Systematic review with meta-analysis: the effects of family history on the risk of Barrett's oesophagus and oesophageal adenocarcinoma

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Funding information
This research received no specific grant from any funding agency in the public, commercial or not-for-profit sectors.

Summary

Background: Current guidelines recommend different screening approaches for individuals with a family history of Barrett's oesophagus (BO) or oesophageal adenocarcinoma (OAC), varying from no screening to screening all individuals with a positive family history.

Aims: To determine evidence-based risk estimates for individuals with a family history of BO or OAC

Methods: We systematically searched Pubmed, Embase and Cochrane Library until October 2020 to identify all studies that reported on the association between family history and the risk of BO and OAC. Pooled summary estimates of adjusted relative risks and prevalence of familial BO/OAC with 95% confidence intervals (CIs) were calculated using a random effects model.

Results: Fourteen studies comprising 16 189 BO/OAC patients were analysed. Familial clustering was seen in 8.84% (95% CI: 5.54-13.82) and 4.37% (95% CI: 2.15-8.69) of patients with BO and OAC, respectively (nine studies). Screening first-degree relatives of BO patients had a diagnostic yield between 12% and 44% for BO (four studies). However, the yield for high-grade dysplasia and OAC was low (<2%). Individuals with a positive family history had a higher risk of having BO (aRR 3.26; 95% CI 1.43-7.40; $I^2 = 46%$; three studies) and OAC (aRR 2.19; 95% CI 1.14-4.21; $I^2 = 48%$; five studies) compared to individuals without a family history.

Conclusions: A verified family history of BO or OAC is a strong risk factor for both BO and OAC. A positive family history could be a clinically meaningful way to identify high-risk individuals who may benefit from early detection strategies.
1 | INTRODUCTION

During the past 30 years, the incidence of oesophageal adenocarcinoma (OAC) has increased up to sixfold in Western countries.\(^1,2\)

With 35,000 new cases in 2018, nearly 50% of the worldwide cases of OAC occur in Europe and North America.\(^3\) OAC still has a poor prognosis with a 5-year survival rate of only 20%, despite improvements in multimodality therapy.\(^4\) The vast majority of patients with OAC present with locally advanced or metastatic disease, as symptoms of early OAC and its precursor lesions are often absent or barely distinct from gastro-oesophageal reflux disease (GERD).\(^1\)

Barrett’s oesophagus (BO) is the major precursor of OAC, increasing the risk of developing OAC by a factor 10-30.\(^5,6\) The population prevalence of BO is estimated to be approximately 1%-2%, which increases to 8%-20% in individuals with long-term GERD.\(^7,8\) Unfortunately, in daily practice, >90% of patients with OAC never had prior endoscopy and only a minority of BO patients are currently diagnosed and under surveillance.\(^9\) Hence, identifying patients with BO and early detection of OAC could be potentially helpful in reducing OAC-related mortality.

As the annual risk of OAC in patients with BO is low (0.1%-0.5%), the merits of population-based endoscopic screening are controversial.\(^5,10\) It is therefore important to identify individuals at increased risk for BO and OAC. Already known risk factors for BO and OAC, including increasing age, male gender, Caucasian race, smoking, obesity and GERD, are in this regard helpful.\(^6\)

Although the vast majority of BO and OAC cases are sporadic and caused by somatic mutations, several reports of families with multiple affected relatives suggest that there may be an underlying genetic susceptibility.\(^11-14\) However, as the exact role of genetic factors in the development of BO and OAC has remained largely unclear, OAC is not included in familial risk management guidelines.\(^15\) Clinical guidelines on the other hand suggest a role for endoscopic screening in individuals with a positive family history for BO or OAC.\(^16-19\) Until now, recommendations for screening in these individuals are merely based on expert opinions and on small number of studies, and consequently recommendations vary between guidelines.\(^16-20\)

Precise and valid evidence-based risk estimates for individuals with a family history of BO or OAC are needed to improve genetic counselling, provide rational advice, develop risk prediction models and to determine appropriate screening strategies. Understanding the association between family history and oesophageal metaplasia and related neoplasia may also improve the knowledge on the pathogenesis of BO and OAC and could be key to identify causal underlying germline mutations. The aim of this systematic review and meta-analysis was therefore to determine the prevalence of a positive family history in patients with BO and OAC. We furthermore aimed to assess the prevalence and the risk of BO and OAC in individuals with a positive family history.

