

Use of cytokine-induced killer cell therapy in patients with colorectal cancer: a systematic review and meta-analysis

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ABSTRACT

Background The number of clinical studies evaluating the benefit of cytokine-induced killer cell (CIK) therapy, an adoptive immunotherapy, for colorectal cancer (CRC) is increasing. In many of these trials, CIK therapy was coadministered with conventional cancer therapy. The aim of this review is to systematically assess the available literature, in which the majority were only in Chinese, on CIK therapy for the management of CRC using meta-analysis and to identify parameters associated with successful CIK therapy implementation.

Methods Prospective and retrospective clinical studies which compared CIK therapy to non-CIK therapy in patients with CRC were searched for electronically on MEDLINE, Embase, China National Knowledge Infrastructure, and Wanfang Data databases. The clinical endpoints of overall survival (OS), progression-free survival (PFS), OS and PFS rates, overall response rate (ORR), and toxicity were meta-analyzed using HR and relative ratio (RR), and subgroup analyses were performed using chi-square (χ^2) test and I-squared (I^2) statistics for study design, disease stage, cotherapy type, and timing of administration.

Results In total, 70 studies involving 6743 patients were analyzed. CIK therapy was favored over non-CIK therapy for OS (HR=0.59, 95% CI: 0.53 to 0.65), PFS (HR=0.55, 95% CI: 0.47 to 0.63), and ORR (RR=0.65, 95% CI: 0.57 to 0.74) without increasing toxicity (HR=0.59, 95% CI: 0.16 to 2.25). Subgroup analyses on OS and PFS by study design (randomized vs non-randomized study design), disease stage (Stage I–III vs Stage IV), cotreatment with dendritic cells (DCs) (CIK vs DC-CIK therapy), or timing of therapy administration (concurrent vs sequential with coadministered anticancer therapy) also showed that the clinical benefit of CIK therapy was robust in any subgroup analysis. Furthermore, cotreatment with DCs did not improve clinical outcomes over CIK therapy alone.

Conclusion Compared with standard therapy, patients who received additional CIK cell therapy had favorable outcomes without increased toxicity, warranting further investigation into CIK therapy for the treatment of CRC.

INTRODUCTION

Colorectal cancer (CRC) is the second leading cause of cancer-related death worldwide.¹

WHAT IS ALREADY KNOWN ON THIS TOPIC

⇒ Cytokine-induced killer cell (CIK) therapy is an adoptive immunotherapy used to treat both solid and hematological cancers for over 20 years. It is predominantly used in China, with multiple studies reporting benefits in patients with colorectal cancer (CRC). Despite this, CIK therapy treatment regimens are not widely used, possibly due in part to the majority of the literature about CIK therapy in CRC being reported in Chinese. Further, CIK therapy is commonly combined with other therapies but it is currently not known if there is a specific combination or treatment regimen that is optimal for CRC.

WHAT THIS STUDY ADDS

⇒ We report the most comprehensive systematic review to date of CIK therapy for patients with CRC, combining both Chinese and English language reports. Patients with CRC who received additional CIK therapy had better survival outcomes than with standard therapy alone. We also showed that the addition of dendritic cells to CIK therapy, common for CRC treatment, did not provide any clinical benefit over CIK therapy alone and that CIK therapy is effective whether given concurrently or sequentially to standard treatment regimens.

HOW THIS STUDY MIGHT AFFECT RESEARCH, PRACTICE OR POLICY

⇒ Our systematic review of Chinese and English publications shows that patients with CRC benefit from the addition of CIK therapy to standard treatment protocols and warrants further international studies.

Patients with locally advanced CRC, including regional lymph node metastases, have a 5-year survival of 75%, which reduces to 15% if there are distant metastases.² Survival outcomes for locally advanced and metastatic CRC have steadily improved due to advancements in surgical techniques, perioperative care, and

therapeutic options. However, tumor recurrence and therapy resistance remain a challenge, creating the need for new treatment options.³

During the last decade, immunotherapy has revolutionized cancer treatment, with clinical efficacy established for multiple solid and hematological cancers.⁴ Immune checkpoint inhibitors have provided significant clinical benefit, particularly in solid cancers with a high tumor mutation burden.⁵ In advanced CRC cases, they have become the standard of care for high microsatellite instability/deficient mismatch repair tumors.^{6,7} Adoptive immunotherapy involves the administration of immune cells expanded and modified in ex vivo culture. Most treatments have focused on chimeric antigen receptor T (CAR-T) cell therapy. However, other technologies including dendritic cell (DC) therapy, natural killer (NK) cell therapy, and cytokine-induced killer cell (CIK) therapy are being studied. Unlike immune checkpoint inhibitors, none of the adoptive immunotherapy products are Food and Drug Administration approved for CRC treatment.⁸

