REVIEW



Identifying Novel Biomarkers Ready for Evaluation in Low-Prevalence Populations for the Early Detection of Upper Gastrointestinal Cancers: A Systematic Review

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ABSTRACT

Introduction: Detecting upper gastrointestinal (GI) cancers in primary care is challenging, as cancer symptoms are common, often non-specific, and most patients presenting with these symptoms will not have cancer. Substantial investment has been made to develop biomarkers for cancer detection, but few have reached routine clinical practice. We aimed to

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M. Messenger Leeds Centre for Personalised Medicine and Health, University of Leeds, Leeds, UK identify novel biomarkers for upper GI cancers which have been sufficiently validated to be ready for evaluation in low-prevalence populations.

Methods: We systematically searched MED-LINE, Embase, Emcare, and Web of Science for studies published in English from January 2000 to October 2019 (PROSPERO registration CRD42020165005). Reference lists of included studies were assessed. Studies had to report on second measures of diagnostic performance (beyond discovery phase) for biomarkers (single or in panels) used to detect pancreatic, oesophageal, gastric, and biliary tract cancers. We included all designs and excluded studies with less than 50 cases/controls. Data were extracted on types of biomarkers, populations and outcomes. Heterogeneity prevented pooling of outcomes.

Results: We identified 149 eligible studies, involving 22,264 cancer cases and 49,474 controls. A total of 431 biomarkers were identified (183 microRNAs and other RNAs, 79 autoantibodies and other immunological markers, 119 other proteins, 36 metabolic markers, 6 circulating tumour DNA and 8 other). Over half (n = 231) were reported in pancreatic cancer studies. Only 35 biomarkers had been investigated in at least two studies, with reported outcomes for that individual marker for the same tumour type. Apolipoproteins (apoAII-AT and apoAII-ATQ), and pepsinogens (PGI and

PGII) were the most promising biomarkers for pancreatic and gastric cancer, respectively.

Conclusion: Most novel biomarkers for the early detection of upper GI cancers are still at an early stage of matureness. Further evidence is needed on biomarker performance in low-prevalence populations, in addition to implementation and health economic studies, before extensive adoption into clinical practice can be recommended

Keywords: Biomarkers; Clinical practice; Early detection; Primary care; Upper gastrointestinal cancers

Key Summary Points

We aimed to identify novel biomarkers which had been validated and showed sufficient promise to warrant further evaluation in low-prevalence populations.

We identified 431 unique biomarkers; only 35 of which had been investigated in at least two studies, with outcomes for that individual marker for the same tumour type - four of these were identified as the most promising for future studies.

This review highlights the need for more biomarker studies that consider primary care/community settings as their intended populations.

Findings also indicate we still need better reporting to facilitate knowledge translation; we also need more consistency in the use of biomarkers.

Research collaborations are vital to reduce duplicate efforts and ensure appropriate samples sizes when studying lowprevalence populations.

DIGITAL FEATURES

This article is published with digital features, including a summary slide, to facilitate understanding of the article. To view digital features for this article go to https://doi.org/10.6084/m9.figshare.13214843.

INTRODUCTION

Gastrointestinal (GI) cancers represented more than 25% (4.8 million) of cancer cases and over a third (3.4 million) of cancer-related deaths worldwide in 2018 [1]. Upper GI cancers contribute an important proportion of these, with over 2.1 million new cases of cancers of the stomach, oesophagus, pancreas and biliary tract diagnosed worldwide in 2018 [1, 2]. Prognosis is often poor as upper GI cancers are generally not detected until the disease is advanced and less amenable to curative treatment [1].

Primary care plays a key role in the early detection of upper GI cancers, as more than 90% of patients present with symptoms [3–5], and screening tests for asymptomatic populations are not yet widely established. Early detection of upper GI cancers is challenging, as initial symptoms such as indigestion, abdominal discomfort or fatigue are common, often intermittent, and most patients presenting with them will not have cancer [6, 7].

There is growing demand to improve early cancer detection through better diagnostic and triage approaches, particularly for use in primary care or other community settings where cancer prevalence is low [5]. New diagnostic approaches, applied either among asymptomatic at-risk populations or to triage patients presenting with cancer symptoms, could be transformational. Electronic health records and large population-based surveys have been used to develop cancer risk prediction models to

identify those requiring investigation for cancer [8]; diagnostic pathways have also been implemented in different countries in an effort to improve timely cancer diagnosis [5]. Innovative strategies applying artificial intelligence techniques to imaging and other medical data are also promising [5, 9]. For cancers with nonspecific symptom signatures, like most upper GI cancers, we also need better biomarkers to support diagnostic assessment [10]. Biomarkers such as carcinoembryonic antigen (CEA) and CA19-9 are used in clinical practice predominantly for surveillance following treatment of upper GI cancers [9, 11]. Substantial investment has been made into developing new biomarkers for early cancer detection; most such biomarker research has been conducted in laboratory and specialist clinical settings [12, 13], where cancer prevalence is higher compared to community settings [14, 15].

The distinction between care settings is important, as the diagnostic performance characteristics of a test are strongly determined by the prevalence and severity of the target disease and of other diseases within the study population [14]. In populations in which the prevalence of the target disease is low (e.g. primary care), positive predictive values are lower than in high-prevalence populations seen in specialist cancer centres. Tests evaluated in highprevalence populations tend to have lower sensitivity and higher specificity when used in low-prevalence populations [15, 16]. This is known as the spectrum effect or spectrum bias [14, 15] and has crucial implications for translating results from one care setting to another. To gain an accurate understanding of how a test will perform within a low incidence setting, it must ultimately be evaluated within that setting.

In recognition of this, the CanTest Framework has been developed, proposing a 5-phase translational pathway for diagnostic tests, from new test development to health system implementation in low-prevalence populations [15]. The framework highlights the importance of evaluating not only clinical performance but also the feasibility and acceptability of implementation, patient safety and quality of care, and cost-effectiveness in the chosen clinical

setting. Understanding and addressing these issues is vital, as test performance alone, even if evaluated in the target populations, does not guarantee clinical utility nor improved patient outcomes [12].

This review set out to systematically identify novel biomarkers for the early detection of upper GI cancers which have been validated and show sufficient promise to warrant further evaluation in low-prevalence populations.

METHODS

Search Strategy and Inclusion/Exclusion Criteria

This systematic review followed the Preferred Reporting Items for Systematic Reviews and Meta-Analysis (PRISMA) guidelines [17], and the was registered in **PROSPERO** (CRD42020165005). We searched MEDLINE, Embase, Emcare and Web of Science from 1 January 2000 to 31 October 2019 for primary studies published in English. The search strategy (Online supplementary file 1) was developed with the assistance of a medical librarian and refined until it identified all relevant core publications known by the senior authors. Reference lists of included studies were also screened. Articles that were not available online were ordered via the British Library.

Studies were included if they reported on at least one measure of diagnostic performance: sensitivity, specificity, positive predictive value (PPV), negative predictive value (NPV), false positive, false negative or area under the curve (AUC) for biomarkers used to detect oesophageal, gastric, pancreatic or biliary tract cancers. We included adult populations (mean/median age \geq 18); we accepted individuals aged < 18 if these were outliers in large samples. The search strategy also included terms for lower GI (colorectal and anal) cancers for the purposes of a parallel review of novel biomarkers for the early detection of lower GI cancers, to be reported separately. Non-specified GI cancers, neuroendocrine cancers and studies only reporting on familial populations at risk of hereditary cancers were excluded.

Novel biomarkers were considered both individually and as part of a combination/panel test. Studies reporting only the performance of a single, established biomarker (i.e. CEA and CA19-9 for pancreatic cancer) were not eligible for inclusion [9]. We included studies reporting on performance for established biomarkers if these were in combination with additional novel biomarkers.

We aimed to identify studies situated within Phase 2 (measures of diagnostic accuracy in high-prevalence settings) and Phase 3 of the CanTest framework (measures of diagnostic accuracy or clinical utility, acceptability and feasibility in intended low-prevalence settings) (Fig. 1) [15]. We included studies if they reported more than preliminary measures of performance calculated in a discovery phase; this required additional measures of diagnostic performance in an independent cohort. If no ref-

erences to previous studies evaluating performance were available and the study provided only one set of measures, the study was excluded. Panels with previously investigated biomarkers were included even if the biomarkers had not been investigated as part of a panel. As larger sample sizes are required beyond the biomarker discovery phase [13, 18], studies had to include at least 50 cancer cases and at least one group of 50 non-cancer controls with similar clinical characteristics (healthy, or with non-malignant or pre-malignant conditions). Similar criterion has been adopted by previous reviews that informed our study [13, 19].

We only included biomarkers which are feasible to use in a community setting, i.e. blood (serum and plasma), urine, faecal, salivary or breath samples. Observational studies (cross-sectional or longitudinal, prospective or retrospective) and trials were eligible for inclusion.

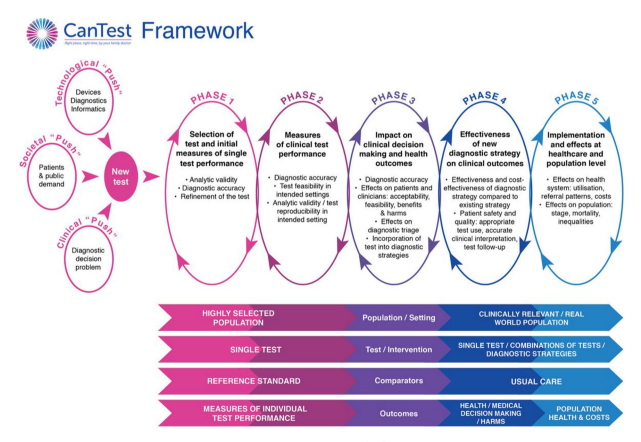


Fig. 1 The CanTest Framework Reproduced with permission from [15]

We included all recruitment settings, as we expected that very few studies would have been carried out in community settings.

We used the online tool Covidence [20] to facilitate title and abstract screening and study selection. Two reviewers (any two of NC, PED, CS, KMM, DB or RB) independently screened titles and abstracts. Then, two reviewers (any two of the above) independently evaluated full-text articles for inclusion. Titles and abstracts of reference lists of included studies were reviewed by one author (NC); full-text articles selected at this stage were independently assessed by two reviewers (any two of NC, PED, RB or DB). Disagreements were resolved by consensus; when this could not be reached a senior, third reviewer (FMW or JE) was consulted.

Data Extraction and Analysis

Data extraction was piloted to ensure consistency and was carried out by one of seven reviewers (NC, PED, RB, DB, JMG, JO and SS). We extracted information on: study characteristics (publication year, country of population of interest, recruitment setting, study aims and design); populations (numbers included, age, sex, tumour staging for cases and health status for controls); biomarkers (type of sample, biomarker name, biomarker category); and summary measures of diagnostic performance (sensitivity, specificity, PPV, NPV, false positives, false negatives and AUC, with 95% confidence intervals when available, for all comparisons). When studies reported on different phases of biomarker development, we only extracted data from the eligible phases (i.e. biomarkers and measures beyond the discovery phase). When studies had more than one eligible phase, we extracted data from all phases. Extracted data were collated and checked for consistency and inaccuracies (NC).