2 | METHODS

2.1 | Search strategy

This systematic review and meta-analysis of studies that investigated the association between a positive family history and risk of BO and related neoplasia was conducted according to the MOOSE (Meta-analysis of Observational Studies in Epidemiology) guidelines and a predetermined protocol (Prospero: CRD42020179348).\(^21\)

We systematically searched the electronic databases of MEDLINE (Pubmed), Embase (Ovid Technologies) and the Cochrane Library from inception to the date of the search (October 2020), without any restrictions. The search terms included three main categories and comprised synonyms for ‘BO’ and ‘OAC’, ‘family’ and ‘risk’ in accordance with previous literature reviews on familial cancer.\(^22,23\) The search strategy was performed in collaboration with an experienced medical librarian. Exact search terms are presented in Table S1. References of eligible articles and reviews on the topic were manually searched for additional articles. An additional literature search excluding the keywords related to family/genetics was performed to possibly identify studies that only reported family history as secondary outcome. We assessed the full text of a randomly selected 5% of identified articles to determine robustness of our search strategy.

2.2 | Study selection

All identified records were exported to the citation management program EndNote X8 (Clarivate Analytics) for deduplication. First, two reviewers (YP and EG) independently screened titles and abstracts. Second, the full text of all included abstracts was assessed by the same reviewers to determine eligibility of each study. Any discrepancies between reviewers in both screening phases were resolved by consensus. Remaining disagreements were resolved through discussion with a third reviewer (PS).

Studies were eligible for inclusion if they permitted quantitative assessment of the association between BO and/or OAC defined by their presence during upper endoscopy and validated by pathology review and family history for these diseases. A positive family history was defined as having any type of family history of BO or oesophageal cancer irrespective whether the family history was verified by assessing medical or pathology reports from reportedly affected relatives. Studies that did not provide the used definition of a positive family history or used another definition of family history (eg family history of gastric cancer or GERD) were excluded. Additionally, we excluded studies that did not describe how assessment of family history was performed or which relatives were assessed. Studies were included if they reported the proportion of patients with BO or OAC that had a positive family history, the proportion of BO and OAC diagnoses in individuals with a positive family history or a measure of association (relative risk [RR], odds ratio [OR] or standardised incidence ratio [SIR]), or provided data for their calculation.
Case reports, reviews, unpublished data and conference abstracts were excluded. Inclusion was not otherwise restricted by study size, language or study type. We reviewed all included studies for their independence of their study population. For studies from the same data source and investigating the same outcome measure, we included articles that best fitted the relevance to the study questions.

2.3 | Data extraction and quality assessment

After identifying relevant studies, two authors (YP and EG) independently abstracted data on study characteristics and quality, patient demographics, definitions and assessment of family history including endoscopic or histological confirmation of affected relatives, and type of outcome measure onto a standardised form. Of all included studies, differences between patients with familial BO/OAC and sporadic cases were also extracted. For each study, the prevalence of BO or OAC in first-degree relatives was calculated by dividing the cases by the total patient group at risk, while the prevalence of a positive family history was calculated by dividing the cases with a positive family history by the total patient group with BO or OAC. Also, risk estimates (RR, OR or SIR) with 95% confidence intervals (CIs) and potential confounding variables considered in the analyses were recorded for each study. Adjusted risk estimates were used if they were adjusted for relevant effect moderators. Otherwise, unadjusted estimates or raw data were collected. ORs and SIRs were considered to be equivalent to RRs, given that the prevalence of BO or OAC among asymptomatic individuals is relatively low. 24

Risk of bias was assessed using the Quality in Prognosis Studies (QUIPS) tool, as recommended by the Cochrane Collaboration for studies of prognostic factors. 25,26 Quality was analysed based on six domains: study participation, study attrition, prognostic factor measurement, outcome measurement, study confounding and statistical analysis and reporting. Studies were classified as either low, moderate or high risk of bias (Table S2). Disagreements in data extraction and quality assessment were resolved through discussion and consensus and in consultation with a third reviewer (PS).

