ORIGINAL RESEARCH

Prognostic value of circulating tumor cells detected with the CellSearch System in patients with gastric cancer: evidence from a meta-analysis

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Background: Circulating tumor cells (CTCs) have been proposed as a marker for predicting the prognosis of cancer. However, the prognostic value of CTCs detected with the CellSearch System in patients with gastric cancer (GC) remains controversial. We performed a meta-analysis of available studies to investigate this topic.

Methods: Two authors systematically searched the studies independently in PubMed, Science Citation Index, Cochrane Database, Embase, and the references in relevant studies (up to September 2017) using keywords. Our meta-analysis was performed in Stata software, version 12.0 (2011; Stata Corp, College Station, TX, USA), with the risk ratio (RR), hazard ratio (HR), and 95% CI as the effect measures. Subgroup analyses and meta-regression were also conducted.

Results: Seven studies (including eight sets of data) containing 579 patients with GC from four countries were included in this meta-analysis. The pooled results showed CTC-positive status detected by the CellSearch System was significantly associated with poor overall survival (HR =2.09, 95% CI [1.71, 2.55], P<0.001, I^2 =31.5%) and progression-free survival (HR =2.11, 95% CI [1.25, 3.57], P=0.005, I^2 =75.6%) of patients with GC, regardless of sampling time. The disease control rate of CTC-positive group was lower than that of CTC-negative group for both baseline and intra-therapy, although no statistical difference existed at both sampling time points (baseline: 69.5% versus 81.8%, RR=0.79, 95% CI [0.54, 1.16], P=0.23, I^2 =68.0%; intra-therapy: 50.0% versus 85.9%, RR=0.24, 95% CI [0.02, 3.13], P=0.28, I^2 =87.4%).

Conclusion: Our meta-analysis demonstrated that CTCs detected with the CellSearch System from the peripheral blood had significant prognostic value and might predict poor response to chemotherapy for patients with GC.

Keywords: circulating tumor cells, gastric cancer, chemotherapy, prognosis, meta-analysis

Introduction

Gastric cancer (GC) ranks the fourth most common cancer and the third leading cause of cancer-associated mortality worldwide, with over 951,600 new cases and 723,100 deaths estimated to have occurred (GLOBOCAN 2012).¹ Although recent diagnosis and treatment advances have improved the clinical outcome of GC patients, the 5-year survival rate is still <30%.² Also, recurrence or metastasis after curative resection is the main reason for GC-related deaths.³ Unfortunately, the lack of effective predictors prevents doctors from identifying recurrence and metastasis timely during therapy and review, which leads to missing the best time of intervention and thus affects the therapeutic effect. Nowadays, circulating tumor cells (CTCs), which shed from

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the primary tumor mass and are circulated in the blood, have gradually been accepted as a new and efficient prognostic marker to evaluate the relapse and metastasis, monitor the therapeutic responsiveness, and predict the prognosis, due to their advantage of earlier and more reproducible indication of disease status than that of current imaging methods.⁴

Numerous studies have shown that the presence of CTCs in the peripheral blood (PB) is a poor prognostic indicator of overall survival (OS) and progression-free survival (PFS) in patients with GC.⁵⁻⁸ In those studies, the methods of detecting CTCs were predominantly immunocytochemistry (ICC)^{5,6} and reverse transcription-polymerase chain reaction (RT-PCR).^{7,8} However, these detection methods vary across laboratories and the optimal cutoff value for CTCs has not yet been confirmed. Currently, the CellSearch System (Veridex, Raritan, NJ, USA), a semi-automated immunological assay for enumeration of CTCs based on the epithelial cell adhesion molecule, is the first and only standardized system approved by the US Food and Drug Administration for applying to detect CTCs in patients with breast, prostate, and colorectal cancer.9-11 Moreover, the prognostic significance of CTCs detected with CellSearch System in patients with breast, prostate, and colorectal cancer had been quantitatively and comprehensively summarized by previous meta-analyses.¹²⁻¹⁴ With regard to GC, although there were many studies designed to find out the relationship between CTCs detected with the CellSearch System and prognosis, the lack of statistical power together with their different study design and results limited its individual clinical value and prognostic effect. Therefore, a combined analysis of available studies on this topic is required to provide a more precise estimate of the prognostic relevance of CTC detection in the PB of patients with GC.