Biomarkers were categorised according to a modified version of Uttley et al.'s classification [19], which included: microRNAs and other RNAs, autoantibodies and other immunological

markers, other proteins (that did not fit into other categories), metabolic markers, circulating tumour DNA, and other biomarkers. Controls were classified as: normal/healthy, having non-malignant, or pre-malignant conditions. Biomarkers and control populations were coded by one author (NC) and checked by other authors (PED, KMM and MM; and PED, FMW and JE, respectively). Controls described as being healthy were coded as such unless studies described underlying conditions. Patients with cancer were ineligible as controls. Full details of the classification of controls are available (online supplementary table S1). Microsoft Excel 2015 and SPSS v.23 (IBM) were used for data extraction and data analysis.

Quality Assessment and Risk of Bias

Risk of bias [21] was not assessed as described in the original protocol, following independent piloting. Appraisal was hindered by the use of diverse methods across studies and incomplete reporting, resulting in a large number of "unclear" assessments. Instead, a list of issues identified in the studies was prepared (Online supplementary file 2). As spectrum bias is a key issue when translating results from high- to lowprevalence populations, all included studies were classified as either single-gate or two-gate designs. In single-gate designs, cases and controls are recruited through a single route of entry and with the same inclusion criteria (e.g. all cases and controls presented with symptoms). In two-gate designs, participants are recruited through different routes and different inclusion criteria exist for cases and controls. In this situation, controls can be either normal/ healthy or with an alternative diagnosis, which can produce symptoms and signs similar to patients with cancer [16]. One author (NC) classified all studies and another (PED) checked the classification. A full description of this classification and how it approaches some of the issues covered by the critical appraisal tool is available (Online supplementary file 3).

Data Synthesis

Included studies were heterogeneous and rarely evaluated the same biomarkers in the same way, often using different cut-off points, populations and/or biomarker combinations in panels. Therefore, we were unable to undertake meta-analysis. Instead, we used narrative synthesis to summarise data across studies [22]. First, we developed an overview of the available evidence, describing key characteristics of included studies, their populations and biomarkers, and outcome measures. Then, we looked for similarities that would allow for subgroup analyses, namely the same biomarker, for the same tumour type, with similar designs, outcome measures and populations.

Compliance with Ethics Guidelines

This article is based on previously conducted studies and does not contain any studies with human participants or animals performed by any of the authors.

RESULTS

Database searches identified 16,597 records; 9172 were retained after removing duplicates. During title and abstract screening, 8179 ineligible records were excluded. The full texts of the remaining 993 records were assessed for eligibility; 731 were excluded (Fig. 2). A total of 262 studies from database searches met inclusion criteria; 25 additional studies were identified in reference lists. Of these, 149 included studies referred to upper GI cancers and were included in our narrative synthesis.

Characteristics of Included Studies

Key characteristics of included studies are described in Table 1 and 2. Most studies recruited participants from a single country (n = 142). China was the most common country (n = 77), followed by Japan and South Korea (n = 15 each), the USA (n = 12) and Germany (n = 9). The most common recruitment settings were

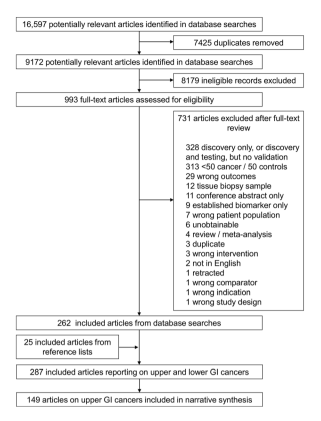


Fig. 2 Study selection

hospital or other secondary care institutions (n = 125), biobanks, reference sets, databases or archived samples (n = 20), general population cohorts or cohorts from population screening programmes (n = 11) and cohorts from previous trials or observational studies (n = 9). Several studies recruited from more than one setting. Gastric cancer was the most commonly investigated tumour type (n = 69), followed by pancreatic (n = 54), oesophageal (n = 24) and biliary tract cancers (n = 3). Four studies investigated more than one type of upper GI cancer (Table 1).

Characteristics of Cases and Controls

Overall, the included studies reported on 22,264 cancer cases (10,589 gastric, 7964 pancreatic, 3258 oesophageal and 290 biliary tract cancers, and 163 oesophago-gastric cancers, not

Table 1 Characteristics of included studies: country, setting and population

Hosp Other Other Auto-	References	Country (normlation)	Settinga		Cases and controls	alo:			
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South Korea X - 147 94 Ub Ub China X - 110 110 10 0 China X - 90 57 57 0 Mongolia X - 50 57 0 752 0 752 Mongolia X - 149 56 187 57 0 752 0 752 0 187 187 187 187 187 187 187 187 187 187 188 189 189 189 189 189 189 189 189 189 189	Chen et al. [25]	China	×	1	87	105	40	65	0
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Mongolia x - 50 752 65 752 South Korea U U 96 187 0 187 China x - 149 74 74 111 China x - 197 74 74 111 China x 6 60 59 59 6 China x - 60 60 60 60 60 China x - 54 54 54 6 6 China x - 60 60 60 60 6 China x - 54 54 54 6 6 China x - 106 55 106 108 108 South Korea x - 106 52 120 108 109 Aban x x x x 109	Dong et al. [28]	China	×	I	06	57	57	0	0
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Japan x - 54 54 54 0 China x - 168 74 74 0 China x - 106 358 160 198 Japan x - 187 561 561 0 Japan x - 380 626 228 291 Japan x - 122 178 79 99 South Korea x - 120 178 179 179 179 Russia - x x 52 104 104 0	Huang et al. [35]	China	×	ı	09	09	09	0	0
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China X - 106 358 160 198 Japan X - 187 561 561 0 Japan X - 380 626 228 291 Japan X - 122 178 79 99 South Korea X - 120 170	Ji et al. [37]	China	×	1	168	74	74	0	0
Japan x - 187 561 561 0 South Korea x - 380 626 228 291 Japan x - 122 178 79 99 South Korea x - 120 Ub Ub Ub Russia - x 52 104 104 0	Juan Cai et al. [38]	China	×	1	106	358	160	198	0
South Korea × - 380 626 228 291 Japan × - 122 178 79 99 South Korea × - 120 120 120 10 10 Russia - × 52 104 104 0	Kaise et al. [39]	Japan	×	ı	187	561	561	0	0
Japan X - 122 178 79 99 South Korea X - 120 Ub Ub Ub Russia - X 52 104 104 0	Kang et al. [40]	South Korea	×	ı	380	979	228	291	107
South Korea \times – 120 U^b U^b U^b Russia – \times 52 104 104 0	Kikuchi et al. [41]	Japan	×	ı	122	178	42	66	0
Russia – \times 52 104 104 0	Kim et al. [42]	South Korea	×	I	120	120	$\Omega_{ m p}$	$\Omega_{ m p}$	0
	Kurilovich et al. [43]	Russia	I	×	52	104	104	0	0

continued
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Table

References	Country (population)	Setting ^a		Cases and controls	rols			
				Cases (N)	Controls (N)	ols (N)		
		Hosp	Other		All	НС	NM	PM
Li et al. [44]	China	×	ı	09	09	09	0	0
Li et al. [45]	China	×	1	62	112	81	0	31
Li et al. [46]	China	×	ı	65	9	99	0	0
Li et al. [47]	South Korea	×	ı	100	100	100	0	0
Li et al. [48]	China	×	ı	234	428	270	0	158
Lim et al. [49]	South Korea	×	ı	100	06	$\Omega_{ m p}$	Ω^{b}	30
Lim et al. [50]	South Korea	×	1	100	100	$\Gamma_{\rm p}$	$\Gamma_{\rm p}$	30
Lin et al. [51]	China	D	Ŋ	51	78	09	18	0
Liu et al. [52]	China	×	ı	142	105	105	0	0
Liu et al. [53]	China	×	1	119	148	66	49	0
Liu et al. [54]	China	×	1	50	90	50	0	0
Meistere et al. [55]	Taiwan, Latvia, Lithuania, Germany	×	×	829	929	929	0	0
Mroczko et al. [56]	Poland	×	1	73	61	61	0	0
Ning et al. [57]	China	×	I	169	75	75	0	0
Oue et al. [58]	Japan	×	I	123	96	92	20	0
Pan et al. [59]	China	×	I	81	130	77	53	0
Park et al. [60]	South Korea	×	1	81	103	32	63	∞
Parvace et al. [61]	Iran	ı	×	50	20	50	0	0
Qin et al. [62]	China	×	×	407	407	407	0	0
Qiu et al. [63]	China	×	I	200	200	200	0	0
Song et al. [64]	China	ı	×	89	89	0	89	0
Su et al. [65]	China	×		82	65	50	6	0

Table 1 continued

References	Country (population)	Setting ^a		Cases and controls	rols			
				Cases (N)	Controls (N)	(N) sl		
		Hosp	Other		All	HC	NM	PM
Sun et al. [66]	China	×	×	332	332	332	0	0
Tsalikidis et al. [67]	Greece	×	ı	66	78	78	0	0
Wang et al. [68]	Taiwan	D	D	170	116	116	0	0
Wang et al. [69]	China	×	ı	72	54	54	0	0
Wang et al. [70]	China	×	×	186	186	186	0	0
Wang et al. [71]	China	×	I	09	120	09	09	0
Werner et al. [72]	Germany	I	×	146	26	26	0	0
Wu et al. [73]	China	×	ı	06	06	06	0	0
Wu et al. [74]	China	×	I	66	132	100	30	2
Wu et al. [75]	China	×	ı	201	318	157	161	0
Yanaoka et al. [76]	Japan	I	×	63	5146	5146	0	0
Yang et al. [77]	South Korea	I	×	290	290	290	0	0
Yang et al. [78]	China	×	1	109	106	0	106	0
Yoon et al. [79]	South Korea	×	×	200	200	200	0	0
Yun et al. [80]	China	×	I	194	376	185	191	0
Zayakin et al. [81]	Latvia, Germany	×	ı	235	367	213	154	0
Zhang et al. [82]	China	×	ı	114	298	187	111	0
Zhang et al. [83]	China	×	×	80	70	0	70	0
Zhang et al. [84]	China	×	ı	80	80	0	80	0
Zhou et al. [85]	China	×	1	50	20	$\Omega_{ m p}$	$\Omega_{\rm p}$	$\Omega_{\rm p}$
Zhou et al. [86]	China	×	ı	71	61	61	0	0
Zhou et al. [87]	China	×	1	70	70	70	0	0

Table 1 continued								
	Country (population)	Setting ^a		Cases and controls	80			
				Cases (N)	Controls (N)	Is (N)		
		Hosp	Other		All	HC NM	NM	PN

References	Country (population)	Setting ^a		Cases and controls	rols			
				Cases (N)	Controls (N)	[S (N)		
		Hosp	Other		All	НС	NM	PM
Pancreatic cancer only								
Akita et al. [88]	Japan	×	ı	116	138	138	0	0
Balasenthil et al. [89]	USA	I	×	86	154	61	93	0
Brand et al. [90]	USA	×	ı	173	120	120	0	0
Cao et al. [91]	China	×	I	156	115	0	57	28
Capello et al. [92]	USA	×	I	73	134	09	74	0
Chung et al. [93]	South Korea	×	I	55	93	70	23	0
Chung et al. [94]	South Korea	×	I	54	80	55	25	0
Deng et al. [95]	China	×	ı	303	640	009	40	0
Duraker et al. [96]	Turkey	×	I	123	173	0	173	0
Firpo et al. [97]	USA	×	×	75	261	150	84	27
Fukutake et al. [98]	Japan	×	I	240	7800	7772	78	0
Gao et al. [99]	China	×	I	70	120	50	20	0
Gold et al. [100]	USA	I	×	53	130	43	82	0
Gold et al. [101]	USA	×	×	298	199	42	120	0
Groblewska et al. [102]	Poland	Ω	Ω	62	99	65	0	0
Guo et al. [103]	China	×	ı	250	300	150	150	0
Honda et al. [104]	Japan, Germany	×	I	319	291	181	110	0
Honda et al. [105]	Japan, USA	×	×	384	342	192	150	0
Honda et al. [106]	Ten European countries ^c	I	×	156	213	213	0	0
Jiang et al. [107]	China	×	I	96	252	200	52	0
Kaur et al. [108]	USA	×	ı	154	167	0	167	0

