REVIEW



Genetic Biomarkers of Barrett's Esophagus Susceptibility and Progression to Dysplasia and Cancer: A Systematic Review and Meta-Analysis

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Abstract Barrett's esophagus (BE) is a common and important precursor lesion of esophageal adenocarcinoma (EAC). A third of patients with BE are asymptomatic, and our ability to predict the risk of progression of metaplasia to dysplasia and EAC (and therefore guide management) is limited. There is an urgent need for clinically useful biomarkers of susceptibility to both BE and risk of subsequent progression. This study aims to systematically identify, review, and meta-analyze genetic biomarkers reported to predict both. A systematic review of the PubMed and EMBASE databases was performed in May 2014. Study and evidence quality were appraised using the revised American Society of Clinical Oncology guidelines, and modified Recommendations for Tumor Marker Scores. Meta-analysis was performed for all markers assessed by more than one study. A total of 251 full-text articles were reviewed; 52 were included. A total of 33 germline markers of susceptibility were identified (level of evidence II-III); 17 were

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included. Five somatic markers of progression were identified; meta-analysis demonstrated significant associations for chromosomal instability (level of evidence II). One somatic marker of progression/relapse following photodynamic therapy was identified. However, a number of failings of methodology and reporting were identified. This is the first systematic review and meta-analysis to evaluate genetic biomarkers of BE susceptibility and risk of progression. While a number of limitations of study quality temper the utility of those markers identified, some—in particular, those identified by genome-wide association studies, and chromosomal instability for progression—appear plausible, although robust validation is required.

Keywords Barrett's esophagus · Esophageal cancer · Esophageal dysplasia · Genetic · Biomarkers

Introduction

The importance of Barrett's esophagus (BE) lies in its increasing prevalence and strong predisposition to esophageal adenocarcinoma (EAC) [1]. Western prevalence is estimated at 0.5-2.0 % [2] (a third of whom are asymptomatic [3]), conferring a 30- to 125-fold increased risk of developing EAC [4]. As EAC becomes more common [5], the benefits of predicting susceptibility to BE and detecting established metaplasia as soon as possible are numerous. Population screening overall is not presently feasible, but identifying both at-risk individuals and established metaplasia sooner provides windows for risk factor modification, chemoprevention, ablation, resection, and surveillance.

However, the natural history of BE is incompletely understood. Overall, the incidence of progression from metaplasia to high-grade dysplasia (HGD) or EAC is approximately 0.26–0.63 % per year [6], and that to EAC alone 0.12 % [7]. However, only a minority of patients progress to low-grade dysplasia (LGD), and of those that do similarly only a minority ultimately progress to HGD or EAC [8]. These progressive grades of dysplasia can help identify patients at particular risk (in particular, those with HGD in whom the risk is considerable) [9], but are insufficient in isolation.

There is therefore an immediate clinical need for biomarkers to predict both susceptibility to BE and progression. A major subtype is genetic variants, both germline and somatic. The potential of the latter, in particular, has been highlighted by a number of recent next-generation sequencing (NGS) studies of both EAC and BE, which have identified a number of candidate genes for further study [10]. However, no systematic reviews have been performed. The aims of this study were firstly to identify and evaluate all genetic markers tested in association with BE susceptibility and progression. Secondly, we aimed to identify markers with statistically significant associations and perform meta-analysis for those assessed by more than one study.

Methods

Inclusion Criteria

Studies testing associations between DNA markers [germline single nucleotide polymorphisms [SNP], somatic single nucleotide variants, insertions/deletions, copy number variants (CNV), loss of heterozygosity (LOH), microsatellite instability (MSI) or chromosomal instability (CIN)] and diagnosis/progression of BE were eligible; diagnosis was defined as endoscopic and histopathological evidence with or without intestinal metaplasia (IM) [13] and progression as histopathological progression from metaplasia to LGD/HGD/EAC or LGD to HGD/EAC, or HGD to EAC during surveillance endoscopies.

Exclusion Criteria

Studies comparing grades of metaplasia/dysplasia/malignancy within samples at one time point were excluded, as were studies comparing grades between patients. Studies using cell line, epigenetic, or expression data were excluded unless patient or DNA-specific data were available.

Literature Search

A search was performed in May 2014 of the PubMed and EMBASE databases, using the MOOSE and PRISMA guidelines [14] and the following term: [((((esophageal OR esophagus OR gastro esophageal)) AND (Barrett's OR metaplasia OR columnar)) AND (genomic OR genetic OR

genome OR pharmacogenetic OR pharmacogenomic OR amplification OR copy OR mutation OR polymorphism OR polymorphic OR variant OR deletion OR insertion OR locus OR loci OR allele OR ploidy OR instability OR biomarker))]. The bibliographies of retrieved articles were also searched.

Study Data

The following data were extracted: study methodology; variants and genes assessed; endpoints; population; and effect size [odds ratio (OR) or hazard ratio (HR)] and variance [standard error or confidence interval (CI)]. For studies not presenting OR, these were calculated using provided allele/variant frequencies. If reference SNP identification numbers were not provided (http://www.ncbi. nlm.nih.gov/dbSNP), these were mapped by searching cited methodology and performing in vitro polymerase chain reaction (http://genome.ucsc.edu), with nucleotide flank BLAST[®] (http://blast.ncbi.nlm.nih.gov).