CIK therapy is an autologous, adoptive immunotherapy generated by expanding a heterogeneous population of immune effector cells from peripheral blood mononuclear cells (PBMCs).⁹ The cell therapy product contains conventional T cells (CD3+CD56-), natural killer (NK)-like T cells (CD3+CD56+), and NK cells (CD3-CD56+).¹⁰ NK-like T cells are considered the main effector cells in CIK therapy, being able to recognize tumor cells in a major histocompatibility complex class I unrestricted manner.^{11,12} Hence, guidelines for CIK therapy patient transfusion require that the cell therapy product contain at least 40% of NK-like T cells.¹³ While CIK therapy is normally combined with conventional chemotherapy, multiple trials which combine CIK therapy with other immunotherapies are being investigated. One of the more popular combinations is combining CIK therapy with autologous DC therapy (DC-CIK therapy) with reports suggesting an improvement in antitumor activity.¹⁴ China has been a leader in CIK therapy trials for multiple solid tumors, and CIK therapy is commonly provided for CRC treatment in some Chinese hospitals.^{15,16}

To date, there is a plethora of publications of varying study quality examining the clinical benefit of CIK therapy for CRC. The latest systematic review investigating the clinical efficacy of CIK therapy with chemotherapy in patients with CRC was published in 2017.¹⁶ Since then more studies have been published that support its clinical benefit,¹⁷⁻²⁰ warranting an updated systematic review to consolidate the evidence for CIK therapy in CRC management. Many of the reports originate in China and are written in Chinese. The objective of this work, therefore, is to systematically assess by meta-analysis the available literature on CIK therapy for the management of CRC, written in either English or Chinese. It includes both prospective and retrospective studies and also analyzed the benefit of parameters commonly modified in trials,

such as the addition of DCs (DC-CIK therapy) or chemotherapy regimens.

METHODS

This systematic review and meta-analysis was performed according to the Preferred Reporting Items for Systematic Reviews and Meta-analysis statement.²¹

Study selection and search

Studies which compared efficacy of CIK therapy, with or without another anticancer treatment, with no treatment or non-CIK anticancer treatment in adult patients with CRC diagnosis were identified on MEDLINE, Embase, China National Knowledge Infrastructure (CNKI), and Wanfang Data databases. CNKI and Wanfang Data were included as there were multiple studies published in Chinese alone, which were not registered with Embase or MEDLINE. The search strategy for Embase and MEDLINE is described in online supplemental tables S1 and S2, respectively. For CNKI and Wanfang Data, the following search keywords were used: “cytokine-induced killer cells,” “CIK,” “rectal cancer,” “colorectal cancer,” “colon cancer,” and “clinical trials.” No limits were placed on the language in which studies were published and the final search was performed in July 2022. Both prospective and retrospective studies with a parallel-arm design were considered, and the CIK therapy arm included patients who received CIK or DC-CIK (CIK/DC-CIK) therapy. Studies that did not report efficacy endpoints were excluded from this systematic review.

Data extraction and quality assessment

Data collection was performed independently by two authors, and discrepancies were resolved by discussion. For studies reported in Chinese, authors who are native to the Chinese language performed the data extraction and translated them into English for collation. The following information was extracted: (1) study characteristics: study design, study site, and recruitment period; (2) patient and disease characteristics: number of patients, age, gender, primary tumor location, and tumor stage; (3) study intervention: type of CIK therapy and non-CIK anticancer therapy received; (4) clinical efficacy endpoints: overall survival (OS); progression-free survival (PFS); 1-year, 3-year, and 5-year OS rates; 1-year, 3-year, and 5-year PFS rates; and overall response rate (ORR); and (5) toxicity.

For studies where patients received curative-intent treatment, disease-free survival (DFS) and DFS rates were extracted as PFS and PFS rates. Risk of bias was assessed for the following domains and graded as high, low, or unclear: (1) random sequence generation, (2) allocation concealment, (3) blinding of participants and personnel, (4) blinding of outcome assessment, (5) imbalance in baseline characteristics, (6) incomplete outcome data, and (7) uniformity of non-CIK/DC-CIK

anticancer treatment administered between intervention and control arms.

