ASSOCIATIONS BETWEEN GASTRO-OESOPHAGEAL REFLUX DISEASE AND A RANGE OF DISEASES: AN UMBRELLA REVIEW OF SYSTEMATIC REVIEWS AND META-ANALYSES

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ABSTRACT

Objective Numerous meta-analyses have revealed the association between gastro-oesophageal reflux disease (GORD) and a range of diseases; however, the certainty of the evidence remains unclear. This study aimed to summarise and assess the certainty of evidence derived from meta-analyses.

Methods Embase, PubMed, Web of Science, Cochrane Databases of Systematic Reviews, CNKI and Wangfang databases from their inception to 22 February 2020 were queried for systematic reviews and meta-analyses on the association between GORD and various diseases. The methodological quality of the included studies was assessed using A Measurement Tool to Assess Systematic Reviews 2 (AMSTAR 2), and evidence certainty was evaluated using the Grading of Recommendations, Assessment, Development, and Evaluation (GRADE) system. Statistical analysis was conducted using Stata V.15.

Results Ten publications with associations between GORD and different types of diseases were included. There was high heterogeneity (I² >75%) among seven independent meta-analyses. Evidence for publication bias in two independent meta-analyses was also observed. According to the AMSTAR 2 approach, the methodological quality was high for 20% of meta-analyses, moderate for 10%, low for 40% and critically low for 30%. Based on GRADE approach, the certainty of evidence was high for the association between GORD and higher risk of chronic obstructive pulmonary disease (COPD) exacerbation (OR 5.37; 95% CI 2.71 to 10.64) and higher prevalence of oesophageal adenocarcinoma (OR 4.57; 95% CI 3.89 to 5.37), and it was moderate for the association between GORD and higher chronic rhinosinusitis prevalence (OR 2.16; 95% CI 1.37 to 3.48).

Conclusion The association between GORD and a range of diseases was extensively studied, and our findings revealed a high certainty of evidence of the association between GORD and an increased risk of COPD exacerbation as well as increased prevalence of oesophageal adenocarcinoma. Further investigations using systematic reviews and meta-analyses of high methodological quality that include prospective large clinical trials are needed to fully understand the association between GORD and various diseases.

Strengthened and limitations of this study

► This umbrella review is the first synthesis of systematic reviews and meta-analyses to consider the association between gastro-oesophageal reflux disease (GORD) and diverse diseases.
► These results provide insights into the association between GORD and diverse diseases.
► The associations between GORD and most of diseases observed in this umbrella review may reflect uncertainty.
► Evidence from original observational studies was not discussed in this study, which may result in conclusion bias.

INTRODUCTION

Gastro-oesophageal reflux disease (GORD) is a condition characterised by stomach content reflux causing discomfort and other symptoms.1 The prevalence of GORD is increasing worldwide.2 The oesophagus is located within the chest area and connects the throat and stomach cavities with the neighbouring non-digestive tract organs, namely the larynx, heart and respiratory tract. Reflux of stomach contents (eg, pepsin, gastric acid or bile) into the oesophagus may damage the specialised physiological structure of neighbouring organs. Therefore, GORD is a risk factor for multiple diseases.

The association between GORD and various diseases has been extensively investigated, and studies have revealed that GORD is associated with respiratory diseases (eg, chronic obstructive pulmonary disease (COPD)),3 cardiovascular diseases (eg, coronary heart disease and atrial fibrillation),4 5 mental disorders,6 head and neck diseases (eg, chronic rhinosinusitis (CRS))7 and cancer (eg, laryngeal cancer).8 These findings suggest that GORD threatens physical health as well as causing various diseases that further aggravate...
the economical and psychological burden of patients.\textsuperscript{3,6} Recognising the associations between GORD and various diseases may be important for public health prevention based on the substantial global burden of diseases. However, to our knowledge, no study has comprehensively summarised these reported associations. Therefore, this study summarises the scope of associations between GORD and diverse diseases.

Although many systematic reviews and meta-analyses have shown the association between GORD and various diseases, the certainty of the evidence remains unclear. The risk of bias, scheme design defects, publication bias or inconsistencies in meta-analysis studies may decrease the certainty of evidence. It is important to clarify the certainty of the associations between GORD and the risk of various diseases for diagnosis and treatment. Therefore, the scope of the associations between GORD and diverse diseases needs to be summarised, and the strength and validity regarding these associations should be clarified using validated tools. An umbrella review is a useful tool to systematically search, collect and assess the existing evidence derived from various systematic reviews and meta-analyses on any clinical health outcomes related to a particular exposure.\textsuperscript{9,10} It provides an overview of the range and validity of the reported associations.

This study aimed to perform an umbrella review of meta-analyses on the association between GORD and various diseases and to assess the strength and validity of such evidence.

