Alcohol intake and gastric cancer: Meta-analyses of published data versus individual participant data pooled analyses (StoP Project)


Contents lists available at ScienceDirect
Cancer Epidemiology
journal homepage: www.elsevier.com/locate/canep

Alcohol intake and gastric cancer: Meta-analyses of published data versus individual participant data pooled analyses (StoP Project)


Contents lists available at ScienceDirect
Cancer Epidemiology
journal homepage: www.elsevier.com/locate/canep

Received 21 August 2017; received in revised form 16 April 2018; accepted 17 April 2018
https://doi.org/10.1016/j.canep.2018.04.009
© 2018 Elsevier Ltd. All rights reserved.
1. Introduction

Systematic reviews have the potential to settle controversies arising from apparently conflicting findings and to answer questions not directly addressed by single studies, as well as to enhance the precision of effect measures [1–4]. Individual participant data pooled analyses are considered more capable of overcoming some of the limitations of systematic reviews and meta-analyses of published data [5], since they allow access to data not previously published and statistical reanalysis based on more homogeneous criteria [3]. However, individual participant data pooled analyses require much more complex and costly management of data, as well as coordination of the underlying consortium of research groups, and the gains in terms of precision and validity of the results may be expected to vary with the topic being addressed. Comparisons of individual participant data pooled analyses with meta-analyses based on the published data from the same studies contribute to understand the extent to which conventional meta-analyses may be biased or lack statistical power, and different results may be expected for distinct research questions.

The World Cancer Research Fund reported evidence of a probable association between alcohol drinking and gastric cancer in April 2016. There were no individual participant pooled analyses for this exposure in that update [6]. The Stomach Cancer Pooling (StoP) Project [7] has recently published a pooled analysis assessing the association between alcohol intake and gastric cancer, based on information from more than 10,000 cases and 26,145 controls (15,600 men, 10,545 women) from Greece [15], Italy (four studies) [16–19], Portugal [20], Russia [21], Spain (two studies) [22,23], Sweden (three studies, two of which were nested in cohort studies) [24,25], China (four studies) [26–29], Iran (three studies) [30–32], Japan [33], Canada [34] and the United States of America (USA) (two studies, one of them unpublished) [35].

The association between alcohol drinking and gastric cancer was estimated through a two-stage modeling approach [8]. Briefly, in the first stage, the association between alcohol drinking and gastric cancer for each study was assessed through multivariable logistic regression models that included, whenever available, terms for age, sex, education/social class, smoking, fruit and vegetable consumption, study center (for multicenter studies), as well as terms for the matching variables, when applicable. In the second stage, the pooled effects estimates were computed using a random-effect models, through the DerSimonian and Laird method [36]. This was performed for the comparison of the following levels of exposure: 1) drinkers vs. non-drinkers; 2) drinkers of less than one drink per day vs. non-drinkers; 3) drinkers of one to four drinks per day vs. non-drinkers; 4) drinkers of three or more drinks per day vs. non-drinkers. Heterogeneity was quantified using the I² statistic [37].

2. Methods

2.1. Individual participant data meta-analysis

The StoP Project is a consortium of case-control studies (including nested case-control within cohort studies), including at least 80 incident, histologically confirmed, gastric cancer cases [7]. The StoP Project received ethical approval from the University of Milan Review Board.

The first release of the StoP Project dataset included 23 case-control studies, comprising 10,290 cases (6,804 men, 3,486 women) and 26,145 controls (15,600 men, 10,545 women) from Greece [15], Italy (four studies) [16–19], Portugal [20], Russia [21], Spain (two studies) [22,23], Sweden (three studies, two of which were nested in cohort studies) [24,25], China (four studies) [26–29], Iran (three studies) [30–32], Japan [33], Canada [34] and the United States of America (USA) (two studies, one of them unpublished) [35].

The association between alcohol drinking and gastric cancer was estimated through a two-stage modeling approach [8]. Briefly, in the first stage, the association between alcohol drinking and gastric cancer for each study was assessed through multivariable logistic regression models that included, whenever available, terms for age, sex, education/social class, smoking, fruit and vegetable consumption, study center (for multicenter studies), as well as terms for the matching variables, when applicable. In the second stage, the pooled effects estimates were computed using a random-effect models, through the DerSimonian and Laird method [36]. This was performed for the comparison of the following levels of exposure: 1) drinkers vs. non-drinkers; 2) drinkers of less than one drink per day vs. non-drinkers; 3) drinkers of one to four drinks per day vs. non-drinkers; 4) drinkers of over four drinks per day vs. non-drinkers. Heterogeneity was quantified using the I² statistic [37].