Data synthesis and analysis

Review Manager 5.4.1²² was used for pooling data at the study level and statistical analysis. For the multi-intervention-arm study, the control arm was split equally into each intervention arm, so that each pairwise comparison can be entered separately. Pooled estimates of effect were expressed as a HR calculated using an inverse variance model for OS and PFS, and risk ratios (RRs) were calculated using the Mantel-Haenszel model for survival rates and ORRs. When individual studies did not describe OS and/or PFS HRs and associated 95% CIs, they were estimated from the published Kaplan-Meier curves using a previously described method.^{23 24} The HR and 95% CI were estimated under the assumption of Gaussian distribution for the study that reported median PFS with a p-value.²⁵

As heterogeneity due to clinical diversity was expected to be high, a random-effects model was used for all the quantitative analyses performed in this review. Heterogeneity across studies was further assessed by visual inspection and statistical using chi-square (χ^2) test and I-squared (I^2) statistics for each analysis. A p-value threshold of 0.10 was employed to determine statistical significance for χ^2 test, and I^2 of 30% or less was considered to be a low degree of heterogeneity, 30% to 60% to be a moderate degree, and 60% or more to be a high degree.

Subgroup analyses were carried out on OS and PFS endpoints to investigate possible sources of heterogeneity. The following subgroup analyses were performed in this review: (1) quality of study design: randomized studies versus non-randomized studies, (2) cancer staging: Stage I–III after resection of primary versus Stage IV (unresectable, metastatic, or recurrent) CRC, (3) CIK therapy type: CIK therapy versus DC-CIK therapy, and (4) CIK therapy administration timing in relation to other anticancer therapy: concurrent versus sequential. The subgroup interactions were tested by using the formal statistical test, χ^2 test, with significance set at 10%.

RESULTS

Search results

Through our electronic search, 333 records were identified: 129 from Embase, 38 from MEDLINE, 60 from CNKI, and 106 from Wanfang Data. After removing duplicate publications and studies in which titles and/or abstracts indicated were ineligible, 106 records were assessed in detail. An additional 36 records were excluded for only a single study arm, lack of information on clinical efficacy endpoints of interest, overlapping patient cohorts with another publication, being unable to extract data specific to patients with CRC, inability to locate original abstracts or full-text articles, patients in all study arms receiving CIK therapy, and patients in the control arm being healthy subjects. Thus, 70 studies containing 16

English^{17–20 26–37} and 44 Chinese^{26 38–90} language articles were selected for study synthesis (figure 1).

Study and patient characteristics

Standardized study cohorts are summarized in table 1. Two studies^{18 90} were abstracts with the rest being full-text articles. All studies were single-center studies performed in mainland China. Fifty-four studies^{25–27 30 32 34 35 38–40 42–51 53 55–61 63–73 75 76 78–81 83–89 91} were prospective and 15 studies^{17–20 29 33 36 37 41 52 54 62 74 77 82} were retrospective in nature. Of the prospective studies, 38^{18 19 27 28 31 35 36 38 39 43–46 48 49 51 52 56 58 60–62 64–68 71–74 79–82 84 86 88 90} were randomized controlled studies.

Overall, 6743 patients with CRC, 3203 in CIK therapy (intervention) arm, and 3540 in non-CIK therapy (control) arm were available for analysis. The median age ranged from 43.2 to 80.0 years old with the youngest being 18 and the oldest being 92 years old. For studies which provided the patient's gender, 3592 out of 6017 patients (59.7%) were males. Primary tumor location was reported in 30 studies, with 1657 colon and 1744 rectum cancer patients. Patients with CRC diagnosed with all cancer stages were considered for analysis. Three studies evaluated purely patients with Stage III CRC,^{50 52 63} while 29 studies evaluated patients with Stage IV CRC.^{20 26 28 30 35 37 40 44 46 48 51 57 58 60 66–68 73–76 78 82 83 85 87–89}

The remaining studies considered patients with multiple stages. Among patients with known cancer stages, 3109 (66.6%) of them had Stage IV disease, comprising the largest group followed by 1148 patients (24.5%) with Stage III disease, 375 patients with Stage II disease, and 46 patients with Stage I disease. Cancer staging for the remaining 1672 patients was either unknown or reported in ranges.

Interventions

In 25 studies,^{17 18 20 26 28 29 32–36 38 39 44 45 47–49 54 56 63 69 73 75 85} patients in the intervention arm received CIK therapy, while in 45 studies^{19 25 27 30 31 37 40–43 46 50–53 55 57–62 64–68 70–72 74 76–84 86–89 91} DC-CIK therapy was administered. Chemotherapy was the most common cotreatment with CIK or DC-CIK therapy, being used in 66 studies.^{17–20 25–27 29–53 55–78 80–88 91}

The most commonly used chemotherapy regimens were FOLFOX and XELOX, being administered in 43^{17 18 26 30–32 34–38 40 42 44 47 49 50 54 56–65 68–70 72–75 77 78 81–84 87 90} and 24^{17 27 28 31 32 34 35 37 39–41 46 57 66 69 71 85 86 88} studies, respectively. Other less commonly used regimens included 5-fluorouracil monotherapy in six studies,^{30 33 36 39 53 77} capecitabine monotherapy in seven studies,^{17 33 36 53 56 75 82} FOLFIRI in eight studies,^{19 20 41 46 74 76 81 86} and FOLFOXIRI in two studies.^{74 86} In total, 2847 patients in the intervention arm and 3033 patients in the control arm were confirmed to have received chemotherapy as a part of the study intervention. In 10 studies, local therapy was administered together with CIK/DC-CIK therapy: radiofrequency ablation in three studies,^{28 74 86} radiotherapy in six studies,^{19 47 50 54 56 77} transarterial chemoembolization in