**MATERIALS AND METHODS**

The presentation of this umbrella review was in accordance with the Preferred Reporting Items for Systematic Reviews and Meta-Analyses statement.\textsuperscript{11} This umbrella review protocol has been registered in PROSPERO as CRD42019122264 (https://www.crd.york.ac.uk/prospero/).

**Literature search**

Two researchers (YZ and XL) independently and comprehensively performed computerised searches on Embase, PubMed, Web of Science, CNKI, the Wangfang database and the Cochrane Database of Systematic Reviews to identify systematic reviews and meta-analyses of observational studies on the associations between GORD and diverse diseases. The searched timeframe was from database inception to 22 February 2020. The search strategy involved a combination of medical subject terms, words and free text. The search algorithms employed for each database can be found in online supplemental appendix 1. References from relevant reviews and meta-analyses were also manually screened. All identified publications were managed using EndNote. The two reviewers (LL and JT) independently screened the titles, abstracts and full texts of eligible articles in accordance with the inclusion and exclusion criteria. Any discrepancy was resolved via discussion, and a third researcher (SX) arbitrated all discrepancies that could not be resolved through discussion.

**Eligibility and exclusion criteria**

Studies were included if they met the following criteria: (1) included systematic reviews and meta-analyses of observational studies (any type of observational study design, eg, case–control study and cross-sectional study) with pooled summary effects (eg, relative risks (RRs), ORs and their 95\% CIs); (2) considered GORD as exposure; (3) included studies in which GORD was diagnosed via validated methods (eg, oesophagastroduodenoscopy and 24-hour pH monitoring); and (4) considered the associations (ie, prevalence, incidence or risk) between GORD and diverse diseases (eg, cancer, respiratory disease and digestive disease) as the outcome.

Studies were excluded if they were systematic reviews without meta-analyses. Animal and laboratory studies were also excluded. When two or more meta-analyses were performed for the same review question, the meta-analysis with the largest sample size and the latest date of publication for each outcome was selected to avoid duplicate studies. If the most current meta-analysis was not the one with the largest sample size, then the reason for this discrepancy was explored. In our review, the meta-analysis with the largest sample size was included.

**Data extraction**

Data from each eligible systematic review and meta-analyses were independently extracted by two investigators (LL and CY). The abstracted information was carefully checked by a third investigator (XH). The name of the first author, publication year, outcomes examined, number of cases and participants, number of included studies, study design of the primary study, method of GORD diagnosis and risk of bias assessment tool were extracted using a predesigned data extraction form. For each eligible meta-analysis, the reported pooled summary effects (ORs and their 95\% CIs) and their p values as well as results of subgroup analyses were extracted. Simultaneously, we extracted the I\(^2\) and p values of Cochran’s Q test to evaluate the heterogeneity between meta-analysis studies. The results of publication bias and p values of Egger’s test were extracted to determine whether publication bias existed in the meta-analysis. If two or more outcomes of interest were examined in a systematic review and meta-analysis, each outcome was recorded separately.

**Assessment of methodological quality**

The methodological quality of the included studies was independently assessed by two investigators (DW and CY) using AMSTAR2, a measurement tool used to assess the methodological quality of systematic reviews,\textsuperscript{12} and the results were checked by a third investigator (XH). Inconsistencies were resolved by discussion. AMSTAR 2 contains 16 checklists (seven critical and nine noncritical checklists) for assessing the quality of systematic reviews and meta-analyses. According to AMSTAR 2,
items 2, 4, 7, 9, 11, 13 and 15 were defined as critical domains, and items 1, 3, 5, 6, 8, 10, 12, 14 and 16 were defined as noncritical domains. The rating criteria of the methodological quality of meta-analyses were as follows: high quality when one or no noncritical weakness was found, moderate quality when two or more non-critical weaknesses were found, low quality when one critical flaw with or without non-critical weaknesses was found and critically low quality when two or more critical flaws with or without noncritical weaknesses were found.

Assessment of certainty evidence

In this umbrella review, the GRADE system was used to evaluate the certainties of evidence derived from systematic reviews and meta-analyses of observational studies on the associations between GORD and various diseases. This system includes five factors (risk of bias, inconsistency, indirectness, imprecision and publication bias) for downgrading the certainty of evidence and three factors (large effect, dose response and plausible residual confounding) for upgrading the certainty of evidence. When there were serious or very serious downgrading factors, the certainty of the evidence was lower by one or two levels, respectively. If the effect was large (OR either >2.0 or <0.5) or very large (OR either >5.0 or <0.2), then evidence quality was upgraded by one or two levels, respectively. If there was evidence that the influence of all plausible confounding would reduce a demonstrated effect or suggest a spurious effect, then evidence quality was upgraded by one level. The rating criteria of the certainty of evidence were as follows: (1) the certainty of the primary evidence of an observational study was considered ‘low’; (2) the certainty of evidence was classified ‘very low’ by downgrading one level; (3) the certainty of evidence was rated ‘moderate’ by upgrading one level; and (4) the certainty of evidence was graded ‘high’ by upgrading two levels. GRADE assessment was conducted by two researchers (LL and JT), and the results were checked by a third investigator (XH). Inconsistencies were resolved by discussion.