Evidence Quality

Overall study and evidence quality were evaluated using the revised American Society of Clinical Oncology (ASCO) level of evidence (LOE) scale for biomarker research [15]. This stratifies study quality on the basis of trial design, patients and data, specimen collection, processing and archival, and statistical design and analysis from A to D (supplementary table 1), and uses this in conjunction with subsequent validation to stratify overall LOE for a marker from I to V (supplementary table 2). Methodological quality was appraised using the recommendations for tumor marker score (REMARK) guidelines modified by the authors, which scored study methodology in detail to generate a score from 0 to 17 (GWAS) and 0 to 18 (candidate studies; supplementary table 3). Reported associations were appraised for appropriate correction for multiple comparisons (Bonferroni method, false discovery rate, or multivariate analysis of all markers). If not undertaken, this was performed via post hoc Bonferroni correction. Significance was taken at $p < 5 \times 10^{-8}$ for genome-wide association studies (GWAS). Genomic quality criteria included reporting of genotyping call rate or providing data to allow its calculation, and assessment of Hardy-Weinberg equilibrium for germline variants.

Meta-Analysis

Meta-analysis was undertaken for markers assessed by more than one study, irrespective of correction for multiple comparisons (other than variants assessed by GWAS and non-GWAS due to major methodological differences) using RevMan v5.2 (Copenhagen: The Nordic Cochrane Centre, The Cochrane Collaboration). Sensitivity analyses were performed on the basis of IM-only versus non-IM studies. In the case of studies reporting updated results using cohorts previously reported (i.e., population overlap), the most recent study was used.

Heterogeneity and Bias

Heterogeneity was estimated using I^2 and Chi-square statistics. For moderate heterogeneity ($I^2 \ge 50$ %), a random-effects model was used. Funnel plots were reviewed for analyses of ≥ 5 studies [16]. Statistical assessment of these was not performed due to the low number of studies per variant. Publication bias was corrected using the "trim and fill" method [17].

Results

Literature Search

A total of 1516 articles were identified, of which 218 were duplicates. Therefore, 1298 articles were appraised on the basis of their title and abstract. Then, 251 full-text articles were retrieved, of which 52 met the inclusion criteria (Fig. 1).

Studies Assessing Germline Markers of Susceptibility

A total of 32 studies were included: 2 GWAS, 29 candidate studies, and 1 meta-analysis of 4 candidate studies (supplementary tables 4 and 5). All original studies were LOE C (indicating prospective observational registries, without standardized treatment and follow-up), published between 1999 and 2014. Median modified REMARK scores were 15/17 (GWAS) and 13/18 (candidate studies; range 7-16; supplementary table 6). A number of frequent methodological issues were identified. The most common of these included failure to blind investigators (n = 18;58.1 %), perform/report quality control procedures (19; 61.3 %), and match cases and controls (23; 74.2 %). Similarly, a number of recurrent reporting issues were identified. The most common of these were comparing markers with established risk factors (30; 96.8 %), adjusting for multiple comparisons (22; 71.0 %) and confounding factors (20; 64.5 %), reporting power calculations (15; 48.4 %) and reporting multivariate effect sizes (30; 96.8 %). Of the 27 studies assessing multiple variants, 24 did not perform multivariate analysis (88.9 %). For GWAS, there were a mean 5507 cases and 14,159 controls. For candidate studies, 134 cases and 196 controls. There were a number of cases of study population overlap,

with 13 studies reported on subjects drawn from one of three populations.

Variants Associated with BE Susceptibility

A total of 187 candidate variants/haplotypes were tested. Twenty-eight significant associations were reported, of which 16 were excluded (Table 2). Twelve were therefore associated with BE (Table 1). However, for 2 of these (rs6785049 and rs9344) precise p values were not provided to allow for Bonferroni correction.

The rs1695 (*GSTP1*) was assessed by 4 studies, which underwent meta-analysis by Bull et al. [18]. This calculated an OR of 1.50 (95 % CI 1.16–1.95 p = not presented; LOE II). While derived from a large total cohort (434 cases and 738 controls), none of the four studies adjusted for risk factors; indeed the sole study finding a significant association had only 22 cases [19]. The *GSTM1* null genotype was also assessed by 4 studies and underwent updated meta-analysis in this study. Overall, no association was demonstrated. On sensitivity analysis, a significant negative association was apparent for the 2 studies not requiring IM [17, 20], although the relevance of this is unclear. Meta-analysis was performed for 5 other variants, none of which demonstrated associations (Table 3).

Of the 12 significant candidate associations reported, only rs1695 (*GSTP1*) and rs25487 (*XRCC1*) were assessed by more than one study. This notwithstanding, 5 appear relatively robust on the basis of adjustment for clinical covariates. These include 3 growth factor variants: rs444903 (*EGFR* [20]; notably associated with reflux esophagitis and EAC), rs6214 (*IGF1* [21]), and rs2229765 (*IGF1R*) [22]. Two interleukin variants also appear plausible: rs3212227 (*IL12B*) and rs917997 (*IL18RAP*) [23–25], with the former demonstrated to be independent of all other tested genotypes [26]. A number of other associations were reported in the *IL1* [27, 28], *IL10* [28], *IL18* [29], and *IL23* [30] clusters. Of these, however, only wild-type rs917997 (*IL18RAP*) persisted following correction for multiple comparisons [29].

The remaining 5 candidate variants included associations with 3 caudal homeobox 1 (*CDX1*) variants: rs3776082, rs2237091, and rs717767. The authors demonstrated these variants to be significantly associated with established risk factors for BE: age, gender, and the presence of hiatus hernia. However, multivariate analysis was not performed to demonstrate whether the association of these variants with BE was independent of these. Of the remaining 2, an association was demonstrated for the rs6785049 (*NR112*) variant; however, similarly this was not adjusted for risk factors. This was performed for the rs9344 (*CCND1* [31]) variant, although the *p* value was not published.