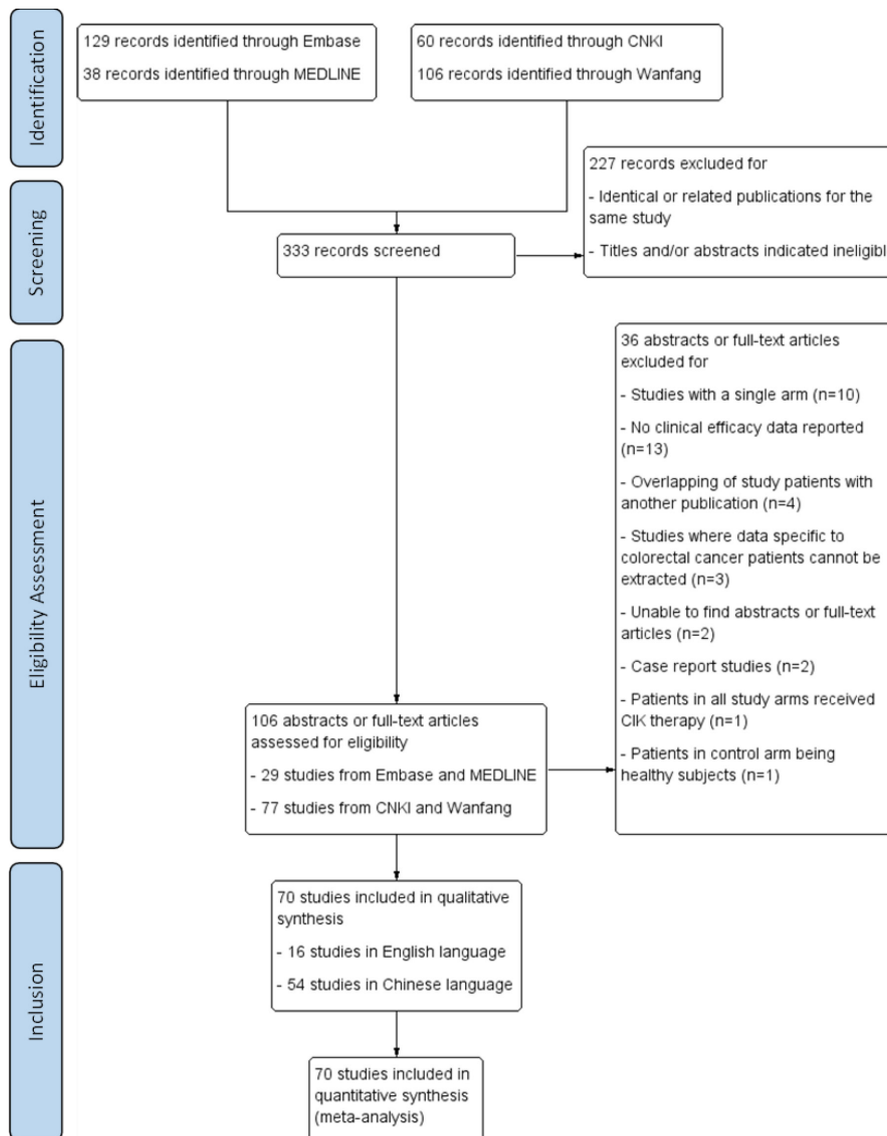


Figure 1 Flow chart of study selection.

one study,⁸⁸ and microwave hyperthermia in one study.⁵⁴ In two studies, some or all patients in the intervention arm received CIK/DC-CIK therapy alone.^{41 79}

Risk of bias assessment

Risk of bias assessment is shown in online supplemental figure 1. Among the 38 studies reported to be prospective randomized controlled studies, only nine studies^{27 32 35 43 59 66 70 78 89} described the method of randomization and no study discussed allocation concealment. None of the included studies provided clarity on the blinding of patients, study personnel, or investigators. However, it was considered unlikely that a lack of blinding would affect the clinical efficacy endpoints evaluated in the review, namely OS, PFS, OS rate and PFS rate, and ORR. All the studies were thus assessed to be at low risk of performance and detection bias secondary to insufficient blinding. Demographic and clinical characteristics of patients were generally well-balanced across the studies. Four studies^{18 27 49 58} were at unclear