Data analysis

We recalculated the summary ORs and corresponding 95% CIs. A random-effects model was used for calculation when $I^2 >50\%$ was observed; otherwise, a fixed-effects model was employed. The heterogeneity among different studies was assessed using the $I^2$ and the $p$ value results of Cochran’s $Q$ test. Publication bias was estimated using Egger’s test. Differences for which $p<0.1$ were deemed statistically significant. No heterogeneity, low heterogeneity, moderate heterogeneity and high heterogeneity were indicated by an $I^2$ of $\leq 25\%$, $25\%$–$50\%$, $50\%$–$75\%$ and $\geq 75\%$, respectively. We also conducted subgroup analysis to explain the sources of heterogeneity. Statistical analyses were conducted using Stata V.15. Figures were prepared using GraphPad Prism V.5.

RESULTS

Description of the meta-analyses

In summary, 4091 articles meeting our search criteria were initially identified from six databases. Twenty-one articles were excluded after reviewing by full text and were shown in online supplemental table S1. Ten articles published between 2007 and 2020 were selected (figure 1). The characteristics of the eligible meta-analyses are summarised in table 1. A total of 92 individual studies, consisting of 14 cross-sectional studies and 78 case–controlled studies, were included. The study estimates pooled per meta-analysis ranged from 2 to 26, and the median of the study estimate was 6. Three meta-analyses included more than 10 primary studies. The number of cases ranged from 82 to 36 503, and the number of participants ranged from 198 to 2 54 978 in the meta-analyses. Associations between GORD and idiopathic pulmonary fibrosis (IPF), asthma, COPD exacerbation, CRS, laryngeal malignancy, oesophageal adenocarcinoma (OAC), Barrett’s oesophagus, non-alcoholic fatty liver disease (NAFLD), obstructive sleep apnea–hypopnoea syndrome (OSAHS) and pharyngeal cancer were found.
Table 1  Description of the included meta-analyses of associations between GORD and diverse diseases

| Study, year | Outcomes | No. of studies | No. of cases | No. of participants | Method of GORD diagnosed | Type of metric* | Effect model | Effect estimate and 95% CI | P value of overall effects | P value of Q test | I²(%) | P value of Egger's test | Quality assessment tool | AMSTAR 2 grading |
|-------------|----------|----------------|--------------|---------------------|---------------------------|------------------|--------------|-----------------------------|--------------------------|----------------|-------|----------------------|----------------------|-------------------|
| **Association between GORD and cancers** | | | | | | | | | | | | | | |
| Rubenstein, 2019 | OAC prevalence | 5 CCS | 566 | 3285 | Symptom questionnaire and interview. | OR | Random | 4.57 (3.89 to 5.36) | <0.05 | 0.36 | 6 | 0.139 | NA | Critically low |
| Zhang, 2014 | Pharyngeal cancer risk | 2 CCS | 4681 | 18,724 | ICD-9 codes, symptomatic of GORD. | OR | Random | 3.76 (0.21 to 67.48) | 0.37 | 0.000 | 94 | 0.771 | NA | Low |
| Parsel, 2018 | Laryngeal malignancy risk | 15 CCS | 36503 | 74,209 | Symptom questionnaire, De Meester's criteria, OGD, pH monitoring; RFS; RSI score. | OR | Random | 2.37 (1.79 to 3.14) | <0.00001 | <0.00001 | 95 | 0.528 | MINORS | Moderate |
| **Association between GORD and respiratory diseases** | | | | | | | | | | | | | | |
| Havemann, 2007 | Asthma prevalence | 7 CSS | 10702 | 25,978 | Questionnaire, physician interview and database review. | OR | Random | 2.27 (1.81 to 2.84) | <0.05 | 0.000 | 85 | 0.062 | NA | Critically low |
| Huang, 2020 | COPD exacerbations risk | 3 CCS | 82 | 198 | pH monitoring or symptom questionnaire. | OR | Fixed | 5.37 (2.71 to 10.64) | <0.00001 | 0.96 | 0 | 0.696 | AHRQ | Low |
| Leason, 2017 | CRS prevalence | 4 CCS | 5391 | 219,670 | Medical records and symptom questionnaire. | OR | Fixed | 2.16 (1.37 to 3.48) | 0.001 | 0.000 | 85.4 | 0.373 | NA | Critically low |
| Bedard Methot, 2019 | IPF risk | 18 CCS | 1760 | 15,574 | Read code, pH monitoring and HARQ score. | OR | Random | 2.94 (1.95 to 4.42) | <0.05 | <0.00001 | 86 | 0.895 | NOS | High |
| Wu, 2018 | OSAHS prevalence | 1 CCS; 3 CSS | 416 | 1042 | GORD questionnaire, pH monitoring, and OGD. | OR | Random | 1.79 (1.00 to 3.22) | 0.05 | 0.03 | 67 | 0.191 | Chorane handbook | Low |
| **Association between GORD and digestive diseases** | | | | | | | | | | | | | | |
| Taylor, 2010 | BO prevalence | 26 CSS | 26 | NA | NA | Questionnaire and interview. | OR | Random | 2.90 (1.86 to 4.54) | <0.05 | 0.0001 | 89 | P† | NA | Low |
| Wijarnpreecha, 2017 | NAFLD risk | 4 CCS; 4 CSS | NA | 31,322 | Endoscopic assessment and reflux symptoms. | OR | Random | 2.07 (1.54 to 2.79) | <0.00001 | <0.00001 | 87 | P‡ | NOS | High |