2.4 | Statistical analysis

The outcomes of this study were pooled prevalences and risk estimates of both BO and OAC based on the presence or absence of a positive family history for BO or oesophageal cancer. We also assessed the proportion of patients with BO or OAC with a positive family history of these disorders. We defined familial clustering as the occurrence of two cases of BO or OAC within one family.

Proportions with 95% CIs were calculated using the method of Wilson. 27 For all outcomes, we pooled logit-transformed prevalences and risk estimates with the corresponding 95% CIs using the generic inverse variance method with a random effects model. 28,29 This model incorporates heterogeneity by giving a weight to each study equal to the inverse of the variance of the effect estimate. Between-study variance in the random effects model was estimated by a restricted maximum-likelihood estimator (REML). 30 The between-study heterogeneity was quantified with the inconsistency index ($I^2$) statistic and tested for significance using Cochrane’s Q-test. Because this test is underpowered to detect moderate degrees of heterogeneity, $P$ values <0.10 were defined as indicating the presence of heterogeneity. 31 $I^2$ values >50% indicated substantial heterogeneity.

Given the observational nature of studies and variation in the effect measures used in individual studies, we anticipated heterogeneity in the analyses. To reduce heterogeneity, a stratified meta-analysis was performed according to used definitions of a positive family history. To further explore heterogeneity, we performed pre-planned sensitivity analyses on study-related variables (study design, study location, verification of a positive family history and adjustment for certain covariates). Between-study sources of heterogeneity were assessed by using sensitivity analyses by stratifying original estimates according to the study characteristics, with $P$ values <0.05 for differences between subgroups being considered as statistically significant. To explore a GERD-independent effect of a positive family history on BO and OAC, we performed sensitivity analyses of studies that adjusted for GERD symptoms or included only patients with GERD.

We did not perform an assessment of publication bias and meta-regression because of the small number of studies (<10) for all effect estimates. Statistical calculations and transformations for proportional outcomes and risk estimates were performed using R version 4.0.0 (R Foundation for Statistical Computing, Vienna, Austria). For all tests (except for heterogeneity), $P$ values of <0.05 were considered statistically significant.

3 | RESULTS

3.1 | Study selection and included studies

Our initial search identified 3030 articles after deduplication. Of these, 2928 were excluded after screening titles and abstracts, leaving 102 articles for full-text assessment (Figure 1). Eighteen studies met our inclusion and exclusion criteria, 32-49 of which two were excluded because of overlapping patient populations. 46,47 Another two studies were excluded because the family history of gastric and oesophageal cancer was combined and no individual data could be extracted. 48,49 The sensitivity analysis on full-text search of 1050 studies on BO and OAC did not reveal additional relevant articles on family history fulfilling the eligibility criteria (Figure S1). Agreement between investigators for assessment of study eligibility was good (kappa statistic = 0.79). 50 Our final data set included 14 unique articles: 9 addressed prevalence of familial BO/OAC, 4 addressed prevalence of BO in first-degree relatives and 7 addressed risk of having/developing BO or OAC. 52-45
Based on the QUIPS tool, a total of three studies were classified as low risk of bias.²⁹,⁴²,⁴⁵