The aim of our study was to conduct a meta-analysis to quantitatively summarize the clinical evidence and prognostic value of CTCs detected with the standardized CellSearch System in GC patients.

Methods

Data sources and search strategy

Our meta-analysis was conducted according to the recommendations of the Preferred Reporting Items for Systematic Reviews and Meta-analyses statement.¹⁵ Two authors (CG Yang and K Zou) systematically searched the literature independently from the following databases: PubMed, MEDLINE, EMBASE, Science Citation Index, and Cochrane Library (up to September 2017). The main keywords and MeSH terms used were "circulating tumor cells", "disseminated tumor cells", "isolated tumor cells", "occult tumor cells", "gastric cancer", "stomach carcinoma", "gastrointestinal cancer", "digestive cancer", and "CellSearch System". An additional search through Google Scholar and the clinical trial registration website was conducted to obtain information identifying other potentially relevant publications. The reference lists of the relevant studies (review studies and included studies) were also checked for potentially relevant articles. Discrepancies were resolved by the third author (ZW Yuan).

Inclusion and exclusion criteria

To be included in the meta-analysis, studies had to match the following criteria: 1) used only the CellSearch System to explore the prognostic role of CTC detection in GC patients; 2) at least one of the outcome measures of interest was reported in the article or could be calculated from the published data; and 3) samples being collected from the PB. When multiple studies were published with the same patient population, we included the most informative study.

Studies were excluded from the meta-analysis if 1) the outcomes of interest were not reported or could not be calculated from the original published data; 2) the samples came from the lymph nodes, bone marrow, mesenteric/ portal blood, or peritoneal cavity; 3) studies were based on overlapping patients; and 4) enough data was not available after contacting the original author or magazine.

Data extraction

Two investigators (CG Yang and K Zou) independently extracted the following data from each eligible study: the first author's name, year of publication, country of patients, characteristics of the study population (ie, number, sex, age), sampling time, cutoff value, positive rate of CTC detection, and prognostic outcomes (OS and PFS), hazard ratio (HR), and disease control rate (DCR) of chemotherapy. According to the Response Evaluation Criteria In Solid Tumors guideline (complete response [CR], partial response [PR], stable disease [SD], and progressive disease [PD]),¹⁶ DCR=(CR+PR+SD)/(CR+PR+SD+PD). For the studies with multiple arms (ie, training set and validation set), each of them was considered as an independent data set. For the studies with multiple time points (ie, baseline and intra-therapy), we used data from "baseline" samples prior to the data from "intra-therapy" samples because these data were usually dependent. If the HR and its 95% CI were not reported directly in the original article, these values were calculated from available data using the methods reported by

Tierney et al.¹⁷ Disagreements between two reviewers were resolved by discussion or consultation.

Quality assessment

Two reviewers (CG Yang and K Zou) used the Newcastle– Ottawa scale, which was recommended by the Cochrane Library for nonrandom controlled studies, to assess the quality of included studies independently.¹⁸ Any disagreements on quality assessment were resolved via comprehensive discussion. Based on the Newcastle–Ottawa scale criterion, a score of 5–9 means high quality, whereas a score of 1–4 means low quality.