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Kim et al. [109] USA X X Controls (N) All HC NM Kiwarani et al. [109] USA X X X 418 220 83 Kowarani et al. [110] Japan X X X 418 220 83 Lec et al. [112] Cach Republic Slovakia X X 397 533 374 159 Lec et al. [113] Cach Republic Slovakia X X 397 533 374 159 Liu et al. [114] Cach Republic Slovakia X X 397 533 374 159 Liu et al. [114] China X X 58 146 102 112 Mascubar et al. [114] China X X 23 470 240 230 Mascubar et al. [118] Germany X X 134 57 54 57 Milloy et al. [121] Japan X X 130 136 34 37 <th>References</th> <th>Country (population)</th> <th>Setting^a</th> <th></th> <th>Cases and controls</th> <th>rols</th> <th></th> <th></th> <th></th>	References	Country (population)	Setting ^a		Cases and controls	rols			
Hosp Other All Hosp Hosp Other					Cases (N)	Contro	ols (N)		
111] USA × × 418 220 Japan × × 98 158 105 Japan × × 98 158 105 China × × 58 146 102 China × - 138 175 68 China × - 172 215 133 China × - 172 215 140 Japan × - 140 97 87 Japan × - 140 97 87 Japan × - 143 276 219 Japan × - 143 276 219 South Korea - × 143 146 74 South Korea - × 143 146 74 South Korea - × 129 146 74 Carada × × 146 96 99 Carada × ×			Hosp	Other		All	НС	NM	PM
Japan X - 98 158 105 Cacch Republic, Slovakia X X 397 533 374 South Korea X - 51 112 0 China X - 138 146 102 China X - 172 68 102 China - X - 175 68 102 China - X - 172 68 103 103 104 1	Kim et al. [109]	USA	×	×	278	418	220	83	115
III] Czech Republic, Slovakia x x 397 533 374 South Korea x - 51 112 0 Taiwan x - 58 146 102 China x - 138 175 68 China x - 172 215 133 China x - 172 240 240 China x - 140 97 87 Japan x - 140 97 87 Japan x - 140 97 87 Japan x x 143 184 184 South Korea - x 191 184 184 South Korea y x 263 185 185 USA x x x x x x x x Japan x x x <td>Kuwatani et al. [110]</td> <td>Japan</td> <td>×</td> <td>ı</td> <td>86</td> <td>158</td> <td>105</td> <td>21</td> <td>32</td>	Kuwatani et al. [110]	Japan	×	ı	86	158	105	21	32
South Korea x - 51 112 0 Taiwan x x 146 102 China x - 138 146 102 China x - 172 215 133 China - x 235 470 240 240 Japan - x 140 97 87 240 Japan - x 79 160 80 80 Japan - x 143 276 219 84 184 184 184 Japan - x 180 184 184 184 184 184 South Korea - x 139 146 74 185 44 South Korea - x 139 146 74 184 184 South Korea - x x x 184 x 184 184 Canada x x x x x x x	LeCalvez-Kelm et al. [111]	Czech Republic, Slovakia	×	×	397	533	374	159	0
Taiwan x x 146 102 China x - 138 146 102 China x - 175 68 133 China x - 175 140 240 240 Japan x - 140 97 87 87 Denmark, USA - x 143 276 219 87 Japan x x 143 276 219 87 146 74 South Korea - x 101 184	Lee et al. [112]	South Korea	×	ı	51	112	0	112	0
China × - 138 175 68 China × - 172 215 133 China - × 235 470 240 Japan × - 140 97 87 Denmark, USA - × 143 276 219 Japan × - 180 184 184 UK - × 101 184 184 South Korea - × 139 146 74 South Korea - × 139 146 74 South Korea 0 0 0 292 165 94 Taiwan × × × 84 99 99 Cacch Republic × × × × 18 48 Japan × × × 84 99 99 Cacch Republic × × ×<	Liao et al. [113]	Taiwan	×	×	58	146	102	4	0
China × - 172 215 133 China - × 235 470 240 Japan × - 140 97 87 Denmark, USA - × 143 276 219 Japan - × 143 276 219 UK - × 143 276 219 South Korea - × 139 146 74 South Korea - × 139 146 74 South Korea - × 139 146 74 Vanada - × 263 136 185 Vanada - × 84 99 99 Czech Republic × - 64 185 48 Japan - × 84 99 99 Vanada - - 64 185 48 Japan - - 53 147 102 Vanada -	Liu et al. [114]	China	×	I	138	175	89	107	0
China - × 235 470 240 Japan - - 140 97 87 Denmark, USA - × 143 160 80 Japan - × 143 276 219 UK - × 101 184 84 184 South Korea - × 101 184 184 184 South Korea - × 101 184 184 184 South Korea - × 129 146 74 USA × × 263 230 185 Canada × × 86 134 86 Cacch Republic × × 64 185 48 Japan × × × 147 102 USA × × × 84 99 99 Japan × × ×	Liu et al. [115]	China	×	ı	172	215	133	82	0
Japan x - 140 97 87 Germany - x 79 160 80 Denmark, USA - x 143 276 219 Japan x - 180 180 84 UK - x 139 146 74 South Korea - x 139 146 74 South Korea - x 165 94 74 Taiwan x x 263 230 185 94 USA x x 86 134 86 99 99 Cach Republic x x x 64 185 48 Japan x x x 147 102 Germany x x 186 220 89 48 x x x 86 99 99 Assam x x	Liu et al. [116]	China	I	×	235	470	240	230	0
Germany - × 79 160 80 Denmark, USA - × 143 276 219 Japan × - 180 84 219 UK - × 101 184 184 184 South Korea - × 139 146 74 74 Taiwan X X 263 230 185 94 USA X X 84 99 99 Canada - X 84 99 99 USA X X 84 99 99 Japan X X 185 48 USA X X 187 102 USA X X 185 48 USA X X 185 48 USA X X 147 102 USA X X X 18 243 128 USA X X X X 243	Matsubara et al. [117]	Japan	×	ı	140	26	87	0	10
Denmark, USA - × 143 276 219 Japan × - 180 84 84 UK - × 101 184 184 184 South Korea - × 139 146 74 74 South Korea U U 292 165 94 74 185 94 94 USA × × × 86 134 86 185 98 Cacch Republic × × 84 99 99 99 Japan × - 53 147 102 USA - × 188 220 89 Germany × × 166 243 128	Mayerle et al. [118]	Germany	I	×	42	160	80	80	0
Japan x - 180 84 UK - x 101 184 184 South Korea - x 139 146 74 Taiwan U U 292 165 94 USA x x 86 134 86 Canada - x 84 99 99 Czech Republic x - 64 185 48 Japan x - 53 147 102 USA - x 188 220 89 Germany x x 116 243 128	Mellby et al. [119]	Denmark, USA	I	×	143	276	219	57	0
UK - × 101 184 184 184 South Korea - × 139 146 74 South Korea U U 292 165 94 Taiwan × × 263 185 185 USA × × 84 99 99 Czech Republic × - 64 185 48 Japan × - 64 185 48 USA - × 188 220 89 Germany × × 116 243 128	Mizuno et al. [120]	Japan	×	ı	180	180	84	96	0
South Korea - × 139 146 74 South Korea U U 292 165 94 Taiwan × × 263 135 185 185 USA × × 84 99 99 99 Canada - × 84 99 99 99 Japan × - 64 185 48 Japan × - 53 147 102 USA - × 188 220 89 Germany × × 116 243 128	O'Brien et al. [121]	UK	I	×	101	184	184	0	0
South Korea U U U D P <th< td=""><td>Park et al. [122]</td><td>South Korea</td><td>I</td><td>×</td><td>139</td><td>146</td><td>74</td><td>72</td><td>0</td></th<>	Park et al. [122]	South Korea	I	×	139	146	74	72	0
Taiwan x x 263 185 USA x x 134 86 Canada - x 84 99 99 Czech Republic x - 64 185 48 Japan x - 53 147 102 USA - x 188 220 89 Germany x x 116 243 128	Park et al. [123]	South Korea	Ω	Ŋ	292	165	94	71	0
USA X X 86 134 86 Canada - X 84 99 99 Czech Republic X - 64 185 48 Japan X - 53 147 102 USA - X 188 220 89 Germany X X 116 243 128	Peng et al. [124]	Taiwan	×	×	263	230	185	45	0
Canada - × 84 99 99 Czech Republic × - 64 185 48 Japan × - 53 147 102 USA - × 188 220 89 Germany × × 116 243 128	Poruk et al. [125]	USA	×	×	98	134	98	48	0
Czech Republic × - 64 185 48 Japan × - 53 147 102 USA - × 188 220 89 Germany × × 116 243 128	Ritchie et al. [126]	Canada	I	×	84	66	66	0	0
Japan × - 53 147 102 USA - × 188 220 89 Germany × × 116 243 128	Rychlikova et al. [127]	Czech Republic	×	ı	64	185	48	137	0
USA $ \times$ 188 220 89 Germany \times \times 116 243 128	Sakai et al. [128]	Japan	×	ı	53	147	102	22	23
Germany $\times \times \times 116$ 243 128	Song et al. [129]	USA	I	×	188	220	68	89	63
	Tachezy et al. [130]	Germany	×	×	116	243	128	115	0

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References	Country (population)	Setting ^a		Cases and controls	rols			
				Cases (N)	Contro	Controls (N)		
		Hosp	Other		All	НС	NM	PM
Talar-Wojnarowska et al. [131]	Poland	×	ı	85	122	90	72	0
Tavano et al. [132]	Italy	×	ı	74	117	117	0	0
Ward et al. [133]	UK	×	ı	75	61	0	61	0
Xu et al. [134]	China	×	ı	156	180	9	57	28
Zhang et al. [135]	China	×	ı	129	278	183	95	0
Zhang et al. [136]	China	×	ı	29	206	145	61	0
Zhong et al. [137]	China	×	ı	183	202	141	61	0
Zhou et al. [138]	China	×	ı	152	207	96	91	20
Zhou et al. [139]	China	×	ı	156	199	163	36	0
Zhou et al. [140]	China	×	1	64	64	64	0	0
Oesophageal cancer only								
Bagaria et al. [141]	India	×	ı	50	50	50	0	0
Bai et al. [142]	China	×	ı	68	125	80	14	31
Bagaria et al. [143]	India	×	ı	50	90	50	0	0
Brockmann et al. [144]	Germany	×	ı	50	150	20	100	0
Huang et al. [145]	China	×	ı	09	09	09	0	0
Jia et al. [146]	China	×	I	101	86	86	0	0
Liao et al. [147]	China	×	I	151	230	194	36	0
Lukaszewicz-Zajac et al. [148]	Poland	×	ı	56	99	99	0	0
Lv et al. [149]	China	×	ı	126	80	80	0	0
Pan et al. [150]	China	×	ı	50	110	09	50	0
Peng et al. [151]	China	×	ı	104	53	53	0	0

