
Yes	10	0.86	(0.79 to 0.94)	52.0	0.027
No	5	1.03	(0.95 to 1.11)	0	0.805
Alcohol consumption					
Yes	6	0.87	(0.84 to 0.98)	49.2	0.008
No	9	0.93	(0.84 to 1.03)	57.2	0.017
Physical activity					
Yes	7	0.87	(0.79 to 0.95)	58.4	0.025
No	8	1.00	(0.90 to 1.12)	10.5	0.348
ВМІ					
Yes	9	0.86	(0.78 to 0.94)	56.9	0.017
No	6	1.03	(0.95 to 1.11)	0	0.897
Diabetes					
Yes	5	0.87	(0.84 to 0.98)	64.9	0.022
No	10	0.97	(0.86 to 1.10)	22.8	0.233

<sup>\*</sup>P value for heterogeneity within each subgroup.

BMI, body mass index; NOS, Newcastle-Ottawa Scale; PSA, prostate-specific antigen; RR, relative risk.

<sup>†</sup>Based on definition in each original study, the prostate cancer categories were classified as follows: (a) localised prostate cancer which included localised or non-aggressive cancers, (b) advanced prostate cancer which included advanced or aggressive cancers, (c) fatal prostate cancer which included fatal/lethal cancers or prostate cancer-specific deaths. ‡Since PSA testing was generally introduced after 1986, studies with follow-up periods that ended before 1986 were classified in the PSA-adjusted group.

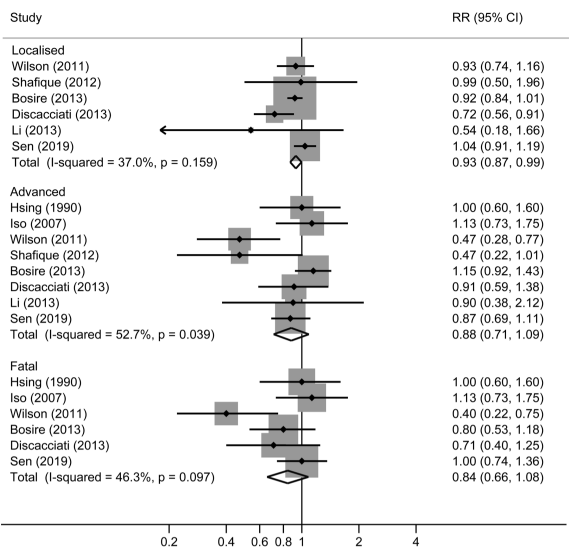


Figure 4 Forest plot for the association between coffee consumption and risk of prostate cancer stratified by cancer stages.

with increased levels of adiponectin plasma, 41 42 which may act as an endogenous insulin sensitiser. 43 Higher adiponectin levels in plasma are supposed to relate to lower concentrations of plasma insulin. 43 In two prospective studies, insulin levels were observed to be directly associated with prostate cancer-specific mortality. 44 45

Coffee is a major contributor of dietary antioxidants such as caffeic acid and chlorogenic acid. <sup>20</sup> A prospective cohort study from the USA found that dietary antioxidants from coffee (eg, caffeic acid) were inversely associated with the risk of total, advanced and lethal prostate cancer.<sup>20</sup> It was suggested that antioxidants protect cells from damage caused by oxidative stress and inflammation, which may further lead to neoplastic transformation in the prostate. 46 Additionally, dietary antioxidants may inhibit prostate cancer progression through suppression of oxidative stress, which might play a critical role during the progression of prostate cancer. 46 Coffee intake was indicated to be related to increased levels in sex hormone-binding globulin (SHBG), as well as total testosterone. 47 48 A

pooled analysis of 18 prospective studies found that SHBG levels may be inversely associated with the risk of prostate cancer. 40 Of note, a nested case-control study found that caffeine or caffeinated coffee intakes were suggested to be associated with an increased level of plasma SHBG. However, such an association was not observed between decaffeinated coffee and plasma SHBG levels. Thus, it was suggested that caffeine may be the key component in coffee, which may be responsible for determining plasma SHBG levels. 48

#### **Strengths and limitations**

A strength of this study was the inclusion of prospective cohort studies. Cohort studies could minimise the risk of selection and recall bias, which is a major concern for case-control design. Besides, large numbers of total cohort members and prostate cancer cases ensure adequate statistical power to detect even a small effect of interest. Furthermore, the dose-response analysis may further lend confidence to the study hypothesis that increased coffee consumption was linearly associated with a lower risk of prostate cancer. Besides,



most of the studies were of high quality with a low risk of bias, which could further lend confidence to the current pooled results.