Statistical analysis

Stata 12.0 software (Stata Corp, College Station, TX, USA) was used for all analyses in our meta-analysis. The estimated HR was used to evaluate the prognostic indicators (PFS and OS) as demonstrated by Parmar et al,¹⁹ and HR >1 denoted more deaths or progression in CTCs-positive arm. The estimated risk ratio (RR) was used to evaluate the DCR. All statistical values were reported with 95% CIs, and the twosided P-value threshold for statistical significance was set at 0.05. Heterogeneity among the studies was calculated with the *Q* test and I^2 statistic, and the I^2 value indicated the degree of heterogeneity.²⁰ A *P*-value < 0.1 for the *Q* statistic and/ or $I^2 > 50\%$ were considered significant heterogeneity, and a random-effects model was used. Otherwise, a fixed-effects model was used.²¹ Furthermore, subgroup analyses were conducted according to the difference in the data retrieved, such as year of publication, country of patients, sampling time, cutoff value, positive rate of CTC detection, and quality of included studies. Meta-regression was performed to explore the potential variables that contributed heterogeneity. Additionally, Galbraith plot was used to further explore which study would contribute substantial heterogeneity to our meta-analysis. Additionally, potential publication bias was detected by Begg's and Egger's test.²² P<0.05 was considered to be significant publication bias.

Results

Baseline characteristics of the included studies

According to the above-mentioned retrieval method, a total of 69 studies were initially identified during a systematic literature search. By screening the titles and abstracts, 61 studies were excluded and 8 potential studies were retrieved. One study was then excluded after full-text articles, as it had insufficient data. Finally, seven studies were eligible for our meta-analysis (Figure 1).

Seven eligible studies, including eight sets of data, contained 579 patients with GC.^{23–29} These studies came from four countries (China, Japan, Korea, and UK) and were published between 2008 and 2016. According to the sampling time points, four studies only assessed CTCs at baseline,^{25–28} one study only assessed CTCs at intra-therapy,²³ and two studies assessed CTCs combined at both time points.^{24,29} HRs for OS and PFS were available in eight sets of data from seven studies^{23–29} and six sets of data from six studies,^{24–29} respectively. Among all the included studies, two studies were of low quality and the other five studies were of high quality. The baseline characteristics and results of quality assessment of included studies are summarized in Table 1.

Impact of CTCs detection on prognostic outcomes (OS and PFS)

The pooled analysis showed that CTC-positive status detected by the CellSearch System in GC patients was associated with poor OS (HR =2.09, 95% CI [1.71, 2.55], P<0.001; Figure 2A), without any statistical significance in between-study heterogeneity (I^2 =31.5%, P=0.177). The HRs for PFS were available in six sets of data from six studies; the result of pooled analysis using the random-effects model showed that CTC positivity was also associated with poor PFS (HR =2.11, 95% CI [1.25, 3.57], P=0.005; Figure 2B), and between-study heterogeneity was obvious (I^2 =75.6%, P=0.001).

Furthermore, we stratified the included studies based on variables (year of publication, country of patients, sampling time, cutoff value, positive rate of CTC detection, and quality of included studies). The results of pooled analysis demonstrated that CTC positivity was a significant prognostic effect for OS and PFS for most of subgroups (Table 2). Additionally, the between-study heterogeneity was eliminated in subgroups by exclusion of studies coming from countries other than Japan, and with a cutoff value ≤ 2 , positive rate of CTC detection $\geq 41.9\%$, or low quality for OS. However, all the above variables did not positively contribute to the heterogeneity for PFS.

Impact of CTC detection on DCR

Only three studies evaluated the correlation between CTCs and DCR in patients receiving adjuvant chemotherapy.^{24,25,29} The pooled results demonstrated that the DCR for the CTC-positive groups was lower than that for the CTC-negative groups at baseline and intra-therapy, although there was no statistical difference (baseline: 69.5% versus 81.8%,



Figure I Flow chart of study selection.

Note: Copyright ©2009. Adapted from Moher D, Liberati A, Tetzlaff J, Altman DG. PRISMA Group. Preferred reporting items for systematic reviews and meta-analyses: the PRISMA statement. *PLoS Med.* 2009;6(7):e1000097.³⁶

Abbreviation: PRISMA, Preferred Reporting Items for Systematic Reviews and Meta-analyses.