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Subject of al. [152] Japan Amort al. [153] Controls (N) Controls (N) Sub or al. [153] Japan x x x x3 9364 9203 161 Wang et al. [153] China x <	References	Country (population)	Setting ^a		Cases and controls	sle			
Japan X X All HC NM China X 2 283 9364 9203 161 China X - 84 154 154 161 China X - 169 154 80 74 China X - 169 134 9 74 China X - 169 134 9 74 China X - 186 18 1 0 China X - 186 186 1 0 China X - 186 186 186 9 0 China X - 150 185 126 59 0 China X - X 88 479 200 0 China X - X 186 186 9 18 China X <th></th> <th></th> <th></th> <th></th> <th>Cases (N)</th> <th>Contro</th> <th>Is (N)</th> <th></th> <th></th>					Cases (N)	Contro	Is (N)		
Japan x x 283 9364 9203 161 China x - 84 154 154 0 China x - 169 134 149 0 China x - 367 134 134 0 China x - 364 229 229 0 China x - 81 81 81 0 China x - 81 81 81 0 China x - 112 112 112 0 China x - 150 185 186 0 0 China x - 150 185 126 59 0 China x - 150 185 126 59 0 China x - 150 128 729 126 126			Hosp	Other		All	HC	NM	PM
China x - 84 154 154 0 China x - 169 154 80 74 China x - 237 134 134 0 China x - 364 229 229 0 China x - 81 81 81 0 China x - 81 81 81 0 China x - 112 112 112 0 China x - 186 186 186 0 0 China x - 112 112 112 0 0 China x - 150 186 186 0 0 0 China x x x 88 126 200 0 China x x x x x x x	Sudo et al. [152]	Japan	×	×	283	9364	9203	161	0
China x - 169 154 80 74 China x - 237 134 134 0 China x - 364 229 229 0 China x - 81 81 81 0 China x - 81 81 81 0 China x - 112 112 112 0 China x - 150 186 186 0 0 China x - 150 112 112 0 0 China x x 88 479 200 0 0 China x x 88 479 200 0 0 China x x x 128 0 65 0 65 China x x x x 128 0 65 0 65 China x x x x x	Wang et al. [153]	China	×	ı	84	154	154	0	0
China x - 237 134 134 0 China x - 70 80 80 0 China x - 81 81 81 0 China x - 62 58 58 0 China x - 81 81 81 0 China x - 186 186 186 0 China x - 112 112 112 0 China x - 150 186 186 0 0 China x - 150 186 186 0 0 0 China x x 8 479 200 0 0 China x x 8 126 78 128 China x x 8 128 128 128 China x x x 128 78 78 China x x<	Xing et al. [154]	China	×	ı	169	154	80	74	0
China x - 70 80 80 0 China x - 364 229 229 0 China x - 62 58 58 0 China x - 81 81 81 0 China x - 112 112 112 0 China x - 150 186 0 0 China x - 150 185 126 50 China x - 153 65 0 65 China x - 153 78 78 China x - 20 0 65 China x - 20 0 65 China x	Xu et al. [155]	China	×	ı	237	134	134	0	0
China x - 364 229 229 0 China x - 81 81 81 0 China x - 81 81 81 0 China x - 186 186 186 0 China x - 112 112 112 0 China x - 150 186 186 0 0 China x - 153 65 0 65 0 65 China x - x 88 479 200 0 0 China x - x 88 479 200 0 65 0 65 0 65 0 65 0 65 0 65 0 65 0 65 0 65 0 65 0 65 0 65 0 65 0 65 0 0 0 0 0 0 0 0	Xu et al. [156]	China	×	ı	70	80	80	0	0
China x - 81 81 81 81 0 China x - 62 58 58 0 China x - 186 186 180 0 China x - 112 112 112 0 China x - 150 185 126 59 China x x 88 479 200 0 China x x 88 479 200 0 China x x x 20 65 0 65 China x x x x x x x x x India x <	Yan et al. [157]	China	×	ı	364	229	229	0	0
China x - 62 58 58 58 0 China x - 81 81 81 0 China x - 186 186 186 0 0 China x - 112 112 112 0 0 China - x 88 479 200 0 0 China x - x 88 479 200 0 0 China x - x 88 479 200 0 0 China x - x 153 65 0 65 0 65 China x - 78 128 0 65 0 65 China x - 78 78 78 78 78 India x - 50 GC 50 50 0 6 W x - 163 GC Or OC 1724 89 89 89	Zhang et al. [158]	China	×	ı	81	81	81	0	0
China x - 81 81 81 0 China x - 186 186 186 186 186 186 186 186 186 186 186 186 186 112 112 0 China - x 88 479 200 0 0 China x - x 88 479 200 0 China x - x 88 479 200 0 China x - x 59 128 0 65 China x - x x 156 78 78 India x - x x x 78 78 78 UK x - 163 GC or OC 172 ⁴ 89 89 89	Zhang et al. [159]	China	×	ı	62	58	58	0	0
China x - 186 186 186 186 0 China x - 112 112 112 0 China x - 150 185 126 59 China x - 59 128 0 65 China x - 78 128 78 78 India x - 50 GC 78 78 UK x - 163 GC or OC 172 ^d 89 82	Zhang et al. [160]	China	×	ı	81	81	81	0	0
China x - 112 112 112 0 China x - 150 185 126 59 China x - 153 65 0 65 0 China x - 59 128 0 128 China x - 78 78 78 India x - 50 GC 78 78 UK x - 163 GC or OC 172 ^d 89 89 82	Zhang et al. [161]	China	×	ı	186	186	186	0	0
China x - 150 185 126 59 China - x 88 479 200 0 China x - 153 65 0 65 Thailand x - 59 128 0 128 China x - 78 78 78 78 India x - 50 GC 50 0 0 WK x - 163 GC or OC 172 ⁴ 89 82	Zhang et al. [162]	China	×	ı	112	112	112	0	0
China x x 479 200 0 China x - 153 65 0 65 Thailand x - 59 128 0 128 China x - 78 156 78 78 India x - 50 GC 50 50 0 UK x - 163 GC or OC 172 ^d 89 82	Zheng et al. [163]	China	×	ı	150	185	126	65	0
China x - 153 65 0 65 Thailand x - 59 128 0 128 China x - 78 78 78 India x - 50 GC 50 0 UK x - 163 GC or OC 172 ^d 89 82	Zhou et al. [164]	China	I	×	88	479	200	0	279
China × - 153 65 0 65 Thailand × - 59 128 0 128 China × - 78 78 78 India × - 50 GC 50 0 UK × - 163 GC or OC 172 ^d 89 82	Biliary tract cancers only								
Thailand x - 59 128 0 128 China x - 78 78 78 India x - 50 GC 50 0 VK x - 163 GC or OC 172 ^d 89 82	Deng et al. [165]	China	×	ı	153	99	0	65	0
China × - 78 156 78 78 India × - 50 GC 50 50 0 VK × - 163 GC or OC 172 ^d 89 82	Leelawat et al. [166]	Thailand	×	ı	59	128	0	128	0
India \times - 50GC $50 50 0$ 50OC $50 $	Wang et al. [167]	China	×	I	78	156	78	78	0
India \times - 50 GC 50 50 0 80 CV	More than one tumour type								
50 OC $VK \qquad \qquad \times \qquad - \qquad 163 \text{ GC or OC} \qquad 172^{\text{d}} \qquad 89 \qquad 82$	Bagaria et al. [168]	India	×	I	50 GC	50	50	0	0
UK \times - 163 GC or OC 172 ^d 89 82					50 OC				
	Markar et al. [169]	UK	×	ı	163 GC or OC	172^{d}	68	82	0

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References	Country (population)	Setting ^a		Cases and controls	ols			
				Cases (N)	Controls (N)	(N)		
		Hosp	Hosp Other		All	НС	NM	PM
Ren et al. [170]	China	×	ı	1049 GC	1019 747	747	272	0
				268 OC				
				160 PaC				
Schneider et al. [171]	Germany	Ŋ	U	122 GC	53	53	0	0
				30 98				

GC gastric cancer, HC healthy control, Hosp hospital, NM non-malignant, OC oesophageal cancer, PaC pancreatic cancer, PM pre-malignant, U unclear, UK United Kingdom, USA United States of America

^a Due to wide variations in health systems across different countries, hospital setting is a broad definition than can encompass secondary and tertiary care. Other setting refers to biobanks, reference sets, databases, or archived samples; general population cohorts or cohorts from population screening programmes; or cohorts b In most of these studies, unclear numbers refer to healthy controls and non-malignant conditions combined (70 controls for [26], 120 controls for [42], 60 from previous trials or observational studies

controls for [49], and 70 controls for [50]). In the case of Zhou et al. [85], it is also unclear whether controls had pre-malignant conditions ^c Denmark, France, Italy, Germany, Greece, Spain, UK, Norway, Sweden & Netherlands

^d Sum of controls does not add up to total number of controls (mismatch in paper)

distinguishing between oesophageal and gastric cancer). The minimum age for cases was 16 while the oldest patient was aged 93. Most cases were male (68%) across all tumour types. Over 50% of cancers had been diagnosed at stages III and IV (median 55.5%, interquartile range 47.0-68.1%; data available for 106 included studies). The included studies reported on 49,474 controls (38,955 normal/healthy, 9042 with non-malignant conditions, 1106 with premalignant conditions, and 371 with either normal or non-malignant conditions). Pancreatitis and gastritis were the most commonly reported non-malignant conditions (online supplementary Figure S1). Over half of the studies (n = 83) investigated more than one type of control population. Normal healthy controls were the majority across all tumour types, except for biliary tract cancers. The minimum age for controls was 16 while the maximum age was 94. Overall, most controls were male (74%); this was the case for all tumour types except for biliary tract cancers.

Types of Biomarkers

Biomarkers were most commonly sampled from blood (145 studies; 107 investigated serum, 33 plasma and 5 both); two studies analysed urine [28, 36], one breath [169] and another saliva [47]. Most studies (n = 128) investigated more than one biomarker. A total of 431 biomarkers were identified (online supplementary table S2). These were most often microRNA and other RNAs (n = 183), other proteins (n = 119) and autoantibodies and other immunological markers (n = 79). Less than a third of studies (n = 44) included biomarkers from different categories. This was often due to use of established biomarkers (proteins CA19-9 and CEA) in combination with novel biomarkers. Studies of pancreatic cancer reported on over half of identified biomarkers (n = 231) (Fig. 3). Only about a fifth (n = 90) of all identified biomarkers were reported in more than one study; 72 of these were reported in more than one study for the same tumour type (Table 3).