This meta-analysis also has several limitations. First, one weakness is that only three databases were searched for eligible studies and other databases, especially non-English databases, were not considered in the literature search. Second, because of the observational design, unmeasured or uncontrolled confounders in the original studies may bias the pooled risk estimate; however, the residual confounding effects from the original studies were difficult to handle in a meta-analysis approach.<sup>49 50</sup> For example, the inverse association between coffee consumption and prostate cancer could be attributed to risk factors related to coffee consumption, such as physical activity and healthy diet. However, most of the original studies have considered or adjusted for these major potential confounders in the analysis. In the sensitivity analysis of restricting the metaanalysis in studies considering most confounders, the strength of association tended to be larger in comparison with the overall association. Third, misclassification of coffee consumption may occur because of the self-reported nature of exposure measurement. However, validation studies by diet records indicated a relatively high validity of coffee consumption measured by food frequency questionnaire. The correlations between questionnaire and diet records were 0.80 in US men, <sup>16</sup> 0.71 in Swedish men <sup>17</sup> and 0.72 in Japanese men. <sup>21</sup> Of note, misclassification of exposure would most likely be non-differential in cohort studies and bias the observed association toward the null. 49 50 Therefore, the true association between coffee consumption and risk of prostate cancer may be even stronger. Fourth, since coffee intake and the incidence of prostate cancer in the USA and Europe are relatively high, most of the studies were conducted in these regions. Since the effect size is small, we should be cautious when generalising the results to other areas, especially where the incidence is relatively low. Last, significant betweenstudy heterogeneity may limit interpretation of the results. The observed heterogeneity may come from various sources. For example, the highest and lowest categories of coffee intake are different in the original studies. Studies with a broader range between the highest and lowest categories were assumed to generate a higher risk estimate. Moreover, the type of coffee and different brewing methods included in the coffee consumption groups differed. The different cohort sizes and follow-up periods may also lead to heterogeneous results. Taken together, due to the significant heterogeneity in the current metaanalysis, the pooled results should be interpreted with caution.

#### **CONCLUSIONS**

This study suggests that increased coffee consumption may be associated with a reduced risk of prostate cancer. Further research is still warranted to explore the underlying mechanisms and active compounds in coffee. If the association is further proved to be a causal effect, men might be encouraged to increase their coffee consumption to potentially decrease the risk of prostate cancer.

**Correction notice** This article has been corrected since it first published. The provenance and peer review statement has been included.

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**Contributors** KW obtained the funding, developed the research design, interpreted the results and also had primary responsibility for the final content. XC, YZ and ZT analysed the data and interpreted the results. XC and KW drafted manuscript. All authors critically reviewed and approved the manuscript.

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Patient consent for publication Not required.

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**Data availability statement** Data are available upon reasonable request. The data are available upon request from the corresponding author (wangkefenguro@sina. com)

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# **PRISMA 2009 Checklist**

Section/topic	#	Checklist item	Reported on page #
TITLE	•		
Title	1	Identify the report as a systematic review, meta-analysis, or both.	P1, L1-2
ABSTRACT			
Structured summary	2	Provide a structured summary including, as applicable: background; objectives; data sources; study eligibility criteria, participants, and interventions; study appraisal and synthesis methods; results; limitations; conclusions and implications of key findings; systematic review registration number.	P2, 3, L20-46
INTRODUCTION			
Rationale	3	Describe the rationale for the review in the context of what is already known.	P5, L84-87
Objectives	4	Provide an explicit statement of questions being addressed with reference to participants, interventions, comparisons, outcomes, and study design (PICOS).	P5, L88-91
METHODS			
Protocol and registration	5	Indicate if a review protocol exists, if and where it can be accessed (e.g., Web address), and, if available, provide registration information including registration number.	NA
Eligibility criteria	6	Specify study characteristics (e.g., PICOS, length of follow-up) and report characteristics (e.g., years considered, language, publication status) used as criteria for eligibility, giving rationale.	P6, L104-125
Information sources	7	Describe all information sources (e.g., databases with dates of coverage, contact with study authors to identify additional studies) in the search and date last searched.	P7, L127-140
Search	8	Present full electronic search strategy for at least one database, including any limits used, such that it could be repeated.	P7, L127-13
Study selection	9	State the process for selecting studies (i.e., screening, eligibility, included in systematic review, and, if applicable, included in the meta-analysis).	P6, L105-124
Data collection process	10	Describe method of data extraction from reports (e.g., piloted forms, independently, in duplicate) and any processes for obtaining and confirming data from investigators.	P5, L97-98
Data items	11	List and define all variables for which data were sought (e.g., PICOS, funding sources) and any assumptions and simplifications made.	P7, 8, L142-148
Risk of bias in individual studies	12	Describe methods used for assessing risk of bias of individual studies (including specification of whether this was done at the study or outcome level), and how this information is to be used in any data synthesis.	P8, L150-153
Summary measures	13	State the principal summary measures (e.g., risk ratio, difference in means).	P8,