Fig. 1 PRISMA diagram

Of the 5 GWAS variants identified, 4 were identified by the Wellcome Trust Centre Case Control Consortium (WTCCC). Two were identified by the initial report [32]: rs9257809 within the major histocompatibility complex (MHC; OR 1.21 [1.13–1.28]; $p = 4.09 \times 10^{-9}$) and rs9936833 (related to *FOXF1*; OR 1.14 [1.10–1.19]; $p = 2.74 \times 10^{-10}$). Subsequent replication identified rs3072 (related to *GDF7*: OR 1.14 [1.09–1.18]; $p = 1.80 \times 10^{-11}$) and rs2701108 (related to *TBX5*: OR 0.90 [0.86–0.93]; $p = 7.50 \times 10^{-9}$) [33]. All remained significant when meta-analyzed with data from the Barrett's and Esophageal Adenocarcinoma Consortium (BEACON) GWAS. The first 2 were also associated with EAC in the BEACON dataset and an independent candidate study [34]. Mechanistically, GWAS arrays rely on linkage disequilibrium between SNPs, by which SNPs act as proxies for others (i.e., genotyping one SNP allows the genotypes of others to be inferred with confidence). This "tagging", however, means that variants identified may be functional or may in fact be bystanders tagging other SNPs [35]. However, the loci overall and related genes are consistent with the roles of immune-mediated inflammation (MHC) and thoracoembryogenesis (*FOXF1*, *GDF7* and *TBX5*) in BE.

The BEACON GWAS identified 3 loci associated with either BE or EAC, although none reached the genome-wide

Table	: 1 Reported germline n	narkers of B	larrett's esophagus suscept	tibility									
Mark	er		Individual studies						Association (meta-analysis	s/indiv	idual	study)	
LOE	Variant	Gene	Association		EV	LOE	Ethnicity	Methodology					
			Variant	Wild-type Non	6				Phenotype and OR (95 % CI)	I^2	Chi	d	и
III	rs9257809	MHC region	Su et al. [32], Palles et al. [33]		A	C	Caucasian	Multistage GWAS Pro/retro archived	BE ^A 1.21 (1.13–1.28)	NA	NA	4.09×10^{-9}	-
	rs9936833	FOXF1			U				BE ^A 1.14 (1.10–1.19)			2.74×10^{-10}	
	rs3072	GDF7			A				BE ^A 1.14 (1.09–1.18)			1.75×10^{-11}	
	rs2701108	TBX5			Α				BE^{A} 0.90 (0.86–0.93)			7.48×10^{-9}	
	rs2687201	FOXP1			Н				BE ^A 1.16 (1.10–1.23)			4.61×10^{-8}	
Ш	rs3776082	CDXI	Ren et al. [43]		AA	С	Caucasian	Candidate	BE-v-RE 4.02 (1.80-9.00)	NA	NA	<0.005	-
								Prospective					
III	rs2237091	CDXI		Ren et al. [43]	AA	C	Caucasian	Candidate	BE-v-RE 0.28 (0.13-0.59)	NA	NA	<0.005	-
								Prospective					
Ш	rs717746	CDXI	Ren et al. [43]		GG	U	Caucasian	Candidate	BE-v-RE 3.65 (1.73–7.69	NA	NA	<0.005	-
								Prospective					
Ш	rs444903	EGFR	Menke et al. [20]		66	C	Caucasian*	Candidate	BE ^A 3.00 (1.50–6.20)	NA	NA	0.002	-
								NCILU ALCIILVOU					
III	rs917997	IL 18RAP		Babar et al. [29]	20	C	Caucasian**	Candidate Retro archived	BE^{A} 0.59 (0.43–0.80)	NA	NA	6.00×10^{-4}	-
III	rs6785049	PXR	van de Winkel et al. [44]		IJ	C	Caucasian*	Candidate	BE 1.36 (1.03–1.79)	NA	NA	<0.05 (NS)	
								Retro archived					
III	rs6214	IGFI		McElholm	AA	C	Caucasian**	Candidate	BE ^A 0.43 (0.24–0.75)	NA	NA	3.00×10^{-3}	-
				[21]				Retro archived					
III	rs2229765 + BMI > 30	IGFIR	MacDonald et al. [22]		A	C	Caucasian	Candidate	BE ^A 3.11 (1.12–8.63)	NA	NA	<0.05 (NS)	-
								Retro archived					
Π	rs3212227	IL12B	Moons et al. [26]		U	C	Caucasian*	Candidate	BE ^A 1.82 (1.17–2.69)	NA	NA	7.00×10^{-3}	-
								Retro archived					

Mark	er		Individual studies							Association (meta-analysis/indiv	vidual s	tudy)		
LOE	Variant	Gene	Association			EV	LOE	Ethnicity	Methodology					
			Variant	Wild-type	None					Phenotype and OR (95 % CI)	I^2	Chi	d	и
п	rs1695	GSTP	Bull et al. [18]			G	С	Caucasian	Meta-analysis	BE 1.50 (1.16–1.95)	NA	NA	<0.01	4
			Kala et al. [19]			IJ		Caucasian	Candidate (pro)					
					Murphy et al. [45]	IJ		Caucasian	Candidate (retro)					
					Casson et al. [46]	IJ		NP	Candidate (pro)					
					van Lieshout et al. [47]	IJ		Caucasian	Candidate (pro)					
Π	rs9344	CCNDI	Casson et al. [31]			$\mathbf{A}\mathbf{A}$	C	NP	Candidate (pro)	BE^{A} 3.69 (1.46–9.29)	NA	NA	<0.05 (NS)	-
Π	rs25487	XRCCI	Casson et al. [48]		Ferguson et al. [49]	AA	C	NP	Candidate (pro)	BE ^A 0.65 (0.25–1.68)	71 %	3.42	0.370	7
						AA	C	Caucasian	Candidate (retro)					

denote overlapping populations

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threshold for BE alone. However, on subsequent metaanalysis of both GWAS, rs2687201 (*FOXP1*, similarly involved in developmental regulation) was significantly associated with BE alone, in addition to a further variant associated with either BE/EAC [33].