3.3 Prevalence of familial BO and OAC

Nine studies including 1623 BO and 998 OAC patients assessed the prevalence of having at least one first-degree relative with BO or oesophageal cancer.²²,³³-³⁷,⁴⁰,⁴³-⁴⁵ A family history of BO or OAC was present in 1% to 10% of BO or OAC cases, except in one study,³³ in which a higher prevalence of familial BO (29%) and OAC (17%) was reported (Figure 2). The overall pooled prevalence of a positive family history in patients with BO (8.8%; 95% CI: 5.5-13.8; I² = 76%; six studies) was higher than in patients with OAC (4.4%; 95% CI: 2.2-8.7; I² = 75%; seven studies, P = 0.10). Only one study reported the prevalence of a family history of BO in patients with BO (prevalence: 17.1%; 95% CI: 8.1-32.7) and four studies reported the prevalence of a family history of oesophageal cancer in patients with OAC (pooled prevalence: 2.3%; 95% CI: 1.3-3.9; I² = 0%). A significant higher pooled prevalence of familial OAC was found in the three studies³³,³⁵,⁴⁵ that confirmed the positive family history (10.0%; 95% CI: 6.4-15.4; I² = 8%) compared with the four studies³²,³⁶,³⁷,⁴⁰ with no confirmation of family history (2.5%; 95% CI: 1.6-3.9; I² = 8%,

3.2 Study characteristics and quality assessment

Study characteristics and quality of included studies are summarised in Table 1; Figures S2 and S3, and Supporting Information File A. Most of the selected studies were cohort studies (n = 8) and all were performed in Western countries. Ten studies were performed in the United States³²-³⁸,⁴²-⁴⁴ and four in Europe.³⁹-⁴¹,⁴⁵ The index population consisted of BO (n = 3 studies), OAC (n = 5) and BO and OAC combined (n = 6). All diagnoses of the 16 189 included index patients were confirmed using medical records and pathology reports. In all studies including BO patients, BO was defined as >1 cm segment of salmon-coloured mucosa in the oesophagus combined with the presence of intestinal metaplasia in biopsies. The most common used definition of family history was having at least one first-degree relative with BO or oesophageal cancer. A small majority of studies used confirmed family histories by assessing pathology or medical reports of relatives.³³-³⁵,³⁸,³⁹,⁴¹,⁴²,⁴⁵ All other studies assessed family history using self-reports through surveys, interviews or medical reports. Of the seven studies addressing the risk of developing BO or OAC,³³,³⁶,³⁷,³⁹,⁴⁰,⁴²,⁴³ five studies controlled for age of persons at risk,³⁶,³⁹,⁴⁰,⁴²,⁴³ and three for gender of persons at risk.³⁹,⁴⁰,⁴² None of the studies adjusted for family size or reported whether relatedness of study participants was addressed.

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The prevalence for familial BO in the three studies that verified a positive family history was also higher than in the three studies that did not verify family history (10.6%; 95% CI: 5.7-26.6; $I^2 = 90$% vs 7.3%; 95% CI: 5.6-9.3; $I^2 = 0$%), but this difference was not statistically significant ($P = 0.48$).

The study by Chak et al.23 was identified as statistical outlier in the analyses on the prevalence of familial BO. Excluding this study, however, did not result in a significantly decreased pooled prevalence of having a positive family history in BO patients (6.8%; 95% CI: 5.7-8.2; $P = 0.31$), but heterogeneity decreased to 0%.

### 3.4 Characteristics of familial BO and OAC

Of the 14 included articles, five studies addressed differences between patients with familial BO/OAC and sporadic cases. No differences in ethnicity, smoking, alcohol consumption and obesity or body mass index were found. Nonetheless, one study including 26 OAC patients showed a significantly lower proportion of males in the familial OAC group, whereas the remaining studies found no differences in gender between sporadic and familial BO/OAC. Two studies reported a lower age at BO or OAC diagnosis for patients with familial BO/OAC compared with sporadic BO (mean age: 58 vs 64 years). This was not found in three other studies. One study showed that familial BO/OAC patients were younger at onset of heartburn compared with patients without a positive family history.