# **PRISMA 2009 Checklist**

			L155-156
Synthesis of results	14	Describe the methods of handling data and combining results of studies, if done, including measures of consistency (e.g., $I^2$ ) for each meta-analysis.	P8, 9, L156-188

Page 1 of 2

		Page 1 of 2				
Section/topic	#	Checklist item	Reported on page #			
Risk of bias across studies	15	Specify any assessment of risk of bias that may affect the cumulative evidence (e.g., publication bias, selective reporting within studies).	P9, L184-185			
Additional analyses	16	Describe methods of additional analyses (e.g., sensitivity or subgroup analyses, meta-regression), if done, indicating which were pre-specified.	P9, L170-182			
RESULTS						
Study selection	17	Give numbers of studies screened, assessed for eligibility, and included in the review, with reasons for exclusions at each stage, ideally with a flow diagram.	P10, L192-204			
Study characteristics	18	For each study, present characteristics for which data were extracted (e.g., study size, PICOS, follow-up period) and provide the citations.	P10, 11, L206-216			
Risk of bias within studies	19	Present data on risk of bias of each study and, if available, any outcome level assessment (see item 12).				
Results of individual studies	20	For all outcomes considered (benefits or harms), present, for each study: (a) simple summary data for each intervention group (b) effect estimates and confidence intervals, ideally with a forest plot.	P11, L218-219			
Synthesis of results	21	Present results of each meta-analysis done, including confidence intervals and measures of consistency.	P11, L220-227			
Risk of bias across studies	22	Present results of any assessment of risk of bias across studies (see Item 15).	P11, L227-230			
Additional analysis	23	Give results of additional analyses, if done (e.g., sensitivity or subgroup analyses, meta-regression [see Item 16]).	P11, 12, L232-244			
DISCUSSION						
Summary of evidence	24	Summarize the main findings including the strength of evidence for each main outcome; consider their relevance to key groups (e.g., healthcare providers, users, and policy makers).	P14, L259-263			
Limitations	Discuss limitations at study and outcome level (e.g., risk of bias), and at review-level (e.g., incomplete retrieval of identified research, reporting bias).					
Conclusions	26	Provide a general interpretation of the results in the context of other evidence, and implications for future research.	P18,			



# **PRISMA 2009 Checklist**

			L357-361
FUNDING			
Funding	27	Describe sources of funding for the systematic review and other support (e.g., supply of data); role of funders for the systematic review.	P19, L364-366

From: Moher D, Liberati A, Tetzlaff J, Altman DG, The PRISMA Group (2009). Preferred Reporting Items for Systematic Reviews and Meta-Analyses: The PRISMA Statement. PLoS Med 6(7): e1000097. doi:10.1371/journal.pmed1000097

For more information, visit: www.prisma-statement.org.