RR=0.79, 95% CI [0.54, 1.16], *P*=0.23, *I*²=68.0%; intratherapy: 50.0% versus 85.9%, RR=0.24, 95% CI [0.02, 3.13], *P*=0.28, *I*²=87.4%; Figure 3).

The source of heterogeneity

To explore the potential sources of heterogeneity, we conducted a meta-regression that considered the variables of year of publication, country, sampling time, cutoff value, and quality of included studies. As shown in Table 3, in a univariate analysis, none of these variables was significantly correlated with the heterogeneity across studies in OS and PFS. However, the results of meta-regression may be affected by the limited number of studies in the analysis (n=6). Therefore, we used Galbraith plot to further explore the source of heterogeneity in PFS, and the result showed that the training set of the study by Uenosono et al²⁸ might have mainly contributed substantial heterogeneity to PFS (Figure 4A). After omitting this study, the pooled HR was not affected obviously (HR =1.73, 95% CI [1.25, 2.41], P=0.001; Figure 4B), but the heterogeneity for PFS dropped to an insignificant level (from I^2 =75.6%, P=0.001 to I^2 =29.5%, P=0.225; Figure 4C).

Assessment of publication bias

Potential publication bias was assessed by Begg's and Egger's tests. There was no evidence of publication bias for the pooled analysis of OS ($P_{\text{Begg's}}$ =0.266, Figure 5A; $P_{\text{Egger's}}$ =0.175, Figure 5B) and PFS ($P_{\text{Begg's}}$ =1.000, Figure 5C; $P_{\text{Egger's}}$ =0.899, Figure 5D).

Discussion

Although curative surgery in combination with chemoradiotherapy and targeted therapy has greatly developed in recent years, the occurrence of metastasis and relapse of GC is still a great challenge for both surgeons and patients. The presence

Table I Baseline characteristics	of inclue	ded studies									
Reference	Year	Country	Number	Age	Tumor	Sampling time	Cutoff	Positive	End	Treatment regimens	Quality
			(male/female)	(years) ^a	stage ^b			rate, n (%)	point		
Hiraiwa et al ²³	2008	Japan	27 (NR)	NR	≥	Intra-therapy	≥2/7.5 mL	15 (55.6)	SO	Chemotherapy	Low
Matsusaka et al ²⁹	2010	Japan	52 (44/8)	62 (24–78)	≥	Baseline	≥4/7.5 mL	17 (32.7)	OS, PFS	Chemotherapy	High
			51 (NR)	NR		Intra-therapy (2 weeks)	≥4/7.5 mL	7 (17.3)	OS, PFS		
			48 (NR)	NR		Intra-therapy (4 weeks)	≥4/7.5 mL	9 (18.8)	OS, PFS		
Uenosono et al ²⁸ (1) (training set)	2013	Japan	148 (99/49)	NR	Ē	Baseline	≥I/7.5 mL	16 (10.8)	OS, PFS	Surgery	High
Uenosono et al ²⁸ (2) (validation set)	2013	Japan	103 (NR)	NR	≥	Baseline	≥1/7.5 mL	62 (61.8)	SO	Chemotherapy	High
Sclafani et al ²⁷	2014	Я	18 (16/2)	61 (38–78)	≥	Baseline	≥2/7.5 mL	8 (44.4)	OS, PFS	Chemotherapy	Low
Lee et al ²⁵	2015	Korea	95 (63/32)	57 (36–85)	≥	Baseline	≥5/7.5 mL	27 (28.4)	OS, PFS	Chemotherapy	High
Okabe et al ²⁶	2015	Japan	136 (87/49)	NR	≥⊢	Baseline	≥I/7.5 mL	25 (18.4)	OS, PFS	Surgery + chemotherapy	High
Li et al ²⁴	2016	China	136 (89/47)	59 (25–80)	≥	Baseline	≥3/7.5 mL	57 (41.9)	OS, PFS	Chemotherapy	High
			106 (NR)	NR		Intra-therapy	≥3/7.5 mL	26 (24.5)	OS, PFS		
Notes: ^a Median (range). ^b The seventh edit	tion of the	AJCC cancer s	staging.								