2.2. Meta-analysis of published data

2.2.1. Search strategy

The strategy to identify all published reports from the 23 studies included in the first version of the StoP Project database is depicted in Supplementary Fig. 1.

We searched PubMed, from inception to December 31, 2016, and conducted forward citation tracking of the reference provided in the StoP Project presentation paper to identify papers based on the same dataset, through Google Scholar and Web of Science™. The responsible investigators for each study were then asked to confirm if all published reports of results from their study had been included, and no additional
articles were identified.

2.2.2. Data extraction and meta-analysis

Two investigators (AF, SM) evaluated independently the selected studies to extract the following data from the original reports: first author, publication year, country, geographic area, number of cases and controls, period of data collection, definition of alcohol consumption, stratification variables and relative risk estimates (odds and hazard ratios) for the association between alcohol drinking and gastric cancer, along with the corresponding confidence intervals. Preference was given to estimates adjusted for the largest number of confounders, although crude estimates or data to compute them could also be extracted when only these were available.

The assessment of alcohol consumption was mainly done through questionnaires, with cases and controls describing frequency and total amount of alcohol intake. However, the description of alcohol drinking habits varied substantially among the reports. When possible, we chose non-drinkers as the reference category, described in the original reports as “non-drinkers”, “alcohol drinking: no”, “alcohol drinking: never”. In several studies, occasional drinkers or drinkers of small quantities of alcohol (defined according to quintiles of consumption) were included in the reference category. In the case of the study Italy 3 [17], non-drinkers included “individuals whose alcohol intake was less than seven grams per day”, and in the study Spain 2 [23] never drinkers were those who consumed less than one drink per month; the studies China 1 [26] and 4 [29], Greece [15] and Italy 2 [19] had a category of low consumption, either defined by the lowest quintile or the lowest amount of alcohol intake, as reference.

Different units were used to express alcohol drinking: grams per day, grams per week, kg per year, times per day, times per week, drinks per day, servings per week, glasses per week and ml per month. We converted them to drinks per day assuming the following equivalences: 1 drink = 12 g of pure ethanol and 1 ml = 0.8 g of ethanol [38]. In order to identify categories of current alcohol consumption corresponding to the exposure closest to less than one drink per day, one to four drinks per day and more than four drinks per day, we assumed that each category corresponded to an exposure equal to the midpoint of the respective category range and that the open-ended categories had the amplitude of the preceding stratum (e.g.: for surveys reporting ≤2, 2–4, 4–6, ≥6 drinks per day, 1 and 7 were the midpoints assigned to the lowest and highest category, respectively).

Data were also extracted according to cancer location within the stomach. For the purpose of analyses, results referring to “cardia” or “esophagus and gastric cardia” cancers were taken as equivalent to cancer of the gastric cardia, and “distal”, “non-cardia” or “all others” as equivalent to cancers not located in the cardia.

Meta-analyses were conducted following as close as possible the analyses described for the individual participant data pooled analyses. The DerSimonian and Laird method [36] was used to pool the estimates calculated for each study. Heterogeneity was quantified using the I² statistic [37].

2.3. Comparison between meta-analyses of published data and of individual participant data

The meta-analyses of published data and individual participant data were compared regarding the number of studies included, the estimates obtained and corresponding precision, as well as heterogeneity of results. For each of these items, the ratios of the values obtained in conventional meta-analysis and individual participant data pooled analyses (ratio MA/StoP) were computed, assuming the latter as the gold standard.
### Table 1
Comparison between meta-analyses performed with data from the published reports of the Stomach Cancer Pooling (StoP) studies and individual participant data meta-analyses regarding the number of studies, summary estimates, and corresponding precision and heterogeneity.