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## MOOSE Guidelines for Meta-Analyses and Systematic Reviews of Observational Studies\*

P1, L1-2

		P1, L1-2
	Topic	Page nR2n3eL20-40
Title	Identify the study as a meta-analysis (or systematic review)	
Abstract	Use the journal's structured format	P5, <u>L</u> 84-87
Introduction	Present:	,
	The clinical problem	<del>  P5, L</del> 87-88
	The hypothesis	P5, L88-90
	A statement of objectives that includes the study population, the condition of interest, the exposure or intervention, and the outcome(s) considered	
Sources	Describe:	P5, L96
	Qualifications of searchers (eg, librarians and investigators)	P7, L129-130
	Search strategy, including time period included in the synthesis and keywords	P7, L133-140
	Effort to include all available studies, including contact with authors	P7, L127-128
	Databases and registries searched	
	Search software used, name and version, including special features used (e.g. explosion)	P7, L128-130
	Use of hand searching (e.g, reference lists of obtained articles)	P7, L130-13
	List of citations located and those excluded, including justification	NA
	Method of addressing articles published in languages other than English	NA
	Method of handling abstracts and unpublished studies	P7, L133-13
	Description of any contact with authors	P7, L133-140
Study Selection	Describe	
	Types of study designs considered	P6, L105-106
	Relevance or appropriateness of studies gathered for assessing the hypothesis to be tested	P6,L105-124
	Rationale for the selection and coding of data (eg, sound clinical principles or convenience)	NA NA
	Documentation of how data were classified and coded (eg, multiple raters, blinding, and inter-rater reliability)	P5, <b>L</b> 96-99
	Assessment of confounding (e.g. comparability of cases and controls in studies where appropriate)	P8, L161-167
	Assessment of study quality, including blinding of quality assessors; stratification or regression on possible predictors of study results	P8, L149-15
	Assessment of heterogeneity	F0, L149-150
	Statistical methods (eg, complete description of fixed or random effects models, justification of whether the chosen models account for predictors of study results, dose-response models, or	P9, L183-184
	cumulative meta-analysis) in sufficient detail to be replicated  Present	P8-9,
Results		L155-188
	A graph summarizing individual study estimates and the overall estimate	
	A table giving descriptive information for each included study	Figure 2
	Results of sensitivity testing (eg, subgroup analysis)	Supplementary Document
	Indication of statistical uncertainty of findings	P11-12, L231-24
Discussion	Discuss	
	Strengths and weaknesses	P11-12, L217-24
	Potential biases in the review process (eg, publication bias)	

P16-18, L312-353

P17, L324-341

P17, L320-321 P18, L344-346

Assessment of quality of included studies	P18, L346-354
Consideration of alternative explanations for observed results	
Generalization of the conclusions (ie, appropriate for the data presented and within the domain of the	<del>P18, L</del> 358-361
literature review)	P19, L364-366
Guidelines for future research	
Disclosure of funding source	

<sup>\*</sup>Modified from Stroup DF, Berlin JA, Morton SC, Olkin I, Williamson GD, Rennie D, et al. Meta-analysis of observational studies in epidemiology: a proposal for reporting. Meta-analysis Of Observational Studies in Epidemiology (MOOSE) group. JAMA 2000;283:2008–12. Copyrighted © 2000, American Medical Association. All rights reserved.

**Pubmed (N=116)** 

Search date: up to Sep 21, 2020

Search terms:

("coffee"[MeSH Terms] OR "coffee"[All Fields] OR "coffee s"[All Fields] OR "coffees"[All Fields]) AND ("prostat"[All Fields] OR "prostate"[MeSH Terms] OR "prostate"[All Fields] OR "prostates"[All Fields] OR "prostatism"[MeSH Terms] OR "prostatism"[MeSH Terms] OR "prostatism"[All Fields] OR "prostatitis"[MeSH Terms] OR "prostatism"[All Fields] OR "cancerated"[All Fields] OR "cancerized] OR "cancerized"[All Fields] OR "cancerized] OR "cancerize

Web of Science (N=162)

Search date: up to Sep 21, 2020

Search terms:

TOPIC: (coffee) AND TOPIC: (prostate) AND TOPIC: (cancer OR carcinoma OR neoplasm OR tumor) Indexes=SCI-EXPANDED, CPCI-S, CCR-EXPANDED, IC Timespan=All years

**Embase (N=219)** 

Search date: up to Sep 21, 2020

Search terms:

coffee AND prostate AND (cancer OR carcinoma OR neoplasm OR tumor)

Table 1. Characteristics of cohort studies of coffee consumption and prostate cancer risk included in the meta-analysis