Abbreviations: AJCC, American Joint Committee on Cancer; NR, not reported; OS, overall survival; PFS, progression-free survival.

of CTCs in the PB may provide a reasonable explanation for metastasis and relapse by the "seed and soil" theory.³⁰ From the clinical perspective, CTCs enumeration can supply important prognostic information for the patients on solid tumor. Besides, due to its advantages of saving time and cost and providing comfort and repeatability, CTCs detection from PB can be used as an effective tool for monitoring therapeutic effect. Several previous meta-analyses have reported the prognostic and clinicopathologic significance of CTCs detection from PB in patients with GC; however, the methods of detecting CTCs were mainly RT-PCR and ICC in the studies included by them.^{31–33} Recently, the clinical utility of CTC detection in the PB with the standardized CellSearch System has been demonstrated in several studies.²⁴⁻²⁶ Therefore, it is essential to conduct a meta-analysis to quantitatively summarize the clinical value of CTCs detected with the standardized CellSearch System in GC patients.

CTCs, CellSearch System, and gastric cancer

To our knowledge, this was the first comprehensive metaanalysis to assess the prognostic and predictive value of CTC detection with the CellSearch System only in GC. Although the previous meta-analysis by Zou et al explored the clinicopathologic significance of CTC detection with cytological techniques (including CellSearch) in GC and the selected studies of our meta-analysis matched part of their included studies,³¹ there are some important differences between the two studies. First, we processed the data more finely compared to the previous study. In fact, the study by Uenosono et al included two independent sets of data (training set and validation set) and both of them had to be incorporated into the meta-analysis;²⁸ however, Zou et al did not do so, which affected the accuracy of their results. More importantly, we further explored the between-study heterogeneity through subgroup analysis, meta-regression, and Galbraith plot and eventually concluded that the training set of Uenosono et al²⁸ was the main source of the heterogeneity in PFS. In a previous study, the between-study heterogeneity was not obvious, however, as it did not include the training set of Uenosono et al.²⁸ As we all know, when between-study heterogeneity is obvious, clarifying the source of heterogeneity is the premise of conducting a meta-analysis. Therefore, our results were more reliable than the previous study, although both studies came to the same conclusion. In our meta-analysis, the results suggested that patients with CTC positivity have poorer OS and PFS than CTC-negative patients. Moreover, these results were also found in most of the subgroups when we stratified the included studies based on variables (Table 2). Additionally, the pooled analysis showed the DCR of CTC-positive groups was lower than that of CTC-negative groups for both

A	Study ID		HR (95% CI)	Weight (%)
	Hiraiwa et al ²³ (2008) —	\rightarrow	3.35 (1.32, 8.47)	4.57
	Matsusaka et al ²⁹ (2010)		1.54 (0.79, 3.00)	8.92
	Uenosono et al ²⁸ (1) (2013) -	•	2.77 (1.82, 4.21)	22.53
	Uenosono et al ²⁸ (2) (2013)		2.05 (1.33, 3.17)	20.89
	Sclafani et al ²⁷ (2014)		0.52 (0.18, 1.50)	3.51
	Lee et al ²⁵ (2015)		2.06 (1.26, 3.38)	16.18
	Okabe et al ²⁶ (2015)	•	2.20 (1.16, 4.17)	9.62
	Li et al ²⁴ (2016)		1.97 (1.15, 3.36)	13.79
	Overall (<i>I</i> ² =31.5%, <i>P</i> =0.177)	>	2.09 (1.71, 2.55)	100
	0.118	8.47		
В	Study ID		HR (95% CI)	Weight (%)
	Matsusaka et al ²⁹ (2010)		1.07 (0.52, 2.21)	15.94
	Uenosono et al ²⁸ (1) (2013)	\longrightarrow	7.72 (3.84, 15.53)	16.27
	Sclafani et al ²⁷ (2014)	_	1.06 (0.37, 3.03)	11.93
	Lee et al ²⁵ (2015)	•—	2.58 (1.56, 4.26)	18.91
	Okabe et al ²⁶ (2015)		2.14 (1.19, 3.85)	17.77
	Li et al ²⁴ (2016)		1.54 (0.95, 2.49)	19.18
	Overall (<i>I</i> ² =75.6%, <i>P</i> =0.001)	>	2.11 (1.25, 3.57)	100
		1		