Studies Assessing Germline Variants Associated with Progression

One study was identified (supplementary table 7). LOE was C; modified REMARK score was 13.5/19. This assessed 4 variants in the *IGF* axis; none were associated with progression.

Studies Assessing Somatic Variants Associated with Progression

Sixteen studies were identified, published between 1989 and 2012 (supplementary table 8). LOE was C for 12 and D for 4. Mean modified REMARK score was 13.2 (range 10.5–18.5/19; supplementary table 9). Five classified progression as HGD/EAC, 10 as EAC, and 1 as EAC/CIN. Again, a number of recurrent methodological issues were identified. These included failure to: blind investigators (12; 75.0%), perform appropriate quality control/reproducibility (9; 56.3%), match controls (14; 87.5%), and appropriate power calculations (14; 87.5%). Recurrent reporting issues included failure to: report univariate association effects (9; 60.0%), adjust for risk factors (particularly the presence of dysplasia at baseline; 14; 87.5%), and fully report coefficients of multivariate models (13; 81.3%).

Variants Associated with Progression

Of 7 variants assessed, 5 associations were identified: CIN, CNV (>70 Mbp), *TP53* LOH, *p16* LOH, and mutant *TP53* (Table 4). Meta-analysis was possible for CIN, which was assessed by 11 studies. These defined CIN variably as aneuploidy (4), tetraploidy (1), and aneuploidy/tetraploidy (6). Six of these studies were derived from independent sample archives. However, this was not clear for 4. All studies reported associations of CIN with progression. However, only two adjusted for confounding variables, including the presence of dysplasia [36, 37]. Another two, while not adjusting for dysplasia, did adjust for length of Barrett's segment [38, 39], with CIN remaining significant.

Meta-analysis was performed for both OR and HR of progression to HGD/EAC. Significant associations were demonstrated for both (Table 4; Fig. 2). All studies included patients with IM only. Meta-analyzed OR was 5.98 (2.10–17.1; $p = 8.00 \times 10^{-4}$; n = 5 studies; following exclusion of overlapping studies and correction for publication bias). However, only one of these studies adjusted