											NOS total
				Size of		Exposure	Definition of				quality score;
		Study	Age at	cohort/	No. of	assessment	coffee		Follow-		Risk of bias
Study	Country	period	baseline	controls	cases	methods	consumption	Outcome	up time	Confounder adjustments	(Potential bias)
Ong et al.	UK	2006-2010	37-73	131834	7532	Self-reported	1 cup/day	Total	<5 years	Age, townsend deprivation index,	7 stars; low risk
2019			years			diet survey	increase; no	prostate		top 10 ancestral principal	of bias (exposure
							information on the	cancer		components, smoking status,	misclassification
							highest and lowest			BMI, height, alcohol intake, drink	bias)
							coffee intakes			temperature, overall heath rating,	
										highest qualification. Instrumental	
										variable analyses (SNP	
										instruments) were also used to	
										control confounders	
Sen et al.	Europe	1990s-2015	Mean:	142196	7036	Validated	The highest	Total,	Mean: 14	Stratified by center and age at	9 stars; low risk
2019			52			FFQ	intake: median of	Localized,	years	recruitment in 5 years categories,	of bias
			years.				855 ml/day (no. of	advanced		and adjusted for smoking status,	
							cases: 1271);	prostate		BMI, history of diabetes, alcohol	
							The lowest intake:	cancer		intake, education, physical	
							median of 0			activity, energy intake, as well as	
							ml/day (no. of			calcium, fish, tea, fruit and	
							cases: 396)			vegetable intake.	
Pounis et al	Italy	2005-2010	≥50	6989	100	Validated	The highest	Total	Mean:	Age, energy intake, smoking	8 stars; low risk
2017			years;			FFQ	intake: >3	prostate	4.24 years	habits and BMI	of bias
							cups/day (>90	cancer			

			Mean:				g/day) (no. of				
			67 years				cases: 14);				
							The lowest intake:				
							0-2 cups/day (0-				
							55 g/day) (no. of				
							cases: 45)				
Hashibe et	USA	1992-2011	55-74	46771	3037	Validated diet	Mean coffee	Total	>10 years	Age, sex, race, and education.	8 stars; low risk
al. 2015			years			history	intake is 1.9	prostate			of bias
						questonnaire	cups/day;	cancer			(confounding
							The highest				bias)
							intake: ≥2				
							cups/day (no. of				
							cases: 1731);				
							The lowest intake:				
							<1 cups/day (no.				
							of cases: 889)				
Tverdal et	Norway	1974-1999	20-69	224234	5740	Questionnaire	The highest	Total	Mean:	Age, smoking, BMI, height,	8 stars; low risk
al. 2015			years				intake: ≥9	prostate	17.6 years	physical activity, total cholesterol,	of bias (exposure
							cups/day (no. of	cancer		triglycerides, systolic blood	misclassification
							cases: 642);			pressure, year of examination and	bias)
							The lowest intake:			diabetes	
							none (no. of				
							cases: 389)				
Li et al.	Japan	1995-2005	40-79	18,853	318	Validated	The highest	Total	11 years	Age, education, BMI, time	8 stars; low risk
2013			years			FFQ	intake: ≥3	prostate		engaging in sports or exercise,	of bias
										marital status, time spent walking,	

							cups/day (no. of	cancer		smoking status, family history of	
							cases: 24);	incidence		cancer, job status, total energy	
							The lowest intake:			intake, passive smoking, alcohol	
							none (no. of			drinking, daily consumption of	
							cases: 84)			miso soup	
Discacciati	Sweden	1998-2010	45-79	44,613	3801	Validated	The highest	Localized	13 years	Age, tea, alcohol, BMI, diabetes,	8 stars; low risk
et al. 2013			years			self-	intake: ≥6	and		family history of prostate cancer,	of bias
						administered	cups/day (median	advanced		smoke, physical activity,	
						FFQ	of 1484 g/day)	prostate		education, total energy intake.	
							(no. of cases:	cancer			
							173);	incidence			
							The lowest intake:	Prostate			
							none (median: 0	cancer			
							g/day) (no. of	mortality			
							cases: 129)				
Bosire et al.	USA	1995-2008	50-71	288,391	23335	Validated	The highest	Total	>11 years	Age, race, height, BMI, physical	8 stars; low risk
2013			years			FFQ	intake: ≥6	prostate	(median:	activity, smoking, history of	of bias
							cups/day (no. of	cancer	10.5	diabetes, family history of	
							cases: 787);	incidence	years)	prostate cancer, PSA testing,	
							The lowest intake:			intakes of tomato sauce, alpha-	
							none (no. of			linolenic acid, and total energy	
							cases: 2136)			intake.	
Shafique et	UK	1970-2007	21-75	6017	318	Self-	The highest	Total	37 years	Age at screening, cholesterol,	8 stars; low risk
al. 2012			years			administered	intake: ≥3	prostate	(median:	systolic blood pressure, BMI,	of bias (exposure
			(median			questionnaire	cups/day (no. of	cancer	28 years)	alcohol intake, tea consumption,	misclassification
							cases: 65);	incidence		smoking status, social class.	bias)