Figure 2 Summary estimates of HR for OS and PFS of GC patients with CTC positivity.

Notes: (A) Forest plot of OS. (B) Forest plot of PFS. Weights are from random-effects analysis.

Abbreviations: CTCs, circulating tumor cells; GC, gastric cancer; HR, hazard ratio; OS, overall survival; PFS, progression-free survival.

baseline and intra-therapy, indicating that CTCs detected at baseline or intra-therapy might predict the response to chemotherapy. However, no statistical difference existed in both baseline and intra-therapy time points. This was inconsistent with the results of a previous meta-analysis by Huang et al,³² which might be due to the smaller studies used in our metaanalysis to evaluate the correlation between CTC counts and tumor responses on imaging according to the Response Evaluation Criteria In Solid Tumors criteria.^{24,25,29} Taken together, these results indicated that CTCs detected by the CellSearch System have great clinical value in evaluating prognosis, guiding therapeutic decision, and monitoring therapeutic effect for GC patients. For the CTC-positive patients, more aggressive treatment and frequent efficacious assessments might be needed.

Compared to traditional ICC and RT-PCR techniques, the CellSearch System has the advantages of easy operation, time effectiveness, and higher specificity and reproducibility for

Variable	n	OS				n	PFS			
		HR (95% CI)	P-value ^a	1 ²	P-value ^b		HR (95% CI)	P-value ^a	1 ²	<i>P</i> -value ^b
Year										
> Median	4	1.84 (1.36, 2.49)	<0.001	50.2%	0.111	3	2.02 (1.48, 2.76)	<0.001	8.1%	0.337
\leq Median	4	2.30 (1.77, 2.99)	<0.001	0.7%	0.388	3	2.11 (0.52, 8.51)	0.293	88.8%	< 0.001
Country		. ,					. ,			
Japan	5	2.28 (1.79, 2.91)	<0.001	0.0%	0.552	3	2.61 (0.89, 7.63)	0.081	87.1%	<0.001
Other	3	1.75 (1.24, 2.47)	0.001	64.6%	0.060	3	1.80 (1.14, 2.84)	0.011	40.0%	0.189
Sampling time										
Baseline	7	2.04 (1.67, 2.50)	<0.001	34.7%	0.163	6	2.11 (1.25, 3.57)	0.005	75.6%	0.001
Intra-therapy	I.	3.35 (1.32, 8.47)	0.011	-	_	0	-	_	-	_
Cutoff value ^d										
> Median	3	1.90 (1.38, 2.61)	<0.001	0.0%	0.780	3	1.70 (1.06, 2.73)	0.029	54.1%	0.113
\leq Median	5	2.22 (1.72, 2.86)	<0.001	56.3%	0.058	3	2.72 (0.93, 7.94)	0.067	83.4%	0.002
Positive rate ^e										
\geq Median	4	1.91 (1.41, 2.58)	<0.001	58.9%	0.063	3	1.33 (0.92, 1.94)	0.131	0.0%	0.641
< Median	4	2.24 (1.72, 2.91)	<0.001	0.0%	0.512	3	3.39 (1.67, 6.88)	0.001	76.7%	0.014
Quality										
High	6	2.15 (1.75, 2.65)	<0.001	0.0%	0.770	5	2.32 (1.32, 4.09)	0.004	78.6%	0.001
Low	2	1.49 (0.74, 3.00)	0.264	85.1%	0.010	I	1.06 (0.37, 3.03)	0.914	-	_
Overall	8	2.09 (1.71, 2.55)	<0.001	31.5%	0.177	6	2.11 (1.25, 3.57)	0.005	75.6%	0.001