*Unadjusted OR, unless otherwise specified.
†The meta-analyses did not provide the P value of published bias but mentioned no published bias.
‡The meta-analyses did not provide the P value of published bias but mentioned existed published bias.
AHRQ, Agency for Healthcare Research and Quality; BO, Barrett's oesophagus; CCS, case-controlled studies; COPD, chronic obstructive pulmonary disease; CRS, chronic rhinosinusitis; CSS, cross-sectional study; GORD, gastro-oesophageal reflux disease; HARQ, Hull Airway Reflux Questionnaire Study; IDC-9, International Classification of Diseases; IPF, idiopathic pulmonary fibrosis; MINORS, Methodological Index for Non-randomized Studies; NAFLD, non-alcoholic fatty liver disease; NOS, Newcastle-Ottawa Scale; OAC, oesophageal adenocarcinoma; OGD, oesophagogastroduodenoscopy; OSAHS, obstructive sleep apnoea-hypopnoea syndrome; RFS, Reflux Finding Score; RSI, Reflux Symptom Index.
Summary of the methodological quality of the included meta-analyses
The methodological qualities of the included meta-analyses were assessed using AMSTAR 2, and the detailed results are shown in online supplemental table S2. Five (40%) meta-analyses did not report explicit statements or protocols. Five (40%) meta-analyses did not use comprehensive literature search strategies. Six (50%) of the meta-analyses did not assess the risk of bias in the primary studies. None of the meta-analyses reported the details of funding sources for the included studies. Overall, the methodological qualities of the included meta-analyses were categorised as high for 20% (n=2), moderate for 10% (n=1), low for 40% (n=4) and critically low for 30% (n=3) (table 1).

Certainty of evidence of the associations between GORD and diverse diseases
A total of 10 associations between GORD and different types of diseases were found: cancer (n=3), respiratory diseases (n=5) and digestive diseases (n=2). Nine of these associations had nominal significance. The detailed results of the certainty of evidence are summarised in online supplemental table S3. Overall, the certainties of evidence were graded high, moderate, low and very low for 20% (n=2), 10% (n=1), 10% (n=1) and 60% (n=6) of the associations, respectively (figures 2–4).

The associations between GORD and cancers
Figure 2 shows that the individuals with weekly GORD symptoms have a fivefold increase in the odds of developing OAC (OR=4.57; 95% CI 3.89 to 5.36), and the certainty of evidence was rated as high. A very low certainty of evidence indicated that GORD is associated with an increased risk of laryngeal malignancy by more than twofold (OR=2.37; 95% CI 1.79 to 3.14) (figure 2). However, a very low certainty of evidence showed that there was no significant association between GORD and pharyngeal cancer. The results of subgroup analyses are shown in table 2. The individuals with daily GORD symptoms had a sevenfold increase in odds of developing OAC. By stratifying based on duration of GORD symptoms, the odds of developing OAC were higher among individuals with GORD symptoms of at least 20 years was higher among individuals with GORD symptoms of less than 10–15 years. Based on ethnicity, GORD increased the risk of laryngeal malignancy in Europeans, Asians and Americans, particularly in Asians. GORD significantly increased the prevalence of OAC in Europeans, Australians and Americans. Additionally, GORD increased the risk of laryngeal malignancy by approximately fourfold based on diagnoses of GORD using objective methods. However, there was no association between GORD and laryngeal malignancy based on diagnoses of GORD using subjective methods.

The associations between GORD and respiratory diseases
Figure 3 shows the association between GORD and increased risk of COPD exacerbation (OR=5.37; 95% CI 2.71 to 10.64), and the certainty of evidence was high. The asthma prevalence in individuals with GORD was higher than those without GORD (OR=2.27; 95% CI 1.81 to 2.84), but the certainty of evidence was very low. CRS prevalence increased by more than twofold in patients with GORD (OR=2.16; 95% CI 1.37 to 3.48), and the certainty of evidence was moderate. The summary OR for the association between GORD and IPF was OR=2.94 (95% CI 1.95 to 4.42), but the certainty of evidence was very low. The certainty of the positive association between GORD and OSAHS (OR=1.79; 95% CI 1.00 to 3.22) was very low.