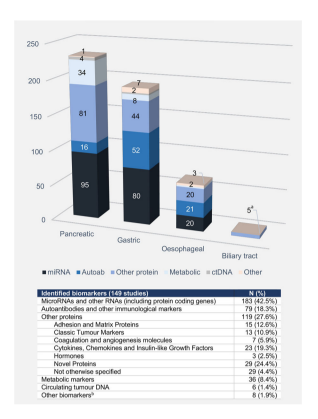


Fig. 3 Types of biomarkers, overall and by tumour type. ^aFive proteins; ^bthese refer to volatile organic compounds and platelets; *autoab* autoantibodies, *ctDNA* circulating tumour DNA, *miRNA* microRNA

Measures of Diagnostic Performance

The most commonly reported measures of performance diagnostic were sensitivity (n = 136). specificity (n = 129)and (n = 123). PPV and NPV were each reported by 40 studies, while false positives and false negatives were least often reported (11 studies each). Outcome data on individual biomarkers were available in most studies (n = 121); the remaining 28 studies only reported on performance for a combination/panel. Over half of the included studies (n = 83) reported on measures of performance for biomarkers both individually and in combinations. Outcome data were not available for all control populations; only 95 studies provided outcome data for cancers versus normal controls, 54 provided outcome data for cancers versus non-malignant controls, and 10 provided measures for cancers

Table 2 Characteristics of included studies: biomarkers and study design

Signature counters only Sample Seample Report Signature counters only Seample Seample<	References	Biomarke	ers										Design	ı	
		Type (N	(Sample			Repo	l H	Sgl	2-gate	
24] 15 16 17 26] 27 27 28 28 29 29 29 29 29 29 29 29 29 29 29 29 29		miRNA	Autoab	Protein	Metab	ctDNA	Other ^a	Serum	Plasma	Other ^b	Ind	Comb	RFD	LGN	TGA
14] 15	Gastric cancers only														
24]	Cai et al. [23]	15	1	1	ı	1	ı	1	×	1	×	1	ı	×	ı
25]	Chen et al. [24]	I	ı	1	ı	ı	ı	×	ı	ı	×	ı	D	D	Ω
[26] 1. Columnation 1. Columnation <td>Chen et al. [25]</td> <td>I</td> <td>ı</td> <td>4</td> <td>ı</td> <td>ı</td> <td>ı</td> <td>×</td> <td>ı</td> <td>1</td> <td>×</td> <td>×</td> <td>ı</td> <td>×</td> <td>×</td>	Chen et al. [25]	I	ı	4	ı	ı	ı	×	ı	1	×	×	ı	×	×
25] 4	Chung et al. [26]	I	1	7	ı	1	ı	×	ı	1	×	×	D	×	n
28] 5. L. C. D. L. C. D. L. C. D. C. C. C. D. C. D. C. D. C. D. C. D. C. D. D. C. D.	Ding et al. [27]	4	ı	1	ı	ı	ı	×	ı	ı	×	×	ı	×	ı
1. [32] 1. [32] <t< td=""><td>Dong et al. [28]</td><td>I</td><td>1</td><td>1</td><td>ı</td><td>1</td><td>ı</td><td>1</td><td>ı</td><td>×</td><td>×</td><td>ı</td><td>ı</td><td>×</td><td>ı</td></t<>	Dong et al. [28]	I	1	1	ı	1	ı	1	ı	×	×	ı	ı	×	ı
39] 1	Gantuya et al. [29]	I	1	2	ı	1	ı	×	ı	ı	×	×	×	ı	ı
1. [32] 1. [32] 2. [4] 1. [4	Gwak et al. [30]	I	1	5	ı	1	ı	×	ı	1	×	ı	ı	ı	×
11. [32] 1. [32] 1. [32] 1. [32] 1. [32] 1. [33] 1. [34] <	He et al. [31]	I	ı	4	ı	ı	ı	×	ı	ı	×	×	D	×	n
[34] 1 5 1 5 1 7 1 7 1 7 1 7 1 7 1 7 1 7 1 7 1 7 1 7 1 7 1 7 1 7 1 7 1 7 1 7 1 7 1 7 1 7 1 1 7 1 1 1 8 1	Hoshino et al. [32]	I	9	7	ı	ı	ı	×	ı	ı	×	×	ı	×	ı
[34] 5 - 2 -	Huang et al. [33]	I	П	~	ı	ı	ı	×	ı	ı	×	ı	1	×	×
[35] 5	Huang et al. [34]	>	I	2	ı	1	ı	×	ı	1	×	×	ı	×	ı
[36] 2	Huang et al. [35]	>	ı	ı	ı	1	ı	×	ı	1	ı	×	D	D	D
1. [38] 2 - </td <td>Iwasaki et al. [36]</td> <td>2</td> <td>I</td> <td>ı</td> <td>ı</td> <td>ı</td> <td>ı</td> <td>1</td> <td>ı</td> <td>×</td> <td>×</td> <td>ı</td> <td>ı</td> <td>×</td> <td>I</td>	Iwasaki et al. [36]	2	I	ı	ı	ı	ı	1	ı	×	×	ı	ı	×	I
X X	Ji et al. [37]	2	ı	1	ı	ı	ı	ı	×	ı	×	ı	ı	MB	ı
	Juan Cai et al. [38]	I	1	3	ı	ı	ı	×	ı	ı	×	ı	ı	MB	MB
	Kaise et al. [39]	I	1	~	ı	ı	ı	×	ı	ı	×	×	ı	×	ı
	Kang et al. [40]	ı	I	1	ı	1	ı	×	ı	1	×	ı	×	ı	ı
1	Kikuchi et al. [41]	I	ı	2	ı	ı	ı	×	ı	ı	×	×	×	ı	ı
1 2	Kim et al. [42]	1	ı	ı	ı	ı	ı	×	ı	ı	×	ı	ı	×	×
	Kurilovich et al. [43]	I	1	2	ı	1	1	×	ı	1	×	×	ı	×	ı

Table 2 continued

Li et al. [44] 3 - - Li et al. [45] 1 - - Li et al. [46] 3 - - - Li et al. [46] 3 - 4 -	7)		Sar	11.		Done	=	Col	2-gate	
3 Autoab 3 3 4 4 4 4 4 4 4 4				Sample		neport		jo Jo	c	
3	Autoab Protein Metab	ctDNA	Other ^a Ser	Serum Plasma	a Other ^b	Ind	Comb	RFD	LGN	TGA
5] 3		I	1	×	ı	×	×	U	U	n
3	1	ı	1	×	ı	×	ı	Ŋ	×	D
13		ı	ı	×	ı	×	1	ı	×	ı
5] 2		ı	1	ı	×	×	×	I	×	I
5] 2 2 1 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2		I	×	I	I	×	×	MB	1	1
5] 2		I	×	I	ı	×	×	D	×	×
5] 2 2		ı	×	I	ı	×	×	MB	×	×
5] 3	1	I	×	×	ı	×	1	D	MB	D
5] 3 - 1 18 - 1 18 - 1 19 - 1 1		ı	×	I	I	ı	×	ı	×	ı
5] 3 - 1 18 - 18 - 18 - 19 - 19 - 19 - 19 -		ı	×	ı	ı	×	×	ı	×	×
5] - 18 56]		ı	1	×	ı	ı	×	ı	×	ı
56]	18	ı	×	ı	ı	ı	×	ı	×	ı
		I	×	×	ı	×	ı	ı	×	ı
	·	I	×	I	ı	×	×	ı	×	ı
3 - 1		ı	×	I	I	×	×	ı	×	×
1 %		ı	×	×	I	×	×	D	×	D
Parvace et al. [61] 3 – – –	1	2	I	×	ı	×	×	ı	×	×
	1	ı	I	×	ı	×	ı	ı	×	ı
Qin et al. [62] – 9 –	- 6	ı	×	I	ı	×	×	ı	×	I
Qiu et al. [63] 4		I	ı	×	ı	×	×	D	Ŋ	D
Song et al. [64] 8 – –		ı	×	I	I	×	×	×	1	I
Su et al. [65] – – 5		ı	×	ı	ı	ı	×	ı	×	×

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References	Biomarker	ers										Design	_	
	Type (N)						Sample			Report	l E	Sgl	2-gate	
	miRNA	Autoab	Protein	Metab	ctDNA	Other ^a	Serum	Plasma	Other ^b	Ind	Comb	RFD	TGN	TGA
Sun et al. [66]	I	1	3	ı	1	1	×	ı	ı	×	×	MB	ı	ı
Tsalikidis et al. [67]	I	ı		1	1	1	×	1	ı	×	ı	ı	×	ı
Wang et al. [68]	I	ı		ı	ı	ı	×	ı	ı	×	ı	ı	×	ı
Wang et al. [69]	5	ı	ı	ı	ı	ı	×	ı	ı	I	×	D	D	D
Wang et al. [70]	I	9	ı	ı	ı	ı	×	ı	ı	ı	×	ı	×	I
Wang et al. [71]	I	ı	3	ı	ı	ı	×	ı	ı	×	×	Ω	×	D
Werner et al. [72]	I	14	ı	ı	ı	1	×	1	ı	1	×	ı	×	ı
Wu et al. [73]	1	ı	2	1	1	1	×	1	ı	×	ı	Ŋ	Ŋ	D
Wu et al. [74]	ı	ı	4	ı	Ω	ı	×	ı	ı	×	ı	ı	×	×
Wu et al. [75]	I	ı		1	ı	3	×	1	1	×	×	Ŋ	×	D
Yanaoka et al. [76]	I	ı	2	ı	ı	ı	×	ı	ı	×	×	×	ı	ı
Yang et al. [77]	I	I	1	ı	ı	ı	ı	×	ı	×	ı	×	ı	I
Yang et al. [78]	3	ı	~	ı	I	ı	I	×	ı	×	×	I	I	×
Yoon et al. [79]	I	ı	1	ı	ı	ı	×	ı	ı	×	1	I	×	I
Yun et al. [80]	I	I	1	ı	ı	2	×	ı	ı	×	×	MB	×	MB
Zayakin et al. [81]	I	45	ı	ı	ı	1	×	1	ı	ı	×	ı	×	×
Zhang et al. [82]	I	1	ı	9	1	1	×	1	ı	×	×	ı	×	×
Zhang et al. [83]	П	ı	ı	ı	1	1	ı	×	ı	×	ı	ı	ı	×
Zhang et al. [84]	~	ı	4	ı	ı	ı	ı	×	ı	×	×	ı	ı	×
Zhou et al. [85]	1	I	ı	ı	ı	ı	ı	×	ı	×	ı	Ω	Ω	D
Zhou et al. [86]	~	ı	ı	ı	ı	ı	ı	×	ı	ı	×	Ω	Ω	Ŋ
Zhou et al. [87]	1	1	1	1	1	1	1	×	1	×	1	U	U	U

Table 2 continued

References	Biomarkers	srs										Design	ـ ا	
	Type (N)						Sample			Report	E	Sgl	2-gate	
	miRNA	Autoab	Protein	Metab	ctDNA	Other ^a	Serum	Plasma	Other b	Ind	Comb	RFD	LGN	TGA
Pancreatic cancers only														
Akita et al. [88]	I	ı	ı	4	ı	ı	×	ı	ı	×	×	D	D	D
Balasenthil et al. [89]	I	ı	3	ı	1	ı	ı	×	ı	ı	×	×	1	ı
Brand et al. [90]	ı	1	3	1	1	1	×	ı	ı	×	×	ı	×	×
Cao et al. [91]	9	1	1	ı	1	ı	ſ	×	ı	ı	×	D	D	D
Capello et al. [92]	9	ı	7	ı	ı	ı	ı	×	ı	×	×	Ŋ	D	D
Chung et al. [93]	I	2	ı	П	ı	ı	×	ı	ı	×	×	Ŋ	×	D
Chung et al. [94]	ı	_	20	ı	ı	1	×	ı	1	×	×	×	×	ı
Deng et al. [95]	1	1	1	ı	1	ı	×	ı	1	×	ı	Ŋ	Ω	Ŋ
Duraker et al. [96]	ı	1	3	ı	1	ı	×	ı	ı	×	×	Ŋ	Ω	Ŋ
Firpo et al. [97]	I	ı	3	ı	ı	ı	×	I	ı	×	×	MB	×	MB
Fukutake et al. [98]	I	ı	ı	9	ı	ı	ı	×	ı	ı	×	ı	×	×
Gao et al. [99]	1	ı	1	ı	ı	ı	×	I	ı	×	×	D	×	Ω
Gold et al. [100]	I	I	1	ı	I	ı	×	ı	ı	×	ı	ı	×	×
Gold et al. [101]	I	1	1	ı	ı	ı	×	ı	ı	×	×	D	×	D
Groblewska et al. [102]	ı	ı	4	ı	ı	ı	×	ı	ı	×	×	ı	×	ı
Guo et al. [103]	I	ı	2	ı	ı	ı	×	ı	ı	×	×	D	×	D
Honda et al. [104]	ı	ı	4	ı	ı	ı	ı	×	ı	×	×	×	ı	ı
Honda et al. [105]	I	ı	3	1	ı	ı	ı	×	ı	×	×	×	ı	ı
Honda et al. [106]	I	ı	3	ı	ı	ı	ı	×	ı	×	×	×	ı	ı
Jiang et al. [107]	ı	ı	3	1	ı	1	×	ı	ı	×	×	ı	×	×
Kaur et al. [108]	ı	1	1	1	1	1		×	1	×	1	×	1	ı