### 3.5 Prevalence of BO in first-degree relatives

The prevalence of BO in patients with a family history of BO or OAC was assessed in four studies in which relatives of familial BO probands underwent (endoscopic) screening to identify new BO cases. As these studies all used different study populations and screening methods, we judged that the risk estimates were too heterogeneous to provide a pooled estimate for the prevalence of BO and related neoplasia in those with a family history of BO.
FIGURE 2 Forest plot: proportion of patients with (A) Barrett's oesophagus (BO) or (B) oesophageal adenocarcinoma (OAC) having a family history (FH) of BO or oesophageal cancer (EC). *Not included in the pooled analyses of the total group.
family history definition (RR 3.64; 95% CI: 2.57-5.14; \( I^2 = 14\% \); \( P < 0.001 \)). Sensitivity analyses were performed to investigate potential sources of heterogeneity (Table 3). Restricting analyses to studies that accounted for GERD and body mass index, \( I^2 = 84\% \); \( P = 0.48 \).

3.8 | Subgroup analyses

None of the studies considered the effect of demographic characteristics (such as gender and age at diagnosis) on the prevalence of a positive family history or on the risk of BO and OAC. We could not perform subgroup analyses according to the number and age of affected relatives, and the degree of relatedness, as not enough studies provided data on BO or OAC risks in non-first-degree relatives. Three studies assessed the number of affected relatives in

### TABLE 2 Prevalence of Barrett’s oesophagus (BO) and oesophageal adenocarcinoma (OAC) in individuals with a first-degree relative with confirmed BO or OAC

| Author   | Screening population | Characteristics of first-degree relatives | FDRs screened | Diagnoses | BO length (cm) | Prevalence of BO/OAC in FDRs |
|----------|----------------------|-------------------------------------------|--------------|-----------|----------------|-------------------------------|
| Chak     | Only 1 affected FDR  | Mean: 44.6 Male: 43% 77% 43%            | 35           | BO: 2|OAC: 0 LSBO: n = 0 | 5.7% (95% Cl: 1.6-18.6) |
|          | \( \geq 2 \) affected FDRs | Mean: 45.3 Male: 55% 78% 30%           | 27           | BO: 10|OAC: 1 LSBO: n = 5 | 40.7% (95% Cl: 24.5-59.3) |
| Juhasz   | Asymptomatic FDRs   | Mean: 44.4 Male: 61% 0% 0%             | 12           | BO: 1|OAC: 0 LSBO: n = 4 | 8.3% (95% Cl: 1.5-35.4) |
|          | Symptomatic FDRs    | Mean: 44.4 Male: 61% 0% 0%             | 35           | BO: 12|OAC: 0 | 34.3% (95% Cl: 20.8-50.9) |
| Mussetto | FDRs with reflux    | Mean: 52 Male: 44% 100% 0%             | 18           | BO: 8|OAC: 0 Mean: 1.3 | 44.4% (95% Cl: 24.0-67.0) |
| Romero   | FDRs with reflux    | Mean: 52 Male: 44% 100% 0%             | 18           | BO: 8|OAC: 0 Mean: 1.3 | 44.4% (95% Cl: 24.0-67.0) |

Abbreviations: BO, Barrett’s oesophagus; OAC, oesophageal adenocarcinoma; FDR, first-degree relative; LSBO, long-segment Barrett’s oesophagus (BO segment \( \geq 3 \) cm).

\( * \) Obesity was defined as a body mass index \( \geq 27.8 \) kg/m\(^2\) in males and \( \geq 27.3 \) kg/m\(^2\) in females.
familial BO/OAC families. In familial BO/OAC, 10%\(^3\) and 48%\(^3\) of first-degree relatives were reported to be affected.

4 | DISCUSSION

This systematic review and meta-analysis provides evidence that confirmed familial clustering is seen in 10% of individuals with BO and OAC. Individuals with a first-degree relative with BO or oesophageal cancer were three and two times more likely to have BO and OAC compared with individuals without a positive family history respectively. Interestingly, OAC risk almost doubled in individuals with a verified family history of BO and OAC. We showed that offering screening to first-degree relatives of patients with BO or OAC had a diagnostic yield between 12% and 44% for BO, which is higher than in the general population (~2%).\(^7\) However, the yield was low for HGD and OAC. The number of BO diagnoses increased in relatives with GERD symptoms and in individuals with ≥2 affected relatives according to two individual studies.