<table>
<thead>
<tr>
<th>Drinking category</th>
<th>Meta-analysis (MA) of published data</th>
<th>Individual participant data pooled analysis of (StoP)</th>
<th>Ratio MA/StoP</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Number of studies (N)</td>
<td>OR (95% CI)</td>
<td>SE</td>
</tr>
<tr>
<td>Drinkers vs Non-drinkersa</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>All Estimates</td>
<td>18</td>
<td>1.21 (1.07-1.36)</td>
<td>0.06</td>
</tr>
<tr>
<td>Crude Estimates</td>
<td>4c</td>
<td>1.29 (0.82-2.01)</td>
<td>0.23</td>
</tr>
<tr>
<td>Adjusted Estimates</td>
<td>14d</td>
<td>1.19 (1.05-1.36)</td>
<td>0.07</td>
</tr>
<tr>
<td>Cardia</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>All Estimates (Adjusted)</td>
<td>5e</td>
<td>1.10 (0.77-1.58)</td>
<td>0.18</td>
</tr>
<tr>
<td>Non-cardia</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>All Estimates (Adjusted)</td>
<td>5e</td>
<td>1.07 (0.86-1.34)</td>
<td>0.11</td>
</tr>
<tr>
<td>&lt; 1 drink/day vs Non-drinkers</td>
<td></td>
<td>0.95 (0.80-1.14)</td>
<td>0.09</td>
</tr>
<tr>
<td>All Estimates (Adjusted)</td>
<td>4i</td>
<td>1.17 (0.90-1.52)</td>
<td>0.13</td>
</tr>
<tr>
<td>&gt; 4 drinks/day vs Non-drinkers</td>
<td></td>
<td>1.16 (1.00-1.34)</td>
<td>0.09</td>
</tr>
</tbody>
</table>

CI – Confidence Interval; OR – Odds Ratio; SE – Standard Error.

* Reference category includes occasional/light drinkers.

* Corresponding to all StoP Project studies, except Iran 3.

* Corresponding to studies China 2, Iran 2, Greece and Spain 2 providing crude estimates or the necessary information to compute them.

* Corresponding to studies Italy 1, Italy 2, Italy 3, Italy 4, Russia, Sweden 1, Sweden 3, China 1, China 3, China 4, Iran 1, Japan, Canada and USA 1.

* Corresponding to studies Italy 4, Russia, Sweden 3, Canada and USA 1.

* Corresponding to studies Canada, Iran 1, Italy 1, Italy 2, Italy 3, Italy 4, Japan, Russia, Portugal, Spain 1, Spain 2, Sweden 1, Sweden 3, USA 1 and USA 2.

* Corresponding to studies Japan, Italy 1 and Italy 2.

* Corresponding to studies Canada, Greece, Italy 1, Italy 2, Italy 3, Italy 4, Japan, Russia, Portugal, Spain 1, Spain 2, Sweden 2, Sweden 3 and USA 2.
Funnel plots and Egger’s regression asymmetry test were used for the assessment of publication bias [4].

All statistical analyses were performed using STATA® statistical software package version 11.2 (StataCorp., College Station, Texas, USA).

3. Results

3.1. Individual-participant data pooled analysis

Twenty-two out of the 23 studies included in the first version of the StoP Project database had information to compute an estimate of the risk of gastric cancer for drinkers vs. non-drinkers [Odds Ratio (OR) (95% Confidence Interval (CI)): 1.10 (0.99–1.23)] (Fig. 1 and Supplementary Table 1).

An excess of gastric cancer risk was observed in drinkers compared to non-drinkers (Table 1), with the strongest association being observed for drinkers of more than four drinks per day [1.37 (1.19–1.58)].

3.2. Meta-analysis of published data

A total of 192 reports from the set of 23 studies included in the first release of the StoP Project dataset were identified in the systematic literature search: two from Greece, 86 from Italy, eight from Portugal, four from Russia, four from Spain, 29 from Sweden, 22 from China, three from Iran, 18 from Japan, 10 from Canada and six from the USA (Supplementary Table 2).