The results of subgroup analyses are shown in table 2. GORD was associated with an increase in the risk of OSAHS...
based on the diagnoses of GORD using objective methods. However, there was no association between GORD and OSAHS based on the diagnosis of GORD using subjective methods. Based on the age of the participants, GORD was associated with a higher risk of CRS in both adults and children, particularly in children (OR=3.29; 95% CI 2.39 to 4.27). Based on ethnicity, GORD was associated with an increase in the risk of asthma in Europeans, Asians and Americans, particularly in Asians, with a summary OR of 5.56. GORD was associated with a higher prevalence of OSAHS in Americans, but not in Asians, and GORD was associated with an increase in the risk of IPF in Europeans, Asians, Americans and Africans, but not in Canadians.

The associations between GORD and digestive diseases
The prevalence of Barrett’s oesophagus in individuals with GORD was approximately three times higher in those without GORD (OR=2.90; 95% CI 1.86 to 4.54), and the certainty of evidence was rated as low (figure 4). Based on ethnicity, GORD was associated with an increase in the prevalence of Barrett’s oesophagus in Americans by twofold. However, there were no associations between GORD and Barrett’s oesophagus based on studies involving Asians (OR=1.62; 95% CI 0.813 to 3.24) and Europeans (OR=3.00; 95% CI 0.901 to 9.99) (table 2). GORD was significantly associated with an increase in the risk of NAFLD with a very low certainty of evidence (figure 4).

Heterogeneity and publication bias of the included meta-analyses
The heterogeneity and publication bias of the included meta-analyses are summarised in table 1. Eight of the included systematic reviews with meta-analyses showed heterogeneity (I² ≥30%). Among these, seven showed high heterogeneity with an I² of >75%, and one meta-analysis showed moderate heterogeneity with an I² of 67%. The results of subgroup analyses showed that heterogeneity in studies significantly decreased when stratifying by age of participants, study design, ethnicity and diagnostic methods. Two of the included meta-analyses showed statistical evidence of publication bias.

DISCUSSION
Principal findings
The influence of GORD on the development of diverse diseases has been examined in many published meta-analyses. This umbrella review included 10 published meta-analyses that investigated on the association between GORD and 10 types of diseases. Approximately 80% of the meta-analyses reported significant heterogeneity. Two meta-analyses showed statistically significant evidence of publication bias. The methodological qualities were categorised as high only for two meta-analyses and moderate for one. The certainties of evidence were graded as high for the association between GORD and increased risk of COPD exacerbation and for the association between GORD and increased prevalence of OAC.

Comparison with other studies
This umbrella review supported some of the recommendations as well as added related evidence. Recommendations on the contribution of GORD to the development of COPD were included in the guidelines. This information was in accordance with our result that GORD was associated with an increased risk of COPD exacerbation. Therefore, controlling GORD in patients with COPD may contribute to the control of COPD and improve prognosis. Additionally, our findings showed that GORD was associated with an increase in the prevalence of OAC, indicating that GORD may play an important role in the aetiology of OAC, with a high certainty of evidence.

This result indicates that controlling GORD might be beneficial in deterring OAC development. However, Australian guidelines state that the treatment of GORD with proton pump inhibitors (PPIs) does not influence OAC progression. Larger randomised controlled trials should be conducted in the future to confirm this discrepancy. Chinese guidelines have reported GORD as a risk factor of CRS but failed to establish the strength of their evidence. This umbrella review confirmed that the certainty of this evidence may be moderate.