	Mark	er		Individual studie	Ş					Association (meta-analysi	s/indiv	idual s	tudy)	I I
Number Num Num Num	LOE	Variant	Gene	Association		EV	LOE	Ethnicity	Methodology					
				Variant	Wild-type	I				Phenotype and OR (95 % CI)	I^2	Chi ,	~	и
	III	rs3776083	CDXI	Ren et al. [43]		AG	С	Caucasian	Candidate pro	BE-v-RE 1.91 (1.09–3.33)	NA	NA (0.02	
	Ш	rs4769585	CDX2	Ren et al. [43]		CT	C	Caucasian	Candidate pro	BE-v-RE 2.68 (1.20–5.98)	NA	NA (0.02	-
	Π	rs3812863	CDX2	Ren et al. [43]		AG	C	Caucasian	Candidate pro	BE-v-RE 2.53 (1.24–5.14)	NA	NA (0.01	-
II VTRs LL Mode Eadoreds+Holia 1/2 Causaian (Czeck) Candidate pro BE 105 (1.02-1.39) NA NA NA NA NA 11 s169347 LLM LLM Eadoreds+Holia CT/ Causaian (Czeck) Candidate pro BE 105 (1.02-1.39) NA N	Ш	rs1143634	ILIB		Izakovicoka-Holla et al. [27]	H		Caucasian (Czeck)	Candidate pro	BE -v-RE 0.56 (0.33-0.93)	NA	NA (0.016	-
	III	VNTR polymorphism	ILIRN		Izakovicoka-Holla et al. [27]	1/2		Caucasian (Czeck)	Candidate pro	BE 0.62 (0.42–0.94)	NA	NA (0.015	-
III $rs18005871$ $LLBV$ $radvicoka+hola$ T/C $cueasian$ (Czeck) $cai (15)-6.24$) N	Π	rs1800587/ rs16944/ rs1143634/ IL1RN	IL IB/ ILRN		Izakovicoka-Holla et al. [27]	C/T/ C/long		Caucasian (Czeck)	Candidate pro	BE 1.95 (1.09–1.30)	NA	AN AN).03	
III rs909233 TVFB Menke et al. [50] AA C Caucasian Candidate retro BF 1 (10-7.93) NA NA NA Odd III rs2305764 Myo9b Menke et al. [51] GG C Caucasian Candidate retro BF 2 2.96 (1.10-7.93) NA NA 0.03 III rs194518 IL/8 Babar et al. [29] CC Caucasian (USh) Candidate retro BF 2 2.96 (1.10-7.198) NA NA 0.03 III rs194518 IL/8 Babar et al. [29] CC Caucasian (USh) Candidate retro BF 2 2.45 (1.07-1.98) NA NA 0.03 III rs11209026 IL/3 Bradbury et al. GG C Caucasian (USh) Candidate retro BF 2 2.45 (1.07-1.98) NA NA 0.03 III rs11209026 IL/3 GG et al. [30] A Caucasian (USh) Candidate retro BF 2 2.43 (1.37-4.93) NA NA 0.01 III rs11209026 IL/3 Gg	Ш	rs1800587/ rs16944/r s1143634/IL1RN	IL IB/ ILRN		Izakovicoka-Holla et al. [27]	T/C/ C/long		Caucasian (Czeck)	Candidate pro	BE 2.67 (1.15–6.24)	NA	NA AN	0.02	-
III rs2305764 $Myo9b$ Menke et al. [51] GG C Caucasian Candidate retro BE ^A 2.96 (1.10-7.99) NA NA 0.03 III rs194518 $IL/8$ Babar et al. [29] CC Caucasian (Irish) achived BE ^A 1.45 (1.07-1.98) NA NA 0.03 III rs179976 $MMPI$ Bradbury et al. GG Caucasian (Irish) candidate retro BE ^A 1.45 (1.07-1.98) NA NA 0.03 III rs179976 $MMPI$ Bradbury et al. GG Caucasian (Irish) candidate retro BE ^A 1.45 (1.07-1.98) NA NA 0.03 III rs179976 $MMPI$ Bradbury et al. GG Caucasian (USA) candidate retro BE ^A 1.45 (1.07-1.98) NA NA 0.03 III rs1120926 $IL23R$ Gaj et al. [30] A Caucasian (USA) candidate retro BE ^A 1.45 (1.07-1.98) NA NA 0.01 III rs1120926 $IL23R$ Gaj et al. [30] T NP	III	rs909253	TNFB	Menke et al. [50]		AA	C	Caucasian (Netherlands)	Candidate retro archived	BE ^A 1.98 (1.03–3.81)	NA	NA (0.04	
III $rs1946518$ $IL/8$ Babar et al. [29] CC Caucasian (Irish) Candidate retro archived BE^A 1.45 (1.07–1.98) NA NA NA 0.01 III rs1799750 MMPI Bradbury et al. GG Caucasian (USA) Candidate retro BE^A 1.45 (1.07–1.98) NA NA 0.01 III rs1799750 MMPI Bradbury et al. GG Caucasian (USA) Candidate retro BE^A 2.21 (1.08–4.53) NA NA 0.01 III rs11209026 IL23K Gaj et al. [30] A Caucasians (Polish) Candidate pro BE^A 2.21 (1.08–4.53) NA NA 0.01 III rs11209026 IL23K Gaj et al. [30] A Caucasians (Polish) Candidate pro BE^A 2.21 (1.08–4.53) NA NA 0.016 III rs1800566 NOU di Martino et al. T NP (UK) Candidate pro BE^A 2.14 (1.22–372 NA NA NA 0.016 III rs415958 ILIAA Gough et al. [28] <t< td=""><td>III</td><td>rs2305764</td><td>Myo9b</td><td>Menke et al. [51]</td><td></td><td>GG</td><td>C</td><td>Caucasian (Netherlands)</td><td>Candidate retro archived</td><td>BE^A 2.96 (1.10–7.99)</td><td>NA</td><td>NA (</td><td>0.032</td><td>-</td></t<>	III	rs2305764	Myo9b	Menke et al. [51]		GG	C	Caucasian (Netherlands)	Candidate retro archived	BE ^A 2.96 (1.10–7.99)	NA	NA (0.032	-
III $rs1799750$ $MMPI$ Bradbury et al. GG Caucasian (USA) Candidate retro BF^{A} 2.21 (1.08-4.53) NA NA NA 0.03 III $rs11209026$ $IL23R$ Gaj et al. [30] A Caucasians (Polish) Candidate pro BE 3.49 (1.37-8.90) NA NA NA NA 0.01 III $rs11209026$ $IL23R$ Gaj et al. [30] A Caucasians (Polish) Candidate pro BE 3.49 (1.37-8.90) NA NA NA 0.01 III $rs1800566$ $NOOI$ di Martino et al. T NP (UK) Candidate pro BE^{A} -v.FB 0.22 NA NA NA 0.016 III $B7$ HLA Rajendra [54] B7 Asian (India) Candidate pro BE^{Ex} 2.1.4 (1.22-372 NA NA 0.016 $rs16944$ ILIRAV Gough et al. [28] S(UK) Candidate pro BE^{Ex} 2.1.4 (1.22-372 NA NA 0.016 $rs16944$ ILI Rajendra [28] S(UK) Candidate pro BE^{Ex} 2.1.4 (1.22-372) NA NA 0.016 rs1644	III	rs1946518	IL18	Babar et al. [29]		CC		Caucasian (Irish)	Candidate retro archived	BE ^A 1.45 (1.07–1.98)	NA	NA (0.02	
$ \begin{array}{ c c c c c c c c c c c c c c c c c c c$	III	rs1799750	IdWW	Bradbury et al. [52]		GG		Caucasian (USA)	Candidate retro archived	BE ^A 2.21 (1.08–4.53)	NA	NA (0.03	-
$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$	III	rs11209026	IL23R	Gaj et al. [30]		А		Caucasians (Polish)	Candidate pro	BE 3.49 (1.37–8.90)	NA	NA	011	-
III B7 HLA Rajendra [54] B7 Asian (India) Candidate pro BE^{EX} 21.4 (1.22–372 NA NA 0.036 III rs415958/ ILIRA/ Gough et al. [28] CC/A NS (UK) Candidate pro BE-v-RE 9.5 (1.19–75.9) NA NA 0.010 rs16944 ILIB Cough et al. [28] CC/A NS (UK) Candidate pro BE-v-RE 9.5 (1.19–75.9) NA NA 0.010 rs16944 ILIB COUR CC/A NS (UK) Candidate pro BE-v-RE 1.84 NA NA 0.035 III rs1800896 ILIO Gough et al. [28] CC NS (UK) Candidate pro BE-v-RE 1.84 NA NA 0.035 III rs1800896 ILIO Gough et al. [28] CC NS (UK) Candidate pro BE-v-RE 1.84 NA NA 0.035	III	rs1800566	ΙΟÕΝ	di Martino et al. [53]		T		NP (UK)	Candidate pro	BE^{A} -v-RE 0.22 (0.07–0.76)	NA	NA	0.016	
III rs415958/ ILIRA/ Gough et al. [28] NA NA NA NA 0.010 rs16944 ILIB ILIB ILIB Candidate pro BE-v-RE 9.5 (1.19–75.9) NA NA 0.010 rs16944 ILIB ILIB Candidate pro BE-v-RE 9.5 (1.19–75.9) NA NA 0.010 rs16944 ILIB 0.010 Cough et al. [28] CC NS (UK) Candidate pro BE-v-RE 1.84 NA NA 0.035 III rs1800896 ILI0 Gough et al. [28] CC NS (UK) Candidate pro BE-v-RE 1.84 NA NA 0.035	III	B7	HLA	Rajendra [54]		B7		Asian (India)	Candidate pro	BE ^{Ex} 21.4 (1.22–372	NA	NA	0.036	-
III rs1800896 IL10 Gough et al. [28] CC NS (UK) Candidate pro BE-v-RE 1.84 NA 0.035 (1.04-3.28) (1.04-3.28) (1.04-3.28) (1.04-3.28) (1.04-3.28)	Ш	rs415958/ rs16944	ILIRA/ ILIB	Gough et al. [28]		CC/A		NS (UK)	Candidate pro	BE-v-RE 9.5 (1.19–75.9)	NA	NA (0.010	
	Π	rs1800896	1110	Gough et al. [28]		CC		NS (UK)	Candidate pro	BE-v-RE 1.84 (1.04–3.28)	NA	NA).035	