The analyses were performed using information extracted from 21 reports, providing data for 18 of the 23 original studies from StoP. Five studies (Iran 3, Portugal, Spain 1, Sweden 2 and USA 2) had no published data on the relation between alcohol drinking and gastric cancer. The reports from four studies (Greece, Spain 2, China 2 and Iran 2) provided only crude estimates of the association between alcohol drinking and gastric cancer, or the necessary information to compute them. A detailed description of each study and corresponding results included in the conventional meta-analysis is provided in Supplementary Table 2. The corresponding summary OR estimates for the comparison of drinkers vs. non-drinkers are presented in Supplementary Table 1 and depicted in Fig. 1; study specific estimates obtained from published reports more often supported a stronger association between alcohol drinking and gastric cancer.

3.3. Comparison between conventional meta-analyses and individual participant data pooled analyses

Table 1 and Fig. 2 show the comparison between conventional meta-analyses and individual participant data pooled analyses.

Data on the comparison between drinkers and non-drinkers were
available for a larger number of studies (18 of 22). The summary OR obtained with published data was about 10% higher than the one obtained with the StoP data (1.21 vs. 1.10) with a higher standard error (ratio MA/StoP = 1.20) and greater heterogeneity (ratio MA/StoP = 1.17). Sensitivity analyses conducted by removing each study at a time did not meaningfully change the results obtained with the StoP data or the meta-analyses of published data. Among the published reports there were 14 studies providing adjusted estimates; the corresponding summary estimates were also higher than those obtained with the StoP data from the same studies (1.19 vs 1.14).

For specific cancer locations and levels of exposure, the differences between meta-analyses were mainly regarding the number of studies and the precision of the estimates. For gastric cardia and non-cardia cancers, the ratios MA/StoP were 0.33 and 0.28, for the number of studies, respectively, and the standard error was 80% and 38% higher, respectively, than the ones from the individual participant data pooled analyses. Regarding the number of drinks per day, only around one-fifth of the StoP studies had published reports with this information, resulting in more imprecise estimates than the ones from the individual participant pooled analyses. For example, only four out of 18 studies had published data for drinkers of one to four drinks per day.

The visual inspection of the funnel plot and Egger’s test (P = 0.001) are suggestive of publication bias only for the meta-analysis of published data (Fig. 3).

4. Discussion

We observed that, for the comparison between drinkers and non-drinkers, the use of published data overestimated the association with gastric cancer. For various categories of exposure considered, the estimates obtained with both published and individual participant data were similar, although the latter were generally less heterogeneous and more precise.

Most previously published meta-analyses addressing the relation between heavy alcohol drinking and gastric cancer reported significant positive associations, particularly with an increasing number of drinks per day [9–11,13,14], although with lower magnitude than the estimates obtained in the present individual participant pooled analyses. Tramacere et al [9], using information from 10 case-control studies, estimated a 1.22 (95% confidence interval: 0.98–1.52) OR for heavy drinkers (defined as those drinking more than four drinks per day) and Bagnardi et al [11], with 11 case-control studies, estimated a similar estimate [Relative risk (RR) (95% CI): 1.22 (0.97–1.54)] for the same amount of drinks. Meta-analyses that assessed the association using information from cohort studies obtained summary estimates lower than the ones from the present analyses, [9,11,12] though in the most recent meta-analysis a statistically significant association was observed only among cohort studies, for categories of heavy drinking [11].

The differences between the summary estimates obtained in the previous meta-analyses and in the present pooled analysis may be explained by differences in the characteristics of the original studies included and by avoidance of publication bias and improved homogeneity. However, comparisons between published and unpublished data reflect essentially publication biases and heterogeneity in data analysis and presentation, and therefore the differences may vary across different sets of studies, according to the extent to which less favorable results are not published or made available with less detail.

For the present study, we obtained the most complete data possible from all the eligible studies of the StoP Project and we were able to set the results based on a reanalysis of the original studies, using uniform criteria, as the reference for comparison with a conventional meta-analysis. Our results could overestimate the differences between the two strategies of synthesis, because authors of a conventional meta-analysis may contact the authors of the original studies seeking additional data to complement those available in the published reports. However, a large proportion of systematic reviews are based only on published data, and attempts to retrieve additional data by contacting the authors from the original studies are often unsuccessful [39,40]. When comparisons between meta-analysis and pooled analysis are based on a different number of studies, disagreements may also reflect differences in the characteristics of the studies considered for each of these strategies of synthesis. In the present study, the OR estimates were very similar for the meta-analysis and the pooled analysis, and the most important differences were in terms of enhanced precision and homogeneity among the pooled analysis.