Japanese guidelines recommended the management of the coexistence of a condition coexisting with GORD when treating asthma. If asthma is not been managed at any step during therapy, then the assessment of the coexisting of a condition with GORD is also recommended. Korean guidelines indicate that GORD occurs in up to 87% of patients with IPF and can contribute to the exacerbation of IPF. The American College of Gastroenterology recommends the routine use of PPIs for Barrett’s oesophagus even in the absence of symptoms. However, we found that the certainties of evidence for the association between GORD and these diseases are ‘low’ or ‘very low’, indicating that these associations should be assessed.
| Subgroup classification | Association between GORD and* | Number of studies | Effect model | OR (95% CI) | P value of Q test | I² (%) |
|-------------------------|-------------------------------|-------------------|--------------|-------------|------------------|--------|
| **Stratified by frequency of GORD symptoms** | | | | | | |
| Daily symptoms | OAC prevalence | 5 CCS | R | 7.40 (4.94 to 11.1) | 0.01 | 71 |
| Weekly symptoms | OAC prevalence | 5 CCS | R | 4.57 (3.89 to 5.36) | 0.04 | 60 |
| **Stratified by duration of GORD symptoms** | | | | | | |
| At least 20 years | OAC prevalence | 4 CCS | R | 5.41 (2.45 to 11.9) | <0.01 | 89 |
| Less than 10–15 years | OAC prevalence | 4 CCS | R | 3.05 (1.53 to 6.08) | <0.01 | 84 |
| **Stratified by diagnostic methods of GORD** | | | | | | |
| Subjective methods | Laryngeal malignancy risk | 5 CCS | F | 1.43 (0.93 to 2.22) | 0.1 | 48 |
| OSAHS prevalence | 2 CSS | F | 1.63 (1.21 to 2.19) | 0.455 | 0 |
| IPF risk | 12 CCS | R | 2.36 (1.82 to 3.05) | 0 | 91.1 |
| Objective methods | OSAHS prevalence | 1 CCS | NA | 1.21 (0.67 to 2.18) | NA | NA |
| IPF risk | 11 CCS | R | 2.80 (1.57 to 5.00) | 0.007 | 59 |
| Laryngeal malignancy risk | 11 CCS | F | 3.82 (2.61 to 5.59) | 0.14 | 32 |
| **Stratified by study design** | | | | | | |
| Prospective studies | Laryngeal malignancy risk | 12 CCS | F | 2.46 (1.57 to 3.85) | NA | 68 |
| Retrospective studies | Laryngeal malignancy risk | 8 CCS | F | 2.68 (1.86 to 3.85) | NA | 97 |
| Cross-sectional studies | NAFLD risk | 4 CCS | F | 1.52 (1.15 to 2.00) | 0.001 | 86 |
| Case–control studies | NAFLD risk | 4 CCS | F | 3.04 (2.27 to 4.06) | 0.7 | 0 |
| **Stratified by age of participants** | | | | | | |
| Adults with GORD | CRS prevalence | 3 CCS | F | 1.66 (1.57 to 1.75) | 0.426 | 0 |
| Children with GORD | CRS prevalence | 1 CCS | NA | 3.20 (2.39 to 4.27) | NA | NA |
| **Stratified by ethnicity of participants** | | | | | | |
| Europeans | Asthma prevalence | 3 CSS | F | 1.98 (1.79 to 2.20) | 0.51 | 0 |
| BO prevalence | 8 CCS | R | 3.00 (0.901 to 9.99) | <0.0001 | 93 |
| OAC prevalence | 2 CCS | R | 5.59 (3.02 to 10.33) | 0.02 | 81 |
| IPF risk | 6 CCS | R | 2.05 (1.69 to 2.49) | 0 | 77.5 |
| Laryngeal malignancy risk | 6 CCS | F | 4.72 (3.16 to 7.06) | 0.233 | 26.9 |
| Asians | IPF risk | 2 CCS | F | 4.28 (1.81 to 10.11) | 0.548 | 0 |
| Asthma prevalence | 2 CSS | R | 5.56 (1.66 to 18.67) | 0.005 | 87.5 |
| BO prevalence | 3 CCS | F | 1.62 (0.813 to 3.24) | 0.14 | 49 |
| Laryngeal malignancy risk | 2 CCS | F | 4.79 (2.26 to 10.15) | 0.78 | 0 |
| OSAHS prevalence | 2 CSS | R | 8.88 (0.15 to 528.17) | 0.005 | 87.6 |
| Americans | BO prevalence | 12 CCS | R | 2.44 (1.42 to 4.23) | <0.0001 | 87 |
| Asthma prevalence | 2 CSS | F | 1.69 (1.60 to 1.77) | 0.64 | 0 |
| OAC prevalence | 3 CCS | F | 4.09 (3.23 to 5.18) | 0.87 | 0 |
| IPF risk | 8 CCS | R | 3.27 (1.83 to 3.79) | 0 | 89.2 |
| Laryngeal malignancy risk | 7 CCS | R | 1.69 (1.21 to 2.36) | 0 | 97.7 |
| OSAHS prevalence | 1 CCS;1CSS | F | 1.53 (1.00 to 2.36) | 0.25 | 25 |

Continued
with caution. Although certainties of evidence for these associations were graded ‘low’ or ‘very low’, they might still provide ideas for clinicians. Therefore, more well-documented research studies should be conducted to confirm these associations.

In a previous review, Chen summarised the findings of primary studies and meta-analyses on the association between GORD and non-digestive tract diseases. However, that review only focused on the association between GORD and non-digestive tract diseases and not that between GORD and digestive tract diseases. Furthermore, it was only a narrative review, not an umbrella review. The methodological quality of meta-analyses and the certainty of evidence of the existing evidence were not evaluated in this study. Our umbrella review not only provides a broad overview of the current existing evidence derived from meta-analyses but also evaluates the methodological quality of the published meta-analyses and the certainty of their evidence.