Table 2 continued

Type (W) miRNA	Autoab	Protein A	Metab	ctDNA	Other ^a	Sample		,	Report	Comb	Sgl	2-gate TGN	
	Autoab	rotein	Metab	ctDNA	Other ^a	Comme		4	Lad	Comb	0.10	LGN	
[111] 						Sci mii	Plasma	Other"	Ind		Kru		TGA
[11] 	1 1 1 1 1 1 1	γ,		ı	1	×	×	ı	×	×	1	×	×
	1 1 1 1 1 1	,	ı	ı	1	×	ı	1	×	×	Ŋ	Ω	D
1 1 1 1 1	1 1 1 1 1	1	ı	3	1	ı	×	1	ı	×	Ω	×	Ω
	1 1 1 1	9	ı	ı	1	×	I	1	×	×	Ŋ	Ω	Ŋ
L	1 1 1	2	ı	ı	ı	×	ı	1	×	×	ı	×	×
L	1 1	1	ı	ı	ı	ı	×	ı	×	×	MB	×	MB
I	I	1	ı	ı	ı	×	1	ı	ı	×	ı	×	×
		11	ı	ı	1	×	ı	1	×	×	ı	×	×
Matsubara et al. [117] – –	1	2	ı	ı	ı	1	×	ı	×	×	D	MB	D
Mayerle et al. [118]	ı	1 9	6	ı	ı	ı	×	ı	ı	×	MB	ı	MB
Mellby et al. [119] 1 5	ς	20 3	3	ı	ı	×	1	ı	ı	×	×	ı	ı
Mizuno et al. [120]	I)	9	ı	ı	ı	×	ı	I	×	ı	×	×
O'Brien et al. [121] 1	I	3	ı	ı	ı	×	ı	ı	×	×	×	ı	ı
Park et al. [122]	ı	- 6	ı	ı	ı	×	ı	ı	×	×	D	MB	D
I	ı	5	ı	ı	ı	×	ı	ı	×	×	Ŋ	×	×
Peng et al. [124]	ı	2	ı	ı	ı	×	ı	ı	×	×	ı	×	×
Poruk et al. [125]	ı	3	ı	ı	ı	×	ı	ı	×	×	ı	×	MB
Ritchie et al. [126]	I	1	_	ı	ı	×	ı	ı	×	×	D	Ŋ	D
Rychlikova et al. [127]	I	4	ı	ı	ı	×	ı	ı	×	×	MB	Ŋ	MB
Sakai et al. [128] 56 -	I	2	ı	ı	ı	×	×	ı	×	×	ı	×	MB
I	3	3	ı	ı	ı	×	ı	ı	×	×	Ŋ	Ŋ	Ω
Tachezy et al. [130]	ı	ı	ı	1	1	×	1	1	×	1	D	×	D

Table 2 continued

Type (N) Sumple Sumple	References	Biomarker	ers										Design		
Sect al. [131]		Type (N	(Sample			Repo	ıı	Sgl	2-gate	
Sea et al. [131] - 1		miRNA	Autoab	Protein	Metab	ctDNA	Other ^a	Serum	Plasma	$Other^b$	Ind	Comb	RFD	TGN	TGA
23 1	Talar-Wojnarowska et al. [131]	ı	1	1	ı	ı	ı	×	ı	ı	×	ı	U	MB	U
c ct al. [148] 10	Tavano et al. [132]	1	ı	_	ı	ı	ı	×	ı	1	×	×	×	ı	ı
s = 1	Ward et al. [133]	ı	ı	_	2	ı	ı	×	ı	1	×	×	Ŋ	D	Ŋ
	Xu et al. [134]	8	ı	1	ı	1	ı	ı	×	ı	×	ı	Ω	Ω	Ŋ
1	Zhang et al. [135]	ı	2	3	_	1	1	×	1	ı	ı	×	Ŋ	D	D
7]	Zhang et al. [136]	ı	ı	1	9	ı	ı	×	ı	1	×	×	U	×	Ŋ
1	Zhong et al. [137]	ı	_		ı	ı	ı	×	ı	ı	×	×	Ŋ	D	D
	Zhou et al. [138]	ı	1	2	ı	ı	ı	×	ı	ı	×	×	×	ı	ı
rs only 1] 66 1	Zhou et al. [139]	ı	ı	2	ı	1	1	×	1	1	×	×	Ω	Ω	Ŋ
1]	Zhou et al. [140]	9	ı	ı	ı	ı	ı	ı	×	ı	ı	×	ı	×	ı
1] 1	Oesophageal cancers only														
3]	Bagaria et al. [141]	ı	ı		ı	ı	ı	×	ı	ı	×	ı	D	D	D
3] . [144]	Bai et al. [142]	1	1	1	ı	ı	I	ı	×	ı	×	×	ı	×	×
5] 5 2 7 7 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8	Bagaria et al. [143]	ı	I	4	ı	ı	ı	×	ı	1	×	ı	ı	×	ı
c et al. [148]	Brockmann et al. [144]	ı	7	2	ı	ı	ı	×	ı	ı	×	ı	ı	×	×
c et al. [148]	Huang et al. [145]	5	ı	ı	ı	ı	ı	×	ı	ı	ı	×	ı	MB	ı
c et al. [148] 4	Jia et al. [146]	1	ı	9	ı	ı	ı	×	ı	ı	ı	×	ı	×	ı
c et al. [148] 2 × × × × × × × × × × × × × × ×	Liao et al. [147]	ı	I	4	1	ı	ı	ı	×	1	×	×	Ω	Ŋ	D
2	Lukaszewicz-Zajac et al. [148]	ı	I	2	ı	ı	ı	×	ı	ı	×	×	ı	×	ı
	Lv et al. [149]	2	ı	ı	ı	ı	ı	×	ı	ı	×	×	ı	×	ı
× × × · · · · · · · · · · · · · · · · ·	Pan et al. [150]	ı	4	ı	ı	ı	ı	×	ı	ı	×	×	Ω	×	D
	Peng et al. [151]	ı	1	1	ı	1	ı	×	1	ı	×	×	ı	MB	I

Table 2 continued

	Type (N)						Sample			Report	ㅂ	Sgl	7-gate	
	miRNA	Autoab	Protein	Metab	ctDNA	Other ^a	Serum	Plasma	Other ^b	Ind	Comb	RFD	TGN	TGA
Sudo et al. [152]	9	I	ı	I	ı	ı	×	ı	ı	ı	×	ı	×	×
Wang et al. [153]	1	ı	1	ı	ı	ı	×	ı	1	×	ı	Ŋ	D	Ω
Xing et al. [154]	2	ı		1	1	1	×	1	ı	×	×	ı	×	×
Xu et al. [155]	ı	V		1	ı	ı	×	1	1	ı	×	ı	×	ı
Xu et al. [156]	ı	5	П	1	1	1	×	1	ı	ı	×	ı	×	1
Yan et al. [157]	ı	ı	1	1	1	ı	×	ı	1	×	1	ı	×	1
Zhang et al. [158]	1	ı	1	1	ı	1	×	1	ı	×	ı	Ŋ	Ŋ	Ω
Zhang et al. [159]	ı	1	1	1	ı	1	×	1	1	×	ı	Ŋ	D	D
Zhang et al. [160]	1	ı	1	1	ı	ı	×	1	1	×	ı	Ŋ	Ŋ	Ω
Zhang et al. [161]	ı	9	1	1	ı	ı	×	1	1	ı	×	ı	×	ı
Zhang et al. [162]	ı	2	1	1	ı	ı	×	1	1	×	×	Ŋ	Ŋ	Ω
Zheng et al. [163]	ı	ı	4	1	ı	1	×	1	ı	×	×	ı	×	×
Zhou et al. [164]	ı	∞	1	ı	ı	ı	×	ı	ı	×	×	ı	ı	×
Biliary tract cancers only														
Deng et al. [165]	ı	ı	4	ı	ı	ı	×	ı	1	×	×	ı	ı	×
Leelawat et al. [166]	ı	ı	2	1	ı	ı	×	1	ı	×	ı	×	ı	ı
Wang et al. [167]	I	ı	4	ı	ı	1	×	ı	ı	×	×	MB	×	ı
More than one tumour type														
Bagaria et al. [168]	I	ı	2	ı	ı	ı	×	ı	ı	×	×	ı	×	ı
Markar et al. [169]	I	ı	I	ı	ı	5	ı	ı	×	ı	×	MB	ı	ı
Ren et al. [170]	I	1	2	ı	ı	ı	×	ı	ı	×	×	D	D	D

Table 2 continued

References	Biomarker	ers										Design	_	
	Type (N)						Sample			Report		Sgl	Sgl 2-gate	
	miRNA	Autoab	Protein	Metab	Metab ctDNA Other ^a S	Other ^a	Serum	Serum Plasma Other ^b	$Other^b$	Ind Comb	Comb	RFD	TGN TGA	$\overline{\mathrm{TGA}}$
Schneider et al. [171]	ı	ı	4	ı	ı	ı	×	ı	ı	×	×	ı	×	ı

autoantibodies and other immunological markers, Combination or panel, ctDNA circulating tumour DNA, Ind individual, MB maybe/likely (design ikely but no sufficient information to make a final decision), metab metabolic markers, RFD reversed-flow design, Sgl single-gate design, TGA two-gate alternative diagnosis, TGN two-gate normal, U unclear

^a Other biomarker type refers to volatile organic compounds or platelets

Other sample refers to urine or volatile organic compounds

versus pre-malignant conditions (online supplementary table S3).

Individual measures of diagnostic performance were available for 35 biomarkers mentioned more than once, for the same tumour type (online supplementary table S4). We were not able to synthesise outcomes further due to heterogeneity in biomarker combinations, in control populations and subgroup analyses, and variations in reported cut-off points and diagnostic accuracy data (see online supplementary table S5 for a textual description of outcomes).