Table 2 Results of subgroup analyses on OS and PFS

Notes: ^aP-value for estimates of HR. ^bP-value for heterogeneity. ^cThe median year of OS and PFS was 2013 and 2014, respectively. ^aThe median cutoff value of OS and PFS was 2 and 2, respectively. ^cThe median positive rate of OS and PFS was 41.9% and 31.7%, respectively. –, not available. **Abbreviations:** HR, hazard ratio; OS, overall survival; PFS, progression-free survival.

CTCs enrichment. Although our meta-analysis of studies had used the CellSearch System to reduce the heterogeneity caused by different detection methods, a significant heterogeneity was still found in our meta-analysis, specifically in the pooled analysis of PFS. In addition, subgroup analyses and metaregression were conducted to explore the potential sources of heterogeneity; however, the results failed to clarify the heterogeneity among the involved studies. Furthermore, the

Study ID	RR (95% CI)	Weight (%)
Baseline		
Li et al ²⁴ (2016)	0.93 (0.81, 1.06)	28.99
Lee et al ²⁵ (2015)	0.52 (0.30, 0.91)	17.81
Matsusaka et al ²⁹ (2010)	0.86 (0.54, 1.35)	20.63
Subtotal (<i>I</i> ²=68.0%, <i>P</i> =0.044)	0.79 (0.54, 1.16)	67.43
Intra-therapy		
Li et al ²⁴ (2016)	• 0.81 (0.65, 1.01)	27.29
Matsusaka et al ²⁹ (2W) (2010)	• 0.08 (0.01, 1.24)	1.78
Matsusaka et al ²⁹ (4W) (2010)	0.14 (0.02, 0.89)	3.51
Subtotal (<i>I</i> ² =87.4%, <i>P</i> =0.000)	0.24 (0.02, 3.13)	32.57
Overall (<i>I</i> ²=79.8%, <i>P</i> =0.000)	0.71 (0.49, 1.03)	100
0.0057	1 175	

Figure 3 Summary estimates of RR for DCR of patients with CTC positivity. Note: Weights are from random-effects analysis.

Abbreviations: CTCs, circulating tumor cells; DCR, disease control rate; RR, risk ratio; W, week.

Variable	OS				PFS			
	Coef	SE	P-value	Adjusted R ²	Coef	SE	P-value	Adjusted R ²
Year	-0.0234	0.0652	0.731	-27.06%	0.0392	0.1679	0.827	-34.46%
Country	-0.0588	0.1123	0.619	-22.55%	-0.1298	0.2703	0.656	-28.19%
Sampling time	-0.4929	0.6007	0.443	-10.16%	-	_	_	-
Cutoff value	-0.0555	0.0822	0.525	-16.89%	-0.1640	0.2014	0.461	-12.55%
Quality	0.3685	0.4615	0.455	-10.11%	0.7849	0.9145	0.439	-5.75%

Table 3 Results of meta-regression on OS and PFS

Notes: Adjusted R², proportion of between-study variance explained. –, not available.

Abbreviations: Coef, coefficient; OS, overall survival; PFS, progression-free survival; SE, standard error.

Galbraith plot showed that heterogeneity in the pooled analysis PFS was mainly caused by the training set of the study by Uenosono et al.²⁸ The reason might be that the positive rate of CTC detection was relatively low, even with more patients included in this study.