### Possible explanations

We found that GORD was positively associated with an increased risk for nine diseases. However, all of the ORs abstracted from the included meta-analyses were not adjusted for confounders (eg, smoking and drinking). Nevertheless, the influence of confounding factors on the results cannot be ruled out, particularly in the analysis of respiratory diseases (eg, IPF and COPD exacerbation). Additionally, estimates with a very wide 95% CI, such as the association between GORD and pharyngeal cancer also affect the accuracy of evidence. Therefore, this association might be inconclusive.

High certainty of evidence was observed for the positive association between GORD and increased risk of COPD exacerbation. Microaspiration of gastric contents (eg, gastric acids, bile and pepsin) reflux may have an important role in the development of this disease. A previous study detected pepsin in the bronchoalveolar fluid of all of the patients with COPD. Interestingly, pepsin levels in the lung could increase pulmonary protein oxidation. Additionally, the trigger of COPD exacerbations may be related to proximal acidic reflux, and extensive proximal acidic reflux may contribute to pulmonary oxidative stress. A previous study demonstrated an increase in protein oxidation in the lungs of patients with GORD compared with healthy individuals.

Therefore, GORD may substantially contribute to inflammatory and oxidative damage in the lung. In addition, dysregulation of vagal nervous system induced by exposure of gastric acids is also a major mechanism explained the association between GORD and COPD exacerbation. Simultaneously, abnormal breathing physiology in patients promote the occurrence of GORD. Both recurrent episodes of COPD and GORD require frequent use of medications. The frequent application of β2-agonists could exacerbate gastro-oesophageal reflux. This situation results in a pathological vicious cycle.

Additionally, a high certainty of evidence showed that GORD is associated with an increased prevalence of OAC, particularly in individuals with daily GORD symptoms and a symptom duration of at least 20 years. This association may be explained by biological changes caused by gastric content reflux. Gastric contents (eg, bile acids, pepsin and trypsin) promote interleukin-8 production from oesophageal epithelial cells that could induce epithelial cell transformation and promote cancer cell proliferation. Additionally, a decrease in pH in the oesophagus plays an important role in the occurrence of cancer. In previous studies, oesophageal tissues are stimulated repeatedly by the refluxate, causing oesophageal tissue to be repeated damaged as well as inducing chronic inflammation, thereby promoting the development of cancer.

The association between GORD and increased prevalence of CRS warrants further investigation. The certainty of evidence was graded as moderate. A previous study established that pepsin may potentiate CRS by damaging mitochondria in nasal epithelial cells, which may explain the mechanism of GORD that is associated with an increase in CRS prevalence.

Subgroup analyses showed that GORD increases the risk of Barrett’s oesophagus in Americans, but not in Europeans and Asians. Differences in lifestyle and ethnicity might explain this difference. Associations without significance also should be interpreted with caution. Although GORDs were not associated with increased risk of Barrett’s oesophagus in Europeans and Asians, these associations might also suggest an association between these two diseases because of the summary ORs of 3.00 and 1.62, respectively.

This study demonstrated that 80% of the meta-analyses had moderate to high heterogeneity ($I^2 \geq 50\%$).

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**Table 2** Continued

| Subgroup classification | Association between GORD and* | Number of studies | Effect model | OR (95% CI) | P value of Q test | I$^2$ (%) |
|------------------------|-----------------------------|------------------|-------------|-------------|-----------------|--------|
| Australians            | OAC prevalence$^{41}$        | 1 CCS            | NA          | 5.48 (4.23 to 7.10) | NA               | NA     |
| Canadians              | IPF risk$^{36}$             | 1 CCS            | NA          | 0.90 (0.30 to 2.69) | NA               | NA     |
| Africans               | IPF risk$^{36}$             | 1 CCS            | NA          | 10.52 (2.27 to 48.76) | NA               | NA     |

BO, Barrett’s oesophagus; CCS, case-controlled studies; CRS, chronic rhinosinusitis; CSS, cross-sectional study; GORD, gastro-oesophageal reflux disease; IPF, idiopathic pulmonary fibrosis; NAFLD, non-alcoholic fatty liver disease; OAC, oesophageal adenocarcinoma; OSAHS, obstructive sleep apnoea–hypopnoea syndrome.
These sources of heterogeneity may be explained by differences in the age of participants, study design, ethnicity, and diagnostic methods. High heterogeneity indicates that some associations may simply be false positives or exaggerated. Therefore, considering the substantial heterogeneity, only the associations between GORD and higher risk of COPD exacerbations and OAC development are genuine, but this does not mean that other associations are not true. We found that two meta-analyses showed statistically significant evidence of publication bias. Publication bias favours the reporting of significant positive results. Therefore, the association between GORD and asthma and NAFLD should be interpreted with caution.