The definition of non-drinkers varied substantially across studies; in some cases they included infrequent drinkers [15,19,23,41,42], while in others the term non-drinker was not further defined [43,44]. As it
was not possible to organize, from the published data, all individuals into a common definition of non-drinker, the specific definition from each study was used, which may have contributed to the observed heterogeneity. This is a common issue when analyzing the effect of alcohol intake and is discussed in several meta-analyses for the association between alcohol and cancer [14,45,46]. Such a lack of a common definition should have led to an underestimation rather than an overestimation of the association with alcohol in the meta-analyses as compared to the pooled analyses. In addition, the latter are expected to yield more precise and valid RR estimates, namely by accounting for the potential confounding effect of socioeconomic status, fruit and vegetables consumption and tobacco smoking, [36] while also decreasing heterogeneity [2,47].

Studies comparing the results from published literature vs. individual participant data [48,49] concluded that, although the direction and magnitude of the associations may differ between methods, the main difference and advantage of pooled analyses is the number of individuals/studies available, allowing for more precise and statistically significant estimates [49]. We have previously compared the same set of reports with the individual data present in the StoP Project database regarding the association between smoking and gastric cancer, and observed that, for all exposure categories considered, StoP included a larger number of studies and had more information available to perform stratified analyses, particularly regarding cancer location and amount of cigarettes smoked per day [50]. The same pattern was observed in the present work, namely regarding the number of drinks consumed per day, with the StoP database including a larger number of studies with this information and, therefore, allowing for a dose-response analysis, as previously shown [36]. In conclusion, the differences between the estimates obtained from published and individual participant data highlight the importance of individual participant pooled data for a more comprehensive and valid appraisal of the evidence available.

Funding

This study was funded by FEDER through the Operational Programme Competitiveness and Internationalization and national funding from the Foundation for Science and Technology – FCT (Portuguese Ministry of Science, Technology and Higher Education) under the Unidade de Investigação em Epidemiologia – Instituto de Saúde Pública da Universidade do Porto (EPIUnit) (POCI-01-0145-FEDER-006862; Ref. UID/DTP/04750/2013); the PhD Grants PD/BD/105823/2014 (Ana Ferro) and SFRH/BD/102585/2014 (Samantha Morais) and the Postdoc grant SFHR/BDP/108751/2015 (Bárbara Peleteiro) co-funded by FCT and the “Programa Operacional Capital Humano” (POCh/FSE), the Fondazione Italiana per la Ricerca sul Cancro (FIRC), the Associazione Italiana per la Ricerca sul Cancro, project no. 16715 (Investigator Grant) and the Italian Ministry of Health (Young Researchers, GR-2011-02347943 to SB). Matteo Rota was supported by a fellowship from the FIRC.

The authors thank the European Cancer Prevention (ECP) Organization for providing support for the project meetings. We also thank all MCC-Spain study collaborators (CIBERESP, ISCIII, ISGlobal, ICO, University of Huelva, University of Oviedo, University of Cantabria, University of León, ibs. Granada, Instituto Salud Pública de Navarra, FISABIO, Murcia Regional Health Authority and cols).

Conflict of interest

There are no conflicts of interest to disclose.

Authorship contribution statement

The author contributions were as follows: AF collected, performed the statistical analysis and interpreted the data, drafted and revised the manuscript. SM collected and interpreted the data and revised the manuscript. MR, CP, PB, RB, CG harmonized the data, as part of the Stomach Cancer Pooling (StoP) Project. ZFZ, KM, HI, JH, KCJ, GYP, DP, MF, JM, RM, WY, HS, DZ, DM,NFL, MK, JV, EMMN, MF, FP, AW, NO, AB, NH, LM, RP, RCK, MHD, AL, PL, PB, SB, EN, BP supplied the data, as part of the StoP Project. CLV and NL supervised the analysis and interpretation of data, and reviewed the manuscript. NL defined the study hypotheses and designed the investigation. All authors contributed to the discussion of the results. All authors read and approved the final version of the manuscript.

Appendix A. Supplementary data

Supplementary material related to this article can be found, in the online version, at doi:https://doi.org/10.1016/j.canep.2018.04.009.

References