Potential heterogeneity may also result from difference in sampling time, standard of CTC positivity, treatment regimens, or clinicopathologic data of included patients. Theoretically, intra-therapy CTC status may provide relatively more information because it reflects the combined information on baseline CTCs and tumor cells' release during surgical manipulation.³⁴ The study by Ikeguchi and Kaibara found that CTC-positive patients had better prognosis than CTC-negative patients if the blood samples were postoperatively collected within 48 hours.³⁵ However, owing to loss of survival microenvironment in PB, these freshly released CTCs may be rapidly apoptotic and release mass tumor antigens, leading to certain degree of detection bias. Additionally, there is no consensus on the optimal cutoff value for CTCs in the PB for predicting the prognosis of patients with GC. Although most studies used the cutoff value of CTCs $\geq 1/7.5$ or $\geq 2/7.5$ mL of PB in our meta-analysis and subgroup analyses also demonstrated that CTC positivity at both cutoff values strongly tended to have an unpleasant prognosis, the optimal cutoff value of defining CTC positivity in GC patients is still not confirmed. The studies by Li et al and Matsusaka et al showed that the cutoff value of CTCs \geq 3/7.5 mL or $\geq 4/7.5$ mL was also associated with poor prognosis in patients with GC.^{24,29} Thus, further well-designed, large-scale multicenter studies are needed to confirm the optimal sampling time and cutoff value of CTC detection. Furthermore, the genomic and functional heterogeneity of CTCs was



Figure 4 Process of exploring the potential sources of heterogeneity on PFS.

Notes: (A) Galbraith plot for PFS. (B) Forest plot for PFS after Uenosono et al²⁸ (1) (2013) is omitted. (C) Change of heterogeneity before and after Uenosono et al²⁸ (1) (2013) is omitted. Weights are from random-effects analysis. P^d value for heterogeneity. Abbreviations: HR, hazard ratio; PFS, progression-free survival; SE, standard error.



Figure 5 Funnel plots of included studies reporting OS and PFS.

Notes: (A) Begg's funnel plot on OS. (B) Egger's funnel plot on OS. (C) Begg's funnel plot on PFS. (D) Egger's funnel plot on PFS.

Abbreviations: HR, hazard ratio; OS, overall survival; PFS, progression-free survival; SE, standard error; SND, standard normal deviate.

also a source of heterogeneity. Different design types of the cohort studies also generated non-negligible heterogeneity.

Limitations

Several limitations remained in our meta-analysis. First, our meta-analysis used the data extracted from the included studies and the original data from the individual patients was hard to get. Moreover, several studies did not provide HRs and we estimated them from the reported data. Second, there was considerable heterogeneity in our study. We addressed the issue of heterogeneity by a rigorous method that used the relatively conservative random-effect model. As is well known, the prognosis of GC patients with and without surgery is different. However, the small number of studies (only the training set of Uenosono et al)²⁸ included in our meta-analysis may have been insufficient to conduct subgroup analysis to explore the different prognostic significance of CTC detection in surgical and nonsurgical groups. Third, the effects of racial disparity of GC patients from different regions on the outcome

need to be considered. In our study, six of seven included studies were reported from Asian countries, which made our findings have no broad representation. Fourth, the total number of patients was relatively small in this meta-analysis. Therefore, although our pooled analysis showed CTC positivity detected with the CellSearch System was an indicator of poor prognosis in GC patients, however, the results should be interpreted with caution and it requires larger samples, multiple regions, and more accurate study data for further verification.

Conclusion

Altogether, our meta-analysis demonstrated that CTCs detected with the CellSearch System from PB had significant prognostic value and might predict poor response to chemotherapy for patients with GC, which indicated CTC detection might have great clinical potential for guiding individualized therapy, monitoring therapeutic effect, and predicting prognosis of GC patients. Further high-quality, well-designed multicenter studies are required to confirm

the more accurate clinical value of CTCs detected with the CellSearch System in GC patients.

Author contributions

BX: concept and study design. CY, KZ, and ZY: publication retrieval and data extraction. CY, KZ, and TG: statistical analyses. CY: manuscript writing. All authors approved the submission, contributed toward data analysis, drafting and critically revising the paper and agree to be accountable for all aspects of the work.

Disclosure

The authors report no conflicts of interest in this work.

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