Hitherto, one systematic review assessed the association between BO risk and family history as well as other risk factors. However, this review devoted a short paragraph on family history and included only four studies on screening first-degree relatives of BO patients.\(^5\) The authors showed a pooled prevalence of BO in individuals with a positive family history of 24%. However, descriptive data on the study populations were not reported, although necessary when applying the results to a specific patient population and informing relatives about their risk. Furthermore, this prevalence is probably an overestimation, as studies were confounded by the inclusion of particularly motivated individuals, relatives with additional risk factors and multiple individuals from multiplex families.

Familial clustering of BO and OAC could imply a genetic predisposition to BO or OAC, but may equally be caused by a common...
environmental exposure in family members or a genetic susceptibility to recognised risk factors such as GERD and obesity. In the current meta-analysis, adjustment for GERD symptoms attenuated the risk of BO and OAC associated with a positive family history. This GERD-dependent effect of family history on BO and OAC was supported by two case-control studies, which suggested a familial predisposition for GERD in relatives of BO patients. No differences in other lifestyle factors between familial and non-familial cases were found. On the other hand, the reported early age of disease diagnosis in familial BO/OAC cases compared with sporadic cases might indicate that relatives share one or more inherited genetic mutations. This is further supported by multiple case reports and a study of inheritance patterns in 70 families, which suggested an autosomal dominant inheritance pattern in familial BO/OAC. Although the exact pathogenesis of BO and OAC still needs to be unravelled, current evidence is in favour of a genetic susceptibility underlying the observed familial clustering at least in families with multiple affected relatives. In patients with a less extensive family history, shared environmental factors (in particular GERD symptoms) may potentially play a more important role. The current meta-analysis showed that the risk estimate of positive family history was lower for OAC than for BO. Although most studies included in our systematic review considered familial BO and OAC to be part of the same genetic trait, familial OAC may at least to some extent be distinct from familial BO, as a previous study speculated about the existence of a non-BO pathway to OAC. It is also plausible that individuals with a family history of OAC may undergo upper endoscopy more often, leading to enhanced detection and

| Category                                      | Studies (n) | RR (95% CI) | P value | I² (%) | References |
|-----------------------------------------------|-------------|-------------|---------|--------|------------|
| **Barrett’s oesophagus**                      |             |             |         |        |            |
| All studies                                   | 3           | 3.26 (1.43-7.40) | 0.005   | 46     | 33,42,43   |
| Study design                                  |             |             |         |        |            |
| Case-control                                  | 1           | 8.08 (2.53-25.78) | <0.001  | NA     | 33         |
| Cohort                                        | 2           | 2.26 (1.12-4.56) | 0.02    | 0      | 42,43      |
| Study location                                |             |             |         |        |            |
| United States                                 | 3           | 3.26 (1.43-7.40) | 0.005   | 46     | 33,42,43   |
| Europe                                        | 0           | NA          | NA      | NA     |            |
| Verification of BO and OAC in relatives        |             |             |         |        |            |
| Confirmed family history                      | 2           | 3.76 (0.85-16.64) | 0.08    | 70     | 33,42      |
| Verbal family history (no confirmation)       | 1           | 2.63 (1.07-6.47) | 0.04    | NA     | 43         |
| Adjusted for age                              | 2           | 2.26 (1.12-4.56) | 0.02    | 0      | 42,43      |
| Adjusted for gender                           | 1           | 1.77 (0.57-5.47) | 0.32    | NA     | 42         |
| Adjusted for GERD                             | 2           | 2.26 (1.12-4.56) | 0.02    | 0      | 42,43      |
| Adjusted for obesity                          | 2           | 2.26 (1.12-4.56) | 0.02    | 0      | 42,43      |
| **Oesophageal adenocarcinoma**                |             |             |         |        |            |
| All studies                                   | 5           | 2.19 (1.14-4.21) | 0.02    | 48     | 33,36,37,39,40 |
| Study design                                  |             |             |         |        |            |
| Case-control                                  | 4           | 1.55 (0.80-3.00) | 0.19    | 0      | 33,36,37,40 |
| Cohort                                        | 1           | 3.60 (2.52-5.14) | <0.001  | NA     | 39         |
| Study location                                |             |             |         |        |            |
| United States                                 | 3           | 1.69 (0.57-4.97) | 0.34    | 29     | 33,36,37   |
| Europe                                        | 2           | 2.57 (1.06-6.24) | 0.04    | 64     | 39,40      |
| Verification of BO and OAC in relatives        |             |             |         |        |            |
| Confirmed family history                      | 2           | 3.64 (2.57-5.14) | <0.001  | 14     | 33,39      |
| Verbal family history (no confirmation)       | 3           | 1.17 (0.55-2.46) | 0.69    | 0      | 35,36,39   |
| Adjusted for age                              | 3           | 1.92 (0.80-4.63) | 0.15    | 71     | 36,39,40   |
| Adjusted for gender                           | 2           | 2.57 (1.06-6.24) | 0.04    | 64     | 39,40      |
| Adjusted for GERD                             | 1           | 1.40 (0.49-4.01) | 0.53    | NA     | 40         |
| Adjusted for obesity                          | 2           | 1.15 (0.52-2.56) | 0.73    | 0      | 36,40      |