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Marker			Individu	al study		Meta-analysis				
			Associat	ion		-				
Variant	Gene	EV	Variant	Wild-type	No association	OR (95 % CI)	I^2	Chi	р	n
Null	GSTM1	Null	Null		Kadioglu et al. [55]	BE 0.83 (0.61-1.12)	27 %	4.12	0.220	4
					Kala et al. [19]	BE ^{no IM} 0.66 (0.44-0.99)	0 %	0.64	0.420	2
					Casson et al. [46]	BE ^{IM} 1.12 (0.70–1.80)	0 %	0.64	0.630	2
					van Lieshout et al. [56]					
rs4880	SOD2	Т	Т		Kadioglu et al. [55],	BE ^{IM} 0.90 (0.65-1.24)	0.00	0.07	0.520	3
					di Martino et al. [49]					
					Murphy et al. [45]					
rs16944	IL1B	AA	AA		Izakovicova-Holla et al. [27],	BE 1.13 (0.63-2.03)	0.00	0.00	0.680	2
					Gough et al. [28]					
rs1052133	OGG1	G	G		Kadioglu et al. [55],	BE ^{A IM} 1.34 (0.59–3.09)	67 %	3.01	0.480	2
					Ferguson et al. [49]					
rs25487	XRCC1	TT	TT	Casson et al. [48]	Ferguson et al. [49]	BE ^{A IM} 0.65 (0.25–1.68)	71 %	3.42	0.370	2
rs13181	ERCC2	CC	CC		Casson et al. [48],	BE ^{A IM} 0.92 (0.57–1.48)	31 %	1.44	0.730	2
					Fergusons et al. [49]					

Table 3 Meta-analyzed markers assessed in association with BE susceptibility

EV effect variant, OR odds ratio, CI confidence interval, IM intestinal metaplasia, BE Barrett's esophagus

for the presence of dysplasia. Meta-analyzed HR was 1.36 (1.26–1.47; n = 2 studies; $p < 1.00 \times 10^{-5}$; following exclusion of one overlapping study). Overall LOE for CIN was II. While more than one study assessed mutant *TP53* (n = 3) and LOH *TP53* (n = 2), meta-analysis was not possible for either. Two studies for each were derived from the same populations, and the third *TP53* study used a different measure of effect size.

Variants Associated with Risk of Progression Following Photodynamic Therapy

Three studies were identified (supplementary table 10). LOE was C for 2 and D for 1. Mean modified REMARK score was 16.2/20 (range 14.5–19; supplementary table 7). For 2 studies, while endoscopic mucosal resection was variably performed, this was only controlled for in 1 study. A total of 6 variants were assessed; while 5 were assessed by 2 studies, meta-analysis could not be performed due to population overlap. CIN at both 4 and 12 months was reported to be associated with risk of progression (Table 5).

Discussion

We believe this review to be the first to identify, synthesize, and evaluate the evidence for genetic markers of BE susceptibility and risk of progression. Thirty-three susceptibility markers were identified; however, just 17 remained significant after correcting for multiple comparisons. Five (rs9257809, rs9936833, and subsequently rs3072, rs2701108, and rs2687201) were derived from GWAS and are therefore most likely to be reproducible. Of the 6 candidate markers assessed by more than one study, meta-analysis was supportive for one (rs1695, *GSTP1*). Five non-meta-analyzed variants affecting either growth factors or inflammatory cytokines appear plausible and therefore represent priorities for validation: rs444903 (*EGFR*), rs6214 (*IGF1*), rs2229765 (*IGF1R*), rs3212227 (*IL12B*), and rs917997 (*IL18RAP*).