			: 48				The lowest intake:				
			years)				none (no. of				
							cases: 139)				
Wilson et al. 2011	USA	1986-2006	40-75 years	47,911	5035	Validated FFQ	The highest intake: ≥6 cups/day (no. of cases: 152); The lowest intake: none (no. of cases: 587)	Total prostate cancer incidence	20 years	Age in months, calendar time, race, BMI at age 21, current BMI, vigorous physical activity, smoking, diabetes, family history of prostate cancer in father or brother, multivitamin use, intakes of processed meat, tomato sauce, calcium, alpha-linolenic acid, supplemental vitamin E, alcohol intake, energy intake, history of	9 stars; low risk of bias
Nilsson et	Sweden	1985-2007	40-60	30,930	653	Validated	The highest	Total	15 years	PSA testing.  Age, BMI, smoking, education,	8 stars; low risk
al. 2010	Sweden	1965-2007	years (median : 50 years)	30,930	033	Semi- quantitative FFQ	intake: ≥4  cups/day (no. of  cases: 209);  The lowest intake: <1 cup/day (no. of  cases: 60)	prostate cancer incidence	(median: 6 years)	recreational physical activity.	of bias
Iso et al. 2007	Japan	1988-1997	40-79 years	43,500	161	Self- administrated questionnaire	The highest intake: ≥2 cups/day (no. of cases: 38);	Prostate cancer mortality	Mean: 8.15 years	Age, area of study	7 stars; low risk of bias (exposure misclassification bias, confounding bias)

							cases: 146);	incidence			bias)
						interview	cups/week (no. of	cancer			(confounding
al. 1989			years			diet recall	intake: ≥5	prostate	17.4 years		of bias
Severson et	USA	1965-1986	46-68	7998	174	FFQ + 24-h	The highest	Total	Mean:	Age	7 stars; low risk
			years)				<3 cups/day.				bias)
			: 51				The lowest intake:		years)		bias, confounding
			(Median				cups/day;	mortality	15.6		misclassification
1990			years				intake: ≥5	cancer	(Mean:		of bias (exposure
Hsing et al.	USA	1966-1986	≥35	17,633	149	FFQ	The highest	Prostate	20 years	Age, tobacco use.	7 stars; low risk
											bias)
							none.				bias, confounding
						questionnaire	The lowest intake:	incidence			misclassification
al. 1994						life-style	cups/day;	cancer			(exposure
Marchand et			years			administered	intake: ≥2.5	prostate	years		risk of bias
Le	USA	1975-1989	≥45	20,316	198	Self-	The highest	Total	Median: 6	Age, ethnicity, income.	6 stars; medium
							cases: 23)				
							0 mg/day (no. of				bias)
							The lowest intake:				bias, confounding
							cases: 122);	incidence			misclassification
			•				mg/day (no. of	cancer	·		(exposure
2000			years				intake: ≥750	prostate	11.6 year		risk of bias
Ellison et al.	Canada	1970-1993	50-84	3400	145	FFQ	The highest	Total	Mean:	Age, wine consumption.	6 stars; medium
							(no. of cases: 47)				
							≤1-2 cup/month				
							The lowest intake:				

The lowest intake:

<1 cups/week (no.

of cases: 22)

BMI, body mass index; CI, confidence interval; FFQ, food frequency questionnaire; NA, not available; PSA, prostate-specific antigen; RR, relative risk;