risk of selection bias due to a lack of patient characteristics information across treatment arms. Imbalance in age, cancer stage, and history of primary cancer resection were noted for three studies,^{17 36 37} and they were similarly assessed to be at unclear risk of selection bias. An unclear risk of performance bias due to uncertainty around uniformity of non-CIK/DC-CIK treatment across the intervention and control arms was identified in 21 studies^{18 20 27 30 33 37 41 42 46 50 51 53 56 60 74 77–79 81 82 86} with all the studies except one failing to adequately describe study interventions or the proportion of patients receiving various interventions. In the remaining one study,⁷⁹ patients in the intervention arm received DC-CIK therapy alone, while those in the control arm received the best supportive care. The risk of attrition bias was rated unclear for 18 studies^{17–20 26 28 30 32 34–38 40 66 77 81 83 88} which did not reveal the number of patients lost in follow-up and for one study³⁶ in which 18.8% of patients withdrew from the study prematurely.

Table 1 Summary of included studies

Study ID	Study type	RCT	Patient (n)	Age, years (range)	Male (%)	Colon primary (%)	Stage	Intervention arm(s) (n)	Control arm (n)	Outcomes
Bian <i>et al</i> ⁸¹	P	Yes	84	39–78	54.8	73.8	I–III	DC-CIK+ChT (42)	ChT (42)	OS, PFS, ORR, SE
Cai <i>et al</i> ³⁸	P	Yes	80	–	62.5	–	II–III	CIK+ChT (40)	ChT (40)	OS, PFS, SE
Cai and Li ³⁹	P	No	72	–	69.4	0.0	I–III	CIK+ChT (40)	ChT (32)	OS, SE
Cai ⁴⁰	P	No	90	30–70	70.0	–	IV	DC-CIK+ChT (45)	ChT (45)	OS, PFS, SE
Chao <i>et al</i> ⁴¹	R	No	66	–	68.2	–	III–IV	DC-CIK±ChT (33)	No treatment or ChT (33)	OS, SE
Chen <i>et al</i> ⁴²	P	Yes	60	30–65	–	100.0	III–IV	DC-CIK+ChT (30)	ChT (30)	OS, PFS, SE
Chen ⁴³	P	Yes	90	32–76	65.0	–	I–III	DC-CIK+ChT (45)	ChT (45)	OS, PFS
Chu <i>et al</i> ⁴⁴	P	Yes	89	–	51.7	–	IV	CIK (30) and CIK+ChT (29)	ChT (30)	OS, PFS, SE
Deng <i>et al</i> ⁴⁵	P	Yes	60	–	–	–	III–IV	CIK+ChT (30)	ChT (30)	OS, PFS, SE
Dong ⁴⁶	P	No	40	18–75	52.5	37.5	IV	DC-CIK+ChT (20)	ChT (20)	OS, PFS, SE
Du <i>et al</i> ²⁶	P	Yes	60	–	53.3	68.3	IV	CIK+ChT (30)	ChT (30)	OS, PFS, SE
Fan <i>et al</i> ⁴⁷	P	Yes	81	32–83	–	0.0	I–III	CIK+ChT+RT (41)	ChT+RT (40)	OS, PFS, ORR, SE
Fang ⁴⁸	P	Yes	52	–	59.6	–	IV	CIK+ChT (26)	ChT (26)	ORR, SE
Feng <i>et al</i> ⁴⁹	P	No	40	51–55	70.0	–	I–III	CIK+ChT (20)	ChT (20)	OS, PFS
Feng and Wang ⁵⁰	P	Yes	50	41–79	66.0	–	III	DC-CIK+ChRT (25)	ChRT (25)	PFS, ORR, SE
Gao <i>et al</i> ²⁷	P	No	26	–	–	–	<IV	DC-CIK+ChT (13)	No treatment (13)	OS, PFS, SE
Guo and Ma ⁵¹	P	Yes	68	41–83	57.4	27.9	IV	DC-CIK+ChT (34)	ChT (34)	ORR
He and Zhang ⁵²	R	–	100	40–80	60.0	–	III	DC-CIK+ChT (50)	ChT (50)	OS, ORR
Jiang ⁵³	P	No	98	22–78	64.3	–	III–IV	DC-CIK+ChT (50)	ChT (48)	ORR
Leng <i>et al</i> ⁵⁴	R	–	90	29–66	57.8	0.0	III–IV	CIK+IMRT+microwave hyperthermia (45)	IMRT+microwave hyperthermia (45)	ORR, SE
Li <i>et al</i> ⁵⁵	P	Yes	40	–	70.0	–	II–III	DC-CIK+ChT (20)	ChT (20)	OS, ORR
Li <i>et al</i> ⁵⁶	P	No	130	–	56.2	43.0	II–III	CIK+ChT+RT (65)	ChT+RT (65)	ORR, SE
Li <i>et al</i> ²⁸	U	–	60	–	63.3	63.3	IV	CIK+RFA (30)	RFA (30)	ORR, SE
Li <i>et al</i> ²⁹	R	–	137	–	60.6	–	II–IV	CIK+ChT (66)	ChT (71)	ORR, SE
Lin <i>et al</i> ³⁰	P	Yes	255	–	54.9	56.1	IV	DC-CIK+ChT (134)	ChT (121)	OS
Liu ⁵⁷	P	Yes	56	32–72	55.4	–	IV	DC-CIK+ChT (28)	ChT (28)	ORR, SE
Liu <i>et al</i> ⁵⁸	P	No	18	28–73	55.6	–	IV	DC-CIK+ChT (9)	ChT (9)	ORR, SE