Only four novel biomarkers were reported on studies adopting a single-gate design (Table 4). Apolipoproteins AII-AT and AII-ATQ had poor sensitivity (range 4–25%) but good AUCs (range 52-94.6%) reported for pancreatic cancer in three studies (same first author for all) [104–106]. Their diagnostic accuracy increased when combined with CA19-9 (sensitivity range 7-95.4%, specificity range 96-98%, AUC range 56–78%). Pepsinogen I (PGI) and PGI/PGII ratio had a wide range of sensitivity and specificity (ranges 27–77.9% and 20.2–92%, respectively) and good AUC (range 70-76%) reported for gastric cancer across four studies [29, 40, 41, 76]. When evaluated with other novel biomarkers (including miR-1290, MIC-1, ULBP2 and CA125), one established biomarker, CA19-9, also showed some promise (sensitivity range 23.1-88%, specificity range 71.6-96.6%, AUC 92–98%) for pancreatic cancer [121, 132, 138]. There were also two studies reporting panels rather than individual biomarkers using a single-gate, reversed-flow design (Table [89, 119].

DISCUSSION

Our systematic review identified 149 studies reporting on 431 different biomarkers for gastric, pancreatic, oesophageal and biliary tract cancers. Only a fifth of biomarkers were reported by more than one study, and from these only four novel biomarkers, apoAII-AT and apoAII-ATQ (pancreatic cancer) and pepsinogen I and II (gastric cancer), plus one established biomarker (CA19-9 combined with other novel biomarkers), were reported with individual

Table 3 Biomarkers investigated more than once, for the same tumour type (number of studies)

Biomarker	Pancreatic cancer	Gastric cancer	Oesophageal cancer	Biliary tract
)	cancer
MicroRNAs and other	MicroRNAs and other RNAs (including protein coding genes)			
miR-21	2 [114, 115]	3 [23, 34, 44]	ı	I
miR-20a	1	3 [23, 52, 86]	1	I
miR-25	2 [95, 115]	2 [46, 86]	I	I
miR-296-5p	ı	2 [35, 69]	I	I
miR-210	ı	2 [61, 86]	I	I
miR-1	ı	2 [23, 52]	I	I
miR-106b	ı	2 [23, 46]	I	I
miR-106b-3p	2 [91, 134]	I	ı	I
miR-126-3p	2 [91, 134]	I	ı	I
miR-1285	2 [91, 134]	I	1	I
miR-132-3p	I	2 [35, 69]	ı	I
miR-16	2 [99, 114]	I	ı	I
miR-214	I	2 [37, 83]	ı	I
miR-221	ı	2 [23, 64]	ı	I
miR-223	ı	2 [44, 85]	I	I
miR-26b-3p	2 [91, 134]	I	I	I
miR-27a	ı	2 [23, 52]	I	I
miR-376c	I	2 [23, 64]	ı	I
miR-423-5p	ı	2 [23, 52]	ı	I
miR-486-5p	2 [91, 134]	I	I	I
miR-744	I	2 [23, 64]	1	I
miR-938	2 [91, 134]	I	I	I
REG3A	7 [97 171]	1		

Biomarker	Pancreatic cancer	Gastric cancer	Oesophageal cancer	Biliary tract
			0 J	cancer
Autoantibodies and other immunological markers	ınological markers			
p53	1	2 [32, 62]	4 [155, 156, 161, 164]	I
C-Myc	1	2 [62, 70]	2 [161, 164]	I
p62	1	2 [62, 70]	2 [161, 164]	I
New York esophageal squamous cell carcinoma 1 (NY-ESO-1 or CTAG1A)	1	I	3 [150, 155, 156]	I
Squamous Cell Carcinoma- Antigen (SCC-Antigen)	1	ı	3 [144, 147, 163]	I
Antibodies against Helicobacter pylori (HpAb)	1	2 [39, 66]		I
BMI-1	I	ı	2 [155, 156]	I
Heat shock protein 70 (HSP70)	1	ı	2 [155, 156]	I
Immunoglobin G galactosylation ratio (IgG- Gal-ratio)	2 [137, 170]	I	I	1
IMP1	I	I	2 [161, 164]	I
Koc	I	I	2 [161, 164]	I
MIC	2 [129, 138]	I	1	I
NPM1	ı	2 [62, 70]	1	I
P16	1	2 [62, 70]	ı	I
Peroxiredoxin 6 (Prx6)	ı	I	2 [155, 156]	I
Other proteins				
CA19-9	35^a	20 ^b	4 [143, 168, 170, 171]	I

Table 3 continued

Biliary tract 2 [165, 167] 9 [141, 143, 144, 147, 148, 163, 168, 170, 171] 2 [165, 167] cancer 4 [142, 144, 147, 163] Oesophageal cancer 9 [25, 30, 46, 48, 53, 57, 59, 74, 171] 2 [144, 171] 8 [29, 33, 39-41, 43, 66, 76] 9 [29, 33, 38–41, 43, 66, 76] 6 [25, 31, 59, 73, 78, 84] Gastric cancer 3 [31, 59, 78] 2 [56, 68] 2 [24, 66] 7 [96, 102, 110, 112, 116, 127, 170] 4 [96, 112, 116, 121] 4 [92, 122, 123, 125] Pancreatic cancer 3 [125, 127, 129] 3 [94, 119, 135] 2 [112, 116] 3 [104–106] 3 [104-106]2 [107, 116] 2 [121, 129] 2 [94, 119] 2 [94, 135] 2 [94, 119] 2 [92, 123] Metalloproteinase 1 (TIMPbinding protein-2 (IGFBP2) Carcinoembryonic antigen Insulin-like growth factor-Apolipoprotein AII-ATQ Alpha-Fetoprotein (AFP) Apolipoprotein AII-AT Interleukin-13 (IL-13) Interleukin-8 (IL-8 or Fable 3 continued Pepsinogen II (PGII) Tissue Inhibitor of Interleukin-6 (IL-6) Interleukin-4 (IL-4) Pepsinogen I (PGI) (apoAII-ATQ) (apoAII-AT) CEACAM-1 Osteopontin CYFRA21-1 CXCL8) Biomarker CA242 (CEA) CA724 CA125

Table 3 continued

Biomarker	Pancreatic cancer	Gastric cancer	Oesophageal cancer	Biliary tract cancer
Matrix metalloproteinase-7 (MMP-7)	1	1	2 [155, 156]	1
Neuron-specific enolase (NSE)	2 [112, 116]	I	I	ı
Trefoil factor 1 (TFF1)	ı	2 [33, 39]	1	I
Trefoil factor 2 (TFF2)	1	2 [33, 39]	I	ı
Trefoil factor 3 (TFF3)	ı	2 [33, 39]	I	ı
Thrombospondin 2 (THBS2) 2 [109, 124]	2 [109, 124]	ı	I	ı
Vascular Endothelial Growth Factor (VEGF)	2 [94, 119]	ı	I	ı
Metabolic markers				
Histidine	3 [98, 118, 120]	ı	I	ı
Alanine	2 [98, 120]	ı	ı	ı
Asparagine	2 [98, 120]	1	ı	ı
Isoleucine	2 [98, 120]	1	I	ı
PC-594	2 [88, 126]	ı	ı	ı
Phosphatidylcholine-C18.0- C22.6	2 [88, 118]	ı	I	ı
Serine	2 [98, 120]	ı	ı	ı
Tryptophan	2 [98, 120]	1	1	ı

^a CA19-9 in pancreatic cancer: [89, 90, 92, 96, 97, 99, 101–103, 105–107, 109, 110, 112–114, 116–118, 121–127, 129, 132, 135, 137–139, 170]

^b CA19-9 in gastric cancer: [25, 27, 30–32, 34, 38, 46, 52, 53, 57–59, 65, 74, 78, 84, 168, 170, 171]

° CEA in gastric cancer: [25, 26, 30–32, 34, 38, 46, 48–50, 52, 53, 56–59, 65, 73–75, 78, 80, 84, 168, 170, 171]

Table 4 Biomarkers reported more than once for the same tumour type and panels adopting a single-gate (reversed-flow) design

References	Recruitment setting	Cases	Controls	Outcomes (Sensitivity, specificity, AUC where available)
1. Measures	of diagnostic performan	nce available for indi	vidual biomarkers, in studies ado	opting a single-gate design
Apolipoprote	in AII-AT/ATQ alone a	and in combination u	vith CA19-9 (pancreatic cancer)	
Honda et al. [106]	EPIC cohort (population-based study)	156 PaC Median age 58.1 (34.9–75.7) 53% male	213 HC Median age 58.0 (34.5–75.4) 53% male (matched to cases)	Measures for months prior to diagnosis (lag times): up to 6 months, > 6–18, 18, > 18–36 and > 36–40 months
		Staging: 13 localised, 73 metastatic, 69 NA		For ApoAII-AT/ATQ alone, 2 cut-off points Sensitivity, range 0.04–0.25 AUC, range 0.52–0.62 For ApoAII-AT/ATQ plus CA19-9, 2 cut-off points Sensitivity, range 0.07–0.57 Specificity, range 0.96–0.98
Honda et al. [105]	Cohort 1: National Cancer Centre Hospital	131 IDACP Mean age 68.8 (9.01) 55% male Staging: most at advanced stages	131 HC Mean age 62.5 (10.8) 52% male	AUC, range 0.56–0.78 Measures for ELISA and mass spectrometric analysis, also according to tumour staging For ApoAII-ATQ/AT alone, 1 cut-off point AUC, range 0.856–0.946
	Cohort 2: Seven Medical Institutions Cohort 3: NCI- EDRN pancreatic reference set	155 IDACP Age and sex NA Staging: majority advanced stages 98 PaC Age and sex NA Staging: all early stages	57 pancreatic disease other than IDACPAge and sex NA62 CP, 31 acute benign biliary obstruction, 61 HCAge and sex NA	For ApoAII-AT/ATQ plus CA19-9, 1 cut-off point each Sensitivity, 95.4% (cohort 2) Specificity, 98.3% (cohort 2)

Table 4 continued

References	Recruitment setting	Cases	Controls	Outcomes (Sensitivity, specificity, AUC where available)
Honda et al.	Cohort 1: National Cancer Hospital	Does not meet criteria as used	to calculate first measures of	Measures provided according to tumour staging
[104]	and Medical University Hospital	to calculate first measures of performance	performance	For ApoAII-AT/ATQ alone, 1 cut-off point
	_	Does not meet	December of the city of	AUC, 0.953 (cohort 3)
	Cohort 2: National Cancer Hospital	criteria as there were only 41 controls	Does not meet criteria as there were only 41 controls	For ApoAII-AT/ATQ plus CIII-0, and CA19-9, 1 cut-off point (cohort 4)
	Cohort 3:	52 PaC	53 HC and 58 CP	Sensitivity, range 91.60-94.20%
	Department of General Surgery	Mean age 63.1 (9.85)	HC mean age 39.1 (15.6), CP 50.3 (8.9)	Specificity, 93.22% (same for all)
		56% male	HC 59% male, CP 74% male	
		Staging NA		
	Cohort 4: Seven Medical Institutions	249 PDAC and 18 other malignant	128 HC, 38 benign tumour/cyst and 14 CP	
		tumour of the pancreas	HC mean 46.6 (16.8), benign tumour/cyst 63.5 (11.0), CP	
		PDAC mean age	60.2 (10.2)	
		64.4 (9.1), other 68.3 (9.7)	HC 65% male, benign tumour/cyst 45% male, CP 86% male	
		PDAC 59% male, other 67% male		
		Staging NA		
Pepsinogen (PGI and PGI/II ratio)	(gastric cancer)		
Gantuya	National Cancer	50 GC (54% w/	752 non-cancer (302 antrum limited CG and/or atrophy and 450 corpus CG and/or atrophy (77% w/ H. pylori Mean age: 53.8 (SD 1, 27–78) 31% male	For PGI, optimal cut-off point
et al. [29]	Centre Hospital	H. pylori) No information		Sensitivity, 70%
				Specificity, 70%
		on age and sex		AUC, 0.76
		Staging NA		For PGI/II ratio, optimal cut- off point
			2 - / V	Sensitivity, 66%
				Specificity, 65%
				AUC, 0.70