Abbreviations: BO, Barrett’s oesophagus; GERD, gastro-oesophageal reflux disease; OAC, oesophageal adenocarcinoma; RR, relative risk.
eradicaiton of early neoplasia. Hence, a positive family history may have a stronger effect on OAC risk than shown in our meta-analysis.

The future of OAC prevention relies on early detection of BO or early-stage OAC in high-risk individuals, followed by surveillance and endoscopic treatment for (dysplastic) BO. Current guidelines recommend different screening approaches for individuals with a family history of BO or OAC, varying from no screening, considering screening in relatives who also have other risk factors, to screening all individuals with a positive family history. Our findings confirm that a verified positive family history is a strong risk factor for BO and OAC. However, the observed pooled risk estimate is lower than the assumed risk of 12 reported by Chak et al., which has become the key reference in most screening guidelines. In our meta-analysis, this study was actually identified as an outlier. Although verification of family history was performed, the study by Chak et al was assessed as moderate risk of bias as it was limited by a small sample size, including only patients from tertiary hospitals, and combining BO, OAC and gastro-oesophageal junction adenocarcinoma for study outcomes. Additionally, another study showed that the number of new BO cases identified in a screening program of asymptomatic individuals with one first-degree affected relative was comparable to the prevalence in the general population. Taken together, the results of this meta-analysis do, in our opinion, not support a strong recommendation to endoscopically screen all first-degree relatives of patients with confirmed BO or OAC, given the number of first-degree relatives involved, the relatively low risk of BO and OAC in these individuals, associated direct and indirect costs, and invasiveness and potential complications of upper endoscopy.

However, this review emphasises the need for gastroenterologists to be aware of familial clustering and highlights the importance of obtaining a careful family history in all patients with BO or OAC. Although this systematic review could not conclusively quantify the higher risk of BO and OAC for individuals with ≥2 affected first-degree relatives, individual studies and multiple case reports have shown an increased risk of BO and OAC in families with multiple affected relatives, suggesting an underlying genetic aetiology. We believe that endoscopic screening of first-degree relatives should particularly be considered in families with ≥2 affected individuals. For individuals with a less extensive family history, endoscopic screening could be considered in first-degree relatives with multiple risk factors, such as age >50 years, male gender, obesity and GERD symptoms. For these individuals, minimally invasive screening options could also be considered.