No germline markers of progression risk were identified. However, 5 somatic markers were reported, plus another of progression following photodynamic therapy. Meta-analysis was possible for CIN, demonstrating significant effects for both HR and OR after correction for publication bias. However, there was considerable heterogeneity regarding definition of CIN, duration, and frequency of follow-up, confounding risk factors (e.g., prevalence of HGD) in addition to minimum follow-up periods. Notably, however, both studies adjusting for HGD [36, 37] did demonstrated convincing effects.

The robustness of the associations between CIN and progression suggests it to be of immediate clinical utility. CIN is a constituent of genomic instability, a state of erroneous progression through the cell cycle. Inaccurate DNA replication, repair, and chromosomal segregation, results in accumulation of genomic errors and is a major factor driving tumorigenesis. CIN is associated with worse stage and prognosis in a range of

Marke	ar		Individual study							Association (meta-analysis/indiv	vidual st	(dpn)		
LOE	Variant	Gene	Association		EV	LOE	Ethnicity	Phenotype	Adjusted for					
			Variant	None					contounders?	Phenotype and OR/HR (95 % CI)	I^2	Chi	d	и
П	CIN	NA	Bird-Lieberman et al. [36]		CIN	C	NP	HGD/ EAC	Yes (plus length, dysplasia)	HGD/EAC OR ^{(A) IM PB} 5.98 (2.10–17.1)	57 %	16.2	8.00×10^{-4}	2
			Sikkema et al. [37]		CIN	C	Cauc	HGD/ EAC	Yes (plus dysplasia)	HGD/EAC HR ^{(A) IM} 1.36 (1.26–1.47)	0 %	0.98	$<1.00 \times 10^{-5}$	7
			Chao et al. [57]*		CIN	C	NP	EAC	No					
			Galipeau et al. [38]*		CIN	C	NP	EAC	Yes (length, not dysplasia)	PB-adjusted for publication bias Fang 2004, Rabinovitch				
			Fang et al. [58]		CIN	C	NP	EAC	No	2001,				
			Maley et al. [39]*		CIN	C	NP	EAC	Yes (length, not dysplasia)	Teodori 1999				
			Rabinovitch et al. [59]*		CIN	C	NP	EAC	Yes (length, not dysplasia)					
			Reid et al. [9]**		CIN	D	NP	EAC	No	Overlapping populations***				
			Teodori et al. [60]		CIN	U	NP	HGD/ EAC	No					
			Reid et al. [8]**		CIN	C	NP	EAC	No					
			James et al. [61]		CIN	D	NP	EAC	No					
Ш	CNV	NA	Paulson et al. [62]		>70 Mbp	C	dN	EAC/CIN	No	EAC/CIN HR 4.90 (1.60-14.8)	NA	NA	4.70×10^{-3} (NS)	1
III	НОЛ	TP53	Galipeau et al. [38]*		НОП	C	NP	EAC	Yes (length, not dysplasia)	EAC HR ^A 5.4 (2.5–12.0)	NA	NA	$<1 \times 10^{-3}$	-
			Maley et al. [39]*		НОТ	C								
			Reid et al. [63]*		НОН	C								
III	НОН	P16	Galipeau et al. [38]*	Maley et al. [39]*	НОН	C	NP	EAC	Yes (length, not dysplasia)	EAC ^A HR 2.4 (1.0–5.5)	NA	NA	0.045	-
					НОН	U	NP	EAC	Yes (length, not dysplasia)					
III	Mutant	TP53		Galipeau et al. [38]*	Mutant	U	NP	EAC	No	EAC HR ^A 1.27 (1.07–1.50)	NA	NA	<0.01 (NS)	-
			Maley et al. [39]*		Mutant	C	dN	EAC	Yes (length, not dysplasia)					

Marker	Individual study						Association (meta-analysis/individual study)
LOE Variant Gene	Association	EV	LOE	Ethnicity	Phenotype	Adjusted for	
	Variant None	1				confounders?	Phenotype and OR/HR I^2 Chi p n (95 % CI)
	Dolan et al. [64]	Mutant	C	NP	EAC	No	No meta possible as study population overlap and differing effect measures
LOE level of evidence, high-grade dysplasia, E	<i>EV</i> effect variant, <i>OR</i> odds ratio <i>AC</i> esophageal adenocarcinoma,	, <i>HR</i> hazard r A adjusted	atio, <i>CI</i>	confidence	interval, CIN	' chromosomal inst	ability, NA not applicable, NP not presented, NS nonsignificant, HGD

,* denote overlapping populations



Fig. 2 Uncorrected and corrected funnel plots for odds ratio of progression and chromosomal instability

tumors including esophageal [40] and has been demonstrated in a quarter of patients with BE [36]. Importantly, this subset appears to be at significantly higher risk of malignant progression, which can be readily demonstrated by flow cytometry. This does, however, remain somewhat imprecise and is unable to distinguish between stable and unstable, simple and complex abnormalities. Despite this, the incorporation of CIN into a biomarker panel such as that reported by Bird-Lieberman et al. [36] (comprising age, CIN, dysplasia, TP53 and Cyclin A expression, sialyl Lewis antigens, Aspergillus oryzae lectin, and binding of wheat germ agglutinin) may provide invaluable information with which to personalize management of BE.