Continued



Table 1 Continued

Study ID	Study type	RCT	Patient (n)	Age, years (range)	Male (%)	Colon primary (%)	Stage	Intervention arm(s) (n)	Control arm (n)	Outcomes
Liu <i>et al</i> ⁶⁰	P	Yes	58	35–80	63.8	–	IV	DC-CIK+ChT (29)	ChT (29)	ORR
Liu <i>et al</i> ⁶¹	P	Yes	80	22–82	61.3	–	I–IV	DC-CIK+ChT (40)	ChT (40)	ORR
Liu <i>et al</i> ⁶²	R	–	90	–	64.4	0.0	I–IV	DC-CIK+ChT (45)	ChT (45)	ORR
Lin <i>et al</i> ⁸¹	P	Yes	70	–	68.6	40.0	II–IV	DC-CIK+ChT (35)	ChT (35)	OS, ORR, SE
Liu ⁵⁹	P	Yes	68	30–79	59.7	69.1	III–IV	DC-CIK+ChT (34)	ChT (34)	OS
Lv <i>et al</i> ⁶³	P	Yes	85	–	51.8	–	III	CIK+ChT (43)	ChT (42)	ORR, SE
Ma ⁶⁴	P	Yes	50	49–77	60.0	–	II–III	DC-CIK+ChT (25)	ChT (25)	ORR, SE
Niu ⁶⁵	P	Yes	50	34–62	64.0	–	III–IV	DC-CIK+ChT (25)	ChT (25)	ORR
Pan <i>et al</i> ²⁰	R	–	252	–	61.5	63.5	IV	CIK+ChT (126)	ChT (126)	OS, PFS
Pan <i>et al</i> ¹⁷	R	–	122	–	63.9	64.8	II–IV	CIK+ChT (60)	ChT (62)	ORR
Peng <i>et al</i> ⁸²	P	Yes	46	–	63.0	–	II–III	CIK+ChT (23)	ChT (23)	ORR
Pu <i>et al</i> ⁶⁶	P	Yes	98	–	38.8	–	IV	DC-CIK+ChT (49)	ChT (49)	SE
Rui <i>et al</i> ⁶⁷	P	Yes	90	18–60	57.8	53.3	IV	DC-CIK+ChT (45)	ChT (45)	OS, ORR
Sun ⁶⁸	P	No	60	41–68	51.7	–	IV	DC-CIK+ChT (30)	ChT (30)	OS, ORR, SE
Wang <i>et al</i> ⁶⁹	P	No	110	–	60.0	63.6	I–III	CIK+ChT (55)	ChT (55)	OS, ORR
Wang <i>et al</i> ⁷⁰	P	Yes	104	32–69	58.7	–	III–IV	DC-CIK+ChT (52)	ChT (52)	ORR
Wang <i>et al</i> ⁷¹	P	Yes	68	–	64.1	–	III–IV	DC-CIK+ChT (34)	ChT (34)	OS, ORR, SE
Wang <i>et al</i> ¹⁸	R	–	377	–	–	–	–	CIK+ChT (97)	ChT (280)	ORR
Weng <i>et al</i> ⁷²	P	Yes	235	–	55.3	56.6	III–IV	DC-CIK+ChT (124)	ChT (111)	ORR
Weng <i>et al</i> ⁷³	P	Yes	96	25–76	59.6	–	IV	CIK+ChT (48)	ChT (48)	ORR
Wu ⁷⁴	R	–	132	–	66.7	51.5	IV	DC-CIK+ChT+RFA (62)	ChT+RFA (70)	OS
Xie <i>et al</i> ¹⁹	R	–	142	–	55.6	–	III–IV	DC-CIK+ChT±RT (71)	ChT±RT (71)	OS, ORR
Xu <i>et al</i> ⁸³	R	–	116	–	46.6	54.3	II–IV	CIK+ChT (32)*	ChT (82)*	OS, ORR
Yan <i>et al</i> ⁷⁵	P	No	114	–	56.1	–	IV	CIK+ChT (72)	ChT (42)	OS, PFS, ORR
Yin ⁷⁶	P	No	80	–	61.3	60.0	IV	DC-CIK+ChT (40)	ChT (40)	OS, PFS, ORR
Ying <i>et al</i> ⁷⁷	R	–	102	20–86	54.9	–	I–III	DC-CIK+ChT+RT (51)	ChT+RT (51)	PFS, ORR
Yuan <i>et al</i> ⁷⁸	P	Yes	42	45–78	73.8	–	IV	DC-CIK+ChT (21)	ChT (21)	OS
Yue ⁷⁹	P	Yes	110	–	47.3	53.6	III–IV	DC-CIK (55)	BSC (55)	OS, PFS, ORR
Zang <i>et al</i> ⁸⁰	P	Yes	90	–	66.7	–	III–IV	DC-CIK+ChT (45)	ChT (45)	OS
Zhang <i>et al</i> ⁸¹	P	Yes	63	24–82	61.9	–	III–IV	DC-CIK+ChT (32)	ChT (31)	OS
Zhang <i>et al</i> ⁸⁴	P	Yes	60	–	63.3	56.7	I–IV	CIK+ChT (30)	ChT (30)	OS, SE