Table 4 continued

References	Recruitment setting	Cases	Controls	Outcomes (Sensitivity, specificity, AUC where available)
Kang et al. [40]	National University Hospital	380 GC (intestinal and diffuse type) Age and sex not	172 BGU, 119 DU, 107 dysplasia Age and sex not available for	Measures according to tumour type only (intestinal or diffuse)
		available for cases	controls only	For PGI, 1 cut-off point
		only No information on staging		Sensitivity, 77.7% (intestinal), 64.7% (diffuse)
				Specificity, 20.2% (intestinal), 20.2% (diffuse)
				For PGI/II ratio, 1 cut-off point
				Sensitivity, 62.3% (intestinal), 55.8% (diffuse)
				Specificity, 61.0% (intestinal), 61.0% (diffuse)
Kikuchi et al. [41]	University Outpatient Clinic	122 GC Age: 68.2 years (9.7)	16 GU or DU, 17 superficial gastritis, 66 CAG, 79 no abnormality	Measures combining normal and non-malignant conditions
		74% male	Age: 56.2 years (14.9)	Negative or positive PG test
		Staging NA	55% male	For PGI and PGI/II ratio, strict or conventional cut-off point
				Sensitivity, 41.3% (strict), 77.9% (conventional)
				Specificity, 90.4% (strict), 61.8% (conventional)
Yanaoka	Employees in annual	63 GC	5146 HC	or PGI and PGI/II ratio, 3 cut-
et al. [76]	health screening programme	Age: 50.3-51.8	Mean age: 49.2 (4.7)	off points
	programme	(mean range)	100% male	Sensitivity, range 27.0–58.7%
		100% male		Specificity, range 73.4–92.0%
		86% early, 14% late stages		

Table 4 continued

References	Recruitment setting	Cases	Controls	Outcomes (Sensitivity,
				specificity, AUC where available)

2. Measures of diagnostic performance available for established biomarkers combined with novel biomarkers not shown above, in studies adopting a single-gate design

CA19-9 (pancreatic cancer)

GIII)) (pui				
O'Brien et al. [121]	UKCTOCS screening cohort	101 PaC Age NA for validation 100% female Staging NA	184 HC Age N/A for validation 100% female	Measures according to time to diagnosis: 0–4 years, 0–2 years; 1–4 years For CA19-9 (4 cut-off points) plus CA125 (3 cut-off points) Sensitivity, range 23.1–53.1% Specificity, range 71.6–92.6%
Tavano et al. [132]	Hospital (Gastroenterology, Surgery & Oncology)	74 PaC Median age 69 (61–76) 54% male Staging NA for validation	117 HC Median age 62 (55–70) 45% male	For CA19-9 plus miR-1290, 1 cut-off point (each) Sensitivity, 83.8% Specificity, 96.6% AUC, 0.923
Zhou et al. [138]	Gastroenterology Department in Hospital	152 PaC Mean age 56 (SD 13.5) 67% male Staging: 5 IA, 12 IB, 36 IIA, 20 IIB, 40 III, 39 IV	96 HC, 91 CP, 20 premalignancies Mean age: HC 58 (7.6), CP 58 (15.0), pre-malignancies 60 (11.3) HC 75% male; CP 57% male; pre-malignancy 75% male	For CA19-9 plus MIC-1 and ULBP2, 1 cut-off point (each) AUC 0.982 (PaC and CP only) For CA19-9 plus MIC-1, 1 cut-off point (each) AUC 0.932 (PaC and CP only) For CA19-9 plus ULBP2, 1 cut-off point (each) AUC 0.953 (PaC and CP only)

Table 4 continued

References	Recruitment setting	Cases	Controls	Outcomes (Sensitivity,
				specificity, AUC where
				available)

3. Measures of diagnostic performance available for a panel only in studies adopting a single-gate design (all reversed-flow)

Different panels (pancreatic cancer)^a

33 1	4			
Balasenthil et al. [89]	NCI-EDRN pancreatic reference set	98 PaC (52 w/o diabetes or pancreatitis) Age and sex not available Staging: 7 IA, 8 IB, 1 II, 40 IIA and 42 IIB	62 CP, 31 acute biliary obstruction, 61 HC (50 w/o diabetes or pancreatitis) Age and sex not available	Measures for PaC vs. HC, PaC vs. CP, PaC w/o diabetes or pancreatitis vs. HC w/o diabetes or pancreatitis, and according to staging For CA19-9 plus TFPI and TNC-FN III-C, 2 cut-off points Sensitivity, range 0.73–0.81 Specificity, range 0.71–0.84 AUC, range 0.75–0.89
Mellby et al. [119]	Patients referred to Medical Centre for symptomatic pancreatic disease	tre for validation (US cohort) sease 143 PaC patients Median age only by staging; range 24–87	219 HC, 57 CP HC median age 63.0 (24–86), CP 55.5 (32–81) HC 53% male, CP 46% male	Measures available for stages I + II combined For 29-panel signature (no established biomarkers):
				Sensitivity, 95% Specificity, 93%
				AUC, 0.963 (PaC vs. HC) and
		57% male		0.840 (Pac vs. CP)
		Staging: 15 I, 75 II, 15 III and 38 IV		

ACG atrophic chronic gastritis, ApoAII-AT/ATQ apolipoprotein AII-AT/ATQ, apoCIII-0 apolipoprotein CIII-0, BGU benign gastric ulcer, DU duodenal ulcer, CG chronic gastritis, CP chronic pancreatitis, EPIC European Prospective Investigation into Cancer and Nutrition, GC gastric cancer, GU gastric ulcer, IDACP invasive ductal adenocarcinoma of pancreas, MIC macrophage-inhibitory cytokine 1, MPV mean platelet volume, NA not available, NCI-EDRN National Cancer Institute Early Detection Research Network, PaC pancreatic cancer, PDAC pancreatic ductal adenocarcinoma, PDW platelet distribution width, PGI/II serum pepsinogen I/II, PPV positive predictive value, TFPI plasma tissue factor pathway inhibitor, NTC-FN III-C tenascin-C, UKCTOCS UK Collaborative Trial of Ovarian Cancer Screening, ULBP2 UL16 binding protein 2

^a Leelawat et al. [166] also adopted a reversed-flow design but was not added as it was the only study investigating CA19-9 for cholangiocarcinoma

measures of diagnostic performance, adopting a recommended single-gate design. Heterogeneity in methods, populations, biomarkers, outcomes and comparisons precluded meta-analysis. Applying novel biomarkers for the early detection of upper GI cancers is therefore at an early stage of matureness: few have been extensively evaluated and evaluations have almost exclusively focussed on high-prevalence populations. Further evaluation of the most promising biomarkers in low-prevalence populations is needed before extensive adoption into routine clinical practice can be recommended.

While other reviews have investigated biomarkers used for early cancer detection [19, 172], few have considered the evidence in the context of future application of tests in lowprevalence populations, the likely target for clinical application [12, 13]. To our knowledge, this is the first review to do so for upper GI cancers. The four novel and one established biomarkers we highlight in this review were evaluated in a mix of high- and low-prevalence populations, including hospital patients, general population cohorts, screening populations (both high and average cancer risk), and patients presenting with symptoms. We did not identify any studies reporting outcomes relevant to feasibility, acceptability, benefits and harms, nor health economics as initially planned in the review protocol (i.e. phase 3 studies and beyond in the CanTest framework). The best performing biomarkers for pancreatic cancer, with an AUC between 56% and 94%, were ApoAII-ATQ/AT alone, CA19-9 plus miR-1290, MIC-1 and ULPB2, and Mellby et al.'s [119] 29-panel signature. These may be ready for trials and other phase 3 studies, single or in combination, in low-prevalence populations. We did not identify any novel biomarkers with similar AUCs for gastric, biliary tract or oesophageal cancers.

A previous review investigating the role of pepsinogens in early detection of gastric cancers reported that they had only moderate capacity to detect gastric cancer [173]. Another review on early pancreatic cancer detection highlighted that no single biomarker has yet translated to clinical use and suggested the use of 'robust panels of biomarkers' [9]. This review

confirms that more research is required before we have sufficient evidence about biomarkers for upper GI cancers to warrant their adoption into clinical practice.

We identified several important methodological limitations within the biomarker studies to date. These include large numbers of biomarkers analysed in parallel during discovery studies, increasing risk of falsely positive results; limited sample sizes; evaluation of "extreme" cases; limited external, independent validation; and selective reporting for validation (several alternatives analyses and combinations, use of several cut-off points and overoptimistic interpretation of the data) [12]. Together with use of two-gate rather than recommended single-gate designs, these could all lead to over-inflated measures of performance. Population characteristics were often provided as supplementary data, with little discussion of potential selection bias and other sources of uncertainty. We also excluded relevant studies when we could not obtain sufficient information on an individual tumour type; this was the case for the CancerSeek tool [174]. Adoption of reporting guidelines [175] and development of early cancer detection collaborations [15, 18] could be useful strategies to address these issues.

This review offers a comprehensive overview of the available evidence. It benefitted from having a multidisciplinary team of experts, a broad search strategy, independent screening, and classifications checked by senior team members. Since meta-analysis was not feasible nor appropriate, we had to use text and tables to synthesise the evidence. We did not include studies investigating biomarkers as part of risk prediction models or risk assessment tools. These studies have strong potential to be used in the community and should be investigated in a separate systematic review. Recent reviews indicate that only including studies in English has minimal impact on review conclusions [176, 177]. We believe this is also the case for this review, particularly due to the overall lack of evidence on biomarkers ready to be evaluated in low-prevalence settings. Although we did not formally appraise risk of bias, we identified several quality and methodological issues, indicating that challenges already highlighted

in the literature persisted over time [12]. Finally, due to the large amount of evidence on biomarker development and evaluation, we believe the field could benefit from a "living systematic review"; this refers to high quality, up-to-date online summaries of evidence which can be constantly updated as new research becomes available [178].

The studies we identified focused on measures of diagnostic performance, which is reasonable given the phase of development for most of them. The CanTest Framework [15] can help guide studies aiming to build much needed evidence on later phases of biomarker development, focussing on impact on clinical decisionmaking, patient, health system and economic outcomes.

CONCLUSION

There is a large body of evidence on biomarkers being developed for the detection of upper GI cancers, but relatively few have yet to demonstrate their validity or clinical utility in settings where cancer prevalence is low. Early detection of colorectal cancer already benefits from biomarkers that can be used across different populations. This is the case for the faecal immunochemical test (FIT), which is recommended for use in primary care in Spain, Australia and the United Kingdom, in addition to being effective at mass population screening programmes, using different cut-off points [179, 180]. It took several decades from FIT development to generate evidence for its costeffectiveness as a screening test for colorectal cancer. Its role in the assessment of patients in primary care with lower GI symptoms is still being evaluated. Biomarkers for upper GI cancer remain in their infancy but there are a few which show promise and require further evaluations. Ultimately, they may be able to contribute to improving outcomes for upper GI cancers through earlier detection.

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Compliance with Ethics Guidelines. This article is based on previously conducted studies and does not contain any studies with human

participants or animals performed by any of the authors.

Data Availability. The datasets generated during and/or analysed during the current study are available from the corresponding author on reasonable request.

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