By contrast, the immediate benefits of germline susceptibility biomarkers are less tangible. As predicted by the "common disease-common variant" hypothesis of complex traits, both GWAS suggested many common variants of small effect to contribute to development of BE. Consequently, germline associations may be weaker and more complex. Whilst this gives the potential to identify novel biology, variants may have little utility in isolation.

A number of recurrent methodological issues were identified, limiting the generalizability of reported variants.

Association (meta-analysis/individual study)

Pable 5 Reported somatic biomarkers of Barrett's esophagus progression to dysplasia/adenocarcinoma following photodynamic therapy

Individual study

Marker

	и	1		re.a
	d	9.00×10^{-3}	2.90×10^{-3}	a EAC esonhao
	Chi	NA		solacis
	I^2	NA		ade dv
	Phenotype HR (95 % CI)	EAC/HGD HR ^A 4.10 (1.30–13.0)	EAC/HGD HR ^A 3.60 (1.05–12.3)	otodynamic therany HGD high-m
Adjusted for	contounders?	Yes		interval DDT nh
Phenotype		HGD		I confidence i
Ethnicity		NS		ard ratio C
LOE		D		HR har
EV		CIN 4 m post-PDT	CIN 12 m post-PDT	tv EV effect variant H
	None			nstahilit
Association	Variant	Dunn et al. [65]		CIN chromosomal i
Gene		NA		dence
Variant		CIN		level of evi
LOE		IV		IOF

adenocarcinoma

Ultimately, validation studies for prioritized variants should be designed with these in mind, with particular emphasis on the interaction between genomic and clinical factors. Other issues to be addressed include disparity as diagnostic criteria; confirmation of IM is a prerequisite in the USA, yet is not required in the UK [41], although this did not alter the findings of this review.

A number of exploratory NGS studies have recently been performed for both BE and EAC. These have served to highlight the mutational complexity of both conditions, providing biological context for markers and their genes (which are often considered in isolation) as well as suggesting new variants and genes for study. Recently, Streppel et al. [11] performed whole genome sequencing (WGS) of one patient, comparing normal squamous epithelium with metaplastic and neoplastic epithelium. This identified somatic nonsense mutations in genes including AT-rich interactive domain 1A (ARID1A), a member of the SWI/SNF family involved in gene expression via chromatin remodeling, which has been independently identified as a driver gene of EAC by of other NGS studies. The authors found ARID1A loss of expression to become progressively more common during the metaplasia-dysplasia-adenocarcinoma sequence, and to be associated with aberrant cellular proliferation and invasion in a knock-down model. NGS studies also provide valuable contextual information as to mutational spectra, as well as clonal and linear evolution to better understand to development of somatic mutations and genomic instability. Recently, Weaver et al. [12] performed WGS of 112 EACs, similarly identifying a number of significantly and recurrently mutated genes. One hundred and seven BE samples were then genotyped, with the notable finding that most such mutations were already present in non-dysplastic epithelium; just TP53 and SMAD4 mutations occurred later, in HGD and EAC. The advent of thirdgeneration sequencing, for example from single cells, will undoubtedly shed yet further light on this process. Additionally, germline rather than somatic mutations in EAC driver genes have been shown to predispose to EAC, although this has yet to be demonstrated in BE [42].

This review has a number of limitations. While we searched two databases using a comprehensive search term, it is possible that relevant publications (including non-English articles) were not identified. As discussed, meta-analysis for CIN was performed within the context of considerably heterogeneity and must be interpreted with caution. Only one meta-analysis (CIN and OR of progression) involved more than 5 studies, with considerable funnel plot asymmetry, largely due to small studies reporting large effect sizes with significant variance. This was interpreted as publication bias, and was adjusted, without altering statistical significance. In addition, the

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"trim and fill" method used to correct for possible bias while widely used does make assumptions regarding the necessity for plot symmetry, while not incorporating study methodology. Unfortunately, the limited number of studies prevented useful meta-regression to assess this further. There are also limitations in using the revised ASCO guidelines, as these do not fully represent the complexity of methodological quality and also disagreement between studies. We therefore used modified REMARK guidelines to provide a further level of criticism. In particular, the ASCO guidelines do not allow for differences in methodology (for example, GWAS versus candidate studies). And more generically, while genomic biomarkers are typically considered in isolation, in reality their utility depends on innumerable variables (including transcriptional, translational and proteomic regulation, and clinical and environmental factors). Consequently, establishing their true utility will require parallel processing and consideration of these contexts.

In conclusion, this review has identified, evaluated, and synthesized the evidence for genomic biomarkers of BE susceptibility and dysplastic/malignant progression. Seventeen germline markers of susceptibility, 5 somatic markers of progression, and 1 marker of relapse following photodynamic therapy were identified. Metaanalysis demonstrated CIN to be a particularly plausible and clinically useful marker of progression and one which can be demonstrated readily. However, the overall evidence base is characterized by widespread methodological issues, which limit the immediate clinical utility of these markers. Consequently, larger studies with more robust design are required to validate these markers, identify novel variants, and incorporate them into clinical practice.

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Compliance with ethical standards

Conflict of interest Mark R. Middleton has received payment for advisory/consulting roles within the last 2 years from Amgen, Bristol-Myers Squibb, GlaxoSmithKline, Merck, Millennium, and Roche, and institutional funding from Amgen, AstraZeneca, Bristol-Myers Squibb, Clovis, Eisai, GlaxoSmithKline, Immunocore, Johnson & Johnson, Merck, Millennium, Novartis, Pfizer, Roche and Vertex. The authors declare no conflicts of interest.

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