Continued

Table 1 Continued

Study ID	Study type	RCT	Patient (n)	Age, years (range)	Male (%)	Colon primary (%)	Stage	Intervention arm(s) (n)	Control arm (n)	Outcomes
Zhang <i>et al</i> ⁸²	R	-	84	-	54.8	-	IV	DC-CIK+ChT (42)	ChT (42)	OS, PFS, ORR, SE
Zhang <i>et al</i> ²⁵	P	No	112	-	52.2	52.0	III-IV	DC-CIK+ChT (65)	ChT (47)	ORR
Zhang ⁸³	P	Yes	118	46-78	49.2	-	IV	DC-CIK+ChT (59)	ChT (59)	OS, PFS
Zhang ⁸⁴	P	No	90	43-73	67.8	51.1	III-IV	DC-CIK+ChT (45)	ChT (45)	SE
Zhao ⁸⁷	P	Yes	30	32-67	70.0	-	IV	DC-CIK+ChT (15)	ChT (15)	ORR
Zhao ³⁵	P	Yes	122	-	67.2	58.7	IV	CIK+ChT (61)	ChT (61)	ORR, SE
Zhao ⁸⁵	P	Yes	90	40-59	63.3	-	IV	CIK+ChT (45)	ChT (45)	ORR, SE
Zhao <i>et al</i> ⁸⁶	P	No	148	-	62.8	-	I-III	DC-CIK+ChT+RFA (73)	ChT+RFA (75)	OS
Zhou ⁸⁹	P	Yes	60	45-80	68.3	78.3	IV	DC-CIK+ChT (30)	ChT (30)	OS, ORR
Zhou <i>et al</i> ⁸⁸	P	No	90	-	-	-	IV	DC-CIK+TACE (45)	TACE (45)	ORR, SE
Zhu <i>et al</i> ³⁶	R	-	96	-	57.3	62.1	I-IV	CIK+ChT (21)	ChT (75)	OS, PFS, ORR, SE
Zhu <i>et al</i> ³⁷	R	-	351	19-92	65.2	30.8	IV	DC-CIK+standard care (100)	Standard care (251)	ORR, SE

*Sixteen and 18 patients in intervention arm and 47 and 35 patients in control arm were treated in adjuvant and palliative setting, respectively. BSC, best supportive care; ChRT, concurrent chemoradiotherapy; ChT, chemotherapy; CIK, cytokine-induced killer cell; DC-CIK, CIK therapy with autologous dendritic cell therapy; IMRT, intensity-modified radiotherapy; n, number; ORR, overall response rate; OS, overall survival; PFS, progression-free survival; P, prospective; R, retrospective; RCT, randomized controlled trial; RFA, radiofrequency ablation; RT, radiotherapy; TACE, transarterial chemoembolization; U, unknown.