The strengths of this analysis include a thorough systematic literature search according to a standardised protocol with well-defined inclusion criteria. We included all available studies without restricting analyses based on study design or language. All index patients had an endoscopic and histological confirmed BO or OAC diagnosis. Furthermore, a rigorous evaluation of study quality was performed, and two authors independently completed study selection and data extraction. Adjusted risk estimates were used to account for the effect of potential confounders. Finally, sensitivity analysis of between-study variation provided insight into data stability of pooled estimates and heterogeneity.

Some limitations of our study need, however, to be mentioned as well. First, although we performed the search strategy in accordance with previous meta-analyses on familial cancer risk, we did not assess the full text of all studies on BO and OAC, making it possible that studies that only reported the influence of a family history on BO or OAC as secondary outcome in the full text may have been missed. However, our sensitivity analysis and references of eligible articles and literature reviews did not reveal any additional articles. Second, results were derived from a combination of cohort and case-control studies with different study populations and heterogeneous nature of family history definitions, resulting in substantial heterogeneity in analyses. Most importantly, only eight studies verified the accuracy of self-reported data or used objective measures to confirm a positive family history. Furthermore, five studies only assessed family history of oesophageal cancer and did not distinguish between adenocarcinoma and squamous cell carcinoma of the oesophagus in relatives. Third, not all studies were adjusted for potential confounders, especially the effect of obesity, GERD and family size. Additionally, some studies combined BO and OAC for study outcomes and multivariable analyses could therefore not be included. Fourth, although reported family history is thought to be fairly accurate, controls may have been more likely to underreport their family history than BO or OAC patients (recall bias). Fifth, literature shows that the higher the number of affected relatives with cancer and the lower the age at cancer diagnosis in relatives, the greater the risks for an individual to develop cancer. Unfortunately, few studies reported on age at diagnosis of index patients or on the affected relatives or adjusted for number of family members. Also, data on BO and OAC risks for individuals with at least two affected first-degree relatives or second-degree relatives were limited and no study reported on the length of the BO segment in relatives. Hence, we were not able to assess BO or OAC risks according to the age of the individual at risk, age at diagnosis of the relative(s), degree of familial relation between the individual and relatives and number of affected relatives.

This review draws attention to the limited number of well-designed studies assessing a positive family history as a predictor of BO and OAC. Future longitudinal studies with well-defined family history criteria determining the risk of BO and OAC according to the number and age of affected relatives are needed to provide additional insight into the impact of family history on BO and neoplastic progression. Additionally, future research should focus on the underlying mechanisms of familial BO and identifying genetic risk factors. This could lead to more individualised screening, surveillance, prevention and treatment strategies for the clinical management of BO and OAC.

In conclusion, this systematic review and meta-analysis shows that familial aggregation is observed in a small but important subgroup of patients with BO and OAC. The currently available evidence identified a verified positive family history as a strong risk factor for BO and OAC. The review emphasises the importance of
obtaining a careful family history in all patients with BO or OAC. A confirmed family history of having at least two affected first-degree relatives and/or family history combined with other risk factors for BO and OAC can be used to identify individuals in which (endoscopic) screening might be considered to prevent OAC-related mortality.

ACKNOWLEDGEMENTS
We thank Ms OY Chan, biomedical information specialist, for optimising the literature search and dr R. Akkermans for the statistical advice.

Declaration of personal interest: Yonne Peters and Evi van Grinsven do not report any conflict of interest. Peter D. Siersema received an unrestricted grant from Pentax (Japan), Norgine (UK), Motus GI (USA) and The eNose Company (Netherlands).

AUTHORSHIP
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Author contributions: Yonne Peters was involved in study concept and design, data acquisition, quality control of data, data analysis and interpretation, statistical analysis, manuscript preparation, editing and review. Evi van Grinsven was involved in data acquisition, quality control of data, data analysis and critical revision of the manuscript. Peter D. Siersema was involved in study concept and design, data interpretation, critical revision of the manuscript and study supervision.

DATA AVAILABILITY STATEMENT
Data sharing not applicable to this article as no datasets were generated or analysed during the current study.

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