**Strengths, limitations and future studies**

This umbrella review has several strengths. It is the first to systematically summarise the associations between GORD and various diseases and to critically evaluate the methodological quality of meta-analyses and the certainty of existing evidence. The strength of an umbrella review has been demonstrated in other articles. In this review, AMSTAR 2 was used to assess the methodological quality of meta-analyses, and the GRADE system was adopted to evaluate the certainty of evidence. AMSTAR 2 and GRADE are validated tools, and their efficiency and reliability have been exhibited in other published studies. Some defects of the existing published meta-analyses were found throughout this umbrella review, which may facilitate the improvement of future studies. Our umbrella review also found gaps in evidence of the associations between GORD and other diseases (eg, rheumatoid arthritis), which should be considered in future studies.

This umbrella review also has several limitations. Although AMSTAR 2 and GRADE are validated tools, the use of other tools such as Confidence in the Evidence from Reviews of Qualitative Research and Risk Of Bias In Systematic reviews could have led to different conclusions regarding the certainty of evidence. We only focused on existing and published systematic reviews with meta-analyses on observational studies, which may have resulted in conclusion bias. Evidence from individual observational studies was beyond the scope of our discussion. We did not use $\tau^2$ and 95% prediction intervals to categorise heterogeneity; instead, we used the Q test p value and the I² results, which might have decreased the accuracy of the GRADE assessment. All the primary studies included in the published meta-analyses were case-control or cross-sectional studies, so the risk of recall bias or selection bias could not be avoided. A large effect size was found in some meta-analyses; all of them were unadjusted for confounders (eg, body mass index, smoking, abdominal obesity, frequent coughing, or frequent use of $\beta_2$ agonists, or *Helicobacter infection*), which may have affected the validity of the evidence. The small sample size and high heterogeneity of meta-analyses may decrease the strength and validity of evidence. The method of diagnosis of GORD includes clinical history and questionnaires, proton pump inhibitor trial, endoscopy and biopsy, pH-metry and ambulatory reflux monitoring. GORD is usually diagnosed by symptoms. The latest studies have shown that the specificity and specificity of GORD diagnosis by symptoms are 64.9% and 71.4%, respectively. Therefore, the diversity of diagnostic methods for GORD results might also decrease the strength and validity of evidence. Lastly, we included the largest and latest meta-analyses, but these might not have had the highest certainty of evidence and thus might have influenced the conclusions generated by this umbrella review.

Several factors should be considered in future studies to achieve high certainty of evidence. Prospective cohort studies on the associations between GORD and various diseases with a larger sample size should be conducted in the future using time-varying exposure (GORD duration, control or treatment). Adjusting for confounders such as age, sex, smoking status, body mass index, frequent use of $\beta_2$ agonists and *H. pylori* infection may assist in rendering the role of GORD in the development of these diseases much clearer in future. Evidence from meta-analyses with publication bias should be reconfirmed in future studies. Most of the meta-analyses had moderate to high heterogeneity, indicating that associations could be inflated or may be false positives. Well-controlled primary studies should be conducted to reduce the heterogeneity of meta-analyses. The association with high heterogeneity should be further explored by conducting meta-analyses with low heterogeneity. We found gaps in evidence derived from systematic reviews and meta-analyses on the associations between GORD and health disorders in this umbrella review because meta-analyses on those topics have not been performed. However, some individual studies have described these associations despite inconsistent conclusions. Thus, future meta-analyses should be conducted to confirm those findings and resolve gaps in information.

**CONCLUSIONS**

This umbrella review systematically summarised the associations between GORD and various diseases derived from systematic reviews and meta-analyses and evaluated their methodological quality of the meta-analyses and the certainty of existing evidence. GORD is associated with oesophageal adenocarcinoma, Barrett’s oesophagus, NAFLD, CRS, laryngeal malignancy, OSAHS, IPF and COPD exacerbation, but it is not associated with the risk of pharyngeal cancer. There was high certainty of evidence that GORD is a risk factor of COPD exacerbation and OAC. Further studies with systematic review and meta-analyses of high methodological quality that included prospective cohort studies with large sample sizes and adjusted confounders are needed to confirm these associations.

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Contributors JT wrote the protocol and registered it in PROSPERO. LL and JT screened the articles according to the eligibility and exclusion criteria. YZ and XL independently searched the databases and imported the yielded articles to EndNote software. LL and CY extracted the data. DW and CY conducted the AMSTAR 2 evaluating. LL and JT conducted the evidence classifications. XH checked the data and results. JX and RC conduct statistical analyses. LL, JT and CY wrote the draft of the paper, and XH revised the paper. SX was contributed in concept design, guidance and arbitrating all discrepancies and was also responsible for the final content.

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Supplemental material This content has been supplied by the author(s). It has not been vetted by BMJ Publishing Group Limited (BMJ) and may not have been properly cited, appropriate credit is given, any changes made indicated, and the use and license their derivative works on different terms, provided the original work is properly cited, otitis media, and laryngeal malignancy: a systematic review of the evidence. Am J Med 2003;115 Suppl 3A:81–9.

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