#### Supplementary Table S3 Quality of cohort studies included in the meta-analysis

Study	Representativeness of the exposed cohort	Selection of the non-exposed cohort	Ascertainment of exposure	Outcome of interest was not present at start of study	Controls for important risk factors <sup>1</sup>	Assessment of outcome	Follow-up long enough for outcomes to occur	Adequacy of follow up of cohorts	Total quality score
Ong et al. 2019	☆	☆	-	☆	<b>\$</b> \$	☆	-	☆	7
Sen et al. 2019	$\stackrel{\wedge}{\sim}$	☆	☆	*	**	$\stackrel{\wedge}{\Rightarrow}$	☆	$\stackrel{\wedge}{\simeq}$	9
Pounis et al 2017	$\stackrel{\wedge}{\sim}$	☆	☆	*	**	$\stackrel{\wedge}{\Rightarrow}$	-	$\stackrel{\wedge}{\simeq}$	8
Hashibe et al. 2015	$\stackrel{\wedge}{\sim}$	☆	☆	*	☆	$\stackrel{\wedge}{\Rightarrow}$	☆	$\stackrel{\wedge}{\simeq}$	8
Terdal et al. 2015	☆	☆	-	$\stackrel{\sim}{\omega}$	**	$\Rightarrow$	☆	☆	8
Li et al. 2013	$\stackrel{\wedge}{\sim}$	☆	☆	☆	**	$\Rightarrow$	☆	-	8
Discacciati et al. 2013	☆	☆	$\stackrel{\sim}{\sim}$	$\stackrel{\sim}{\omega}$	**	$\Rightarrow$	☆	-	8
Bosire et al. 2013	☆	☆	$\stackrel{\sim}{\sim}$	$\stackrel{\sim}{\omega}$	**	$\Rightarrow$	☆	-	8
Shafique et al. 2012	☆	☆	-	$\stackrel{\sim}{\omega}$	**	$\Rightarrow$	☆	☆	8
Wilson et al. 2011	☆	☆	$\stackrel{\sim}{\sim}$	$\stackrel{\sim}{\omega}$	**	$\Rightarrow$	☆	☆	9
Nilsson et al. 2010	☆	☆	$\stackrel{\sim}{\sim}$	-	**	$\Rightarrow$	☆	☆	8
Iso et al. 2007	☆	☆	-	$\stackrel{\sim}{\omega}$	☆	$\Rightarrow$	☆	☆	7
Ellison et al. 2000	☆	☆	_	$\stackrel{\sim}{\omega}$	☆	$\Rightarrow$	☆	-	6
Le Marchand et al. 1994	☆	☆	-	$\stackrel{\sim}{\omega}$	☆	$\Rightarrow$	$\Rightarrow$	-	6
Hsing et al. 1990	☆	☆	-	$\stackrel{\sim}{\omega}$	☆	$\Rightarrow$	$\Rightarrow$	$\Rightarrow$	7
Severson et al. 1989	☆	☆	☆	☆	☆	☆	☆	-	7

<sup>1.</sup> A maximum of 2 stars could be awarded for this item. Studies that included adjustment for age received one star, and studies that included most of the other important confounders such as ethnicity, dietary factors (energy intake, vitamin D, dietary fat etc.), physical activity, body mass index, type 2 diabetes mellitus, alcohol and smoking received an additional star.

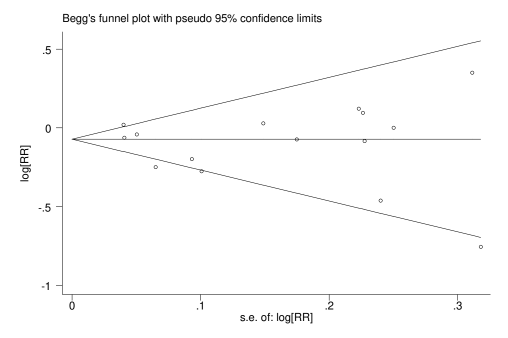


Figure S1 Begg's funnel plot of coffee consumption and prostate cancer risk

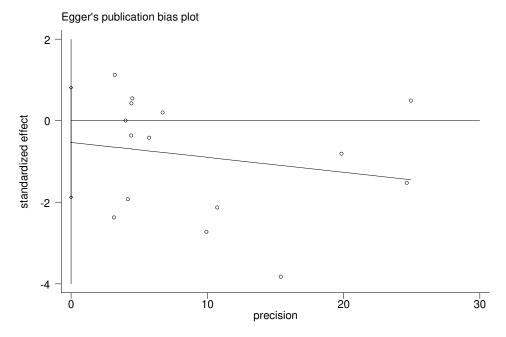


Figure S2 Egger's publication bias plot of coffee consumption and prostate cancer risk