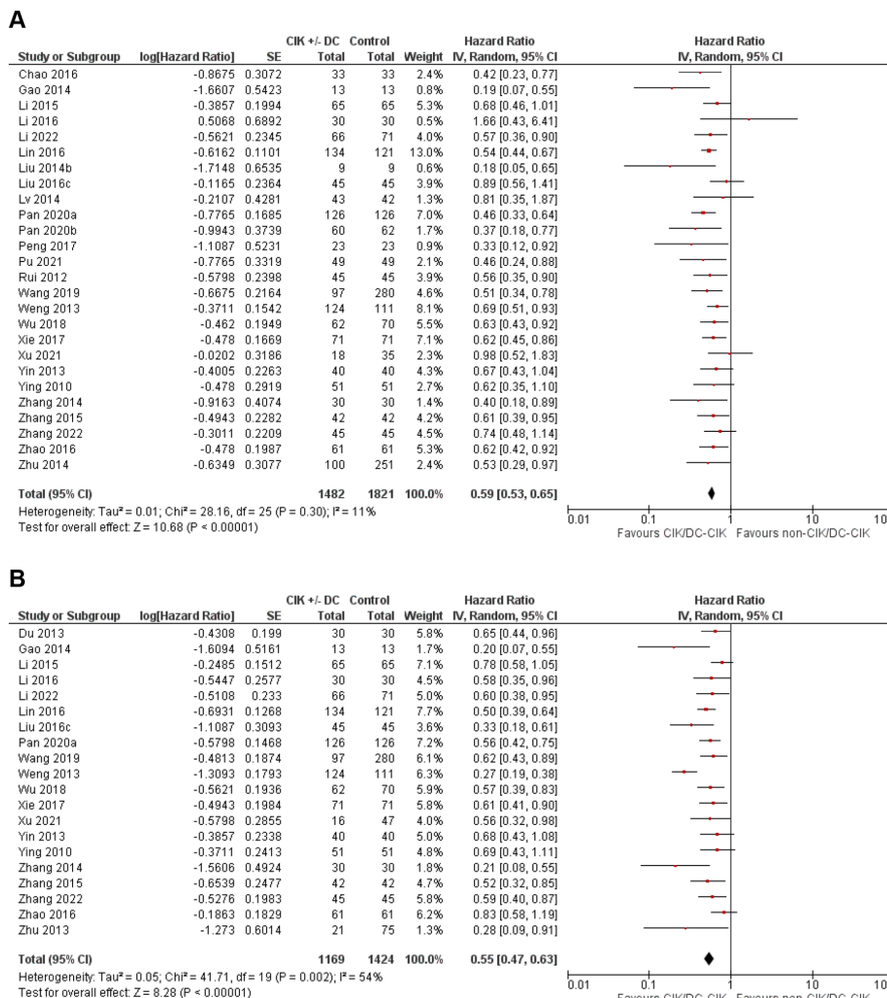


Figure 2 Comparison of CIK/DC-CIK therapy versus non-CIK/DC-CIK therapy for (A) overall survival (OS) and (B) progression-free survival (PFS). Twenty-six studies involving 3,303 patients and twenty studies involving 2,593 patients contributed data to OS and PFS analysis respectively. CIK, cytokine-induced killer cell; DC, dendritic cell.

Overall survival and progression-free survival

There were 26 studies^{17 19 20 27 29 32 33 36 38 39 41 43 47 48 53 55-57 60 63 67 75 77 79 81 86} involving 3303 patients which contributed data to the meta-analysis on OS (figure 2A). The pooled HR was 0.59 (95% CI: 0.53 to 0.65) indicating the OS benefit of CIK/DC-CIK therapy over the control arm. Heterogeneity among the studies was low (I²=11%, p=0.30). For PFS, 20 studies^{18-20 26-30 33-36 56 62 72 74 76 77 82 84} involving 2593 patients contributed the data to the meta-analysis (figure 2B). The pooled HR was 0.55 (95% CI: 0.47 to 0.63), again favoring CIK/DC-CIK therapy. Heterogeneity among the studies was moderate (I²=54%, p=0.002).

Overall survival rates

In total, 27 (2459 patients),^{17 19 20 27 29 32 33 36 38 39 41 43 47-49 53 55-57 60 63 67 75 77 79 81 86} 19 (2167 patients),^{17-20 27-29 33 35 36 38 39 41 48 49 56 67 77 86} and 10 (1401 patients)^{17-20 27 29 33 36 41 56} studies contributed data for 1-year, 3-year, and 5-year OS rate meta-analyses, respectively (online supplemental figure 2). The pooled RR for all the analyses favored CIK/DC-CIK therapy. The 1-year OS rate was 91.7% in the intervention arm and 79.4% in

the control arm with a pooled RR of 0.47 (95% CI: 0.32 to 0.67). Heterogeneity among the studies was moderate (I²=51%, p=0.002). The 3-year OS rate was 67.7% in the intervention arm and 51.8% in the control arm with a pooled RR of 0.67 (95% CI: 0.59 to 0.77). There was a moderate level of heterogeneity among the studies (I²=32%, p=0.09). The 5-year OS rate was 61.2% in the intervention arm and 45.5% in the control arm with an RR of 0.69 (95% CI: 0.54 to 0.88). Heterogeneity among the studies was high (I²=73%, p=0.0001).

Progression-free survival rates

We identified 10 (1166 patients),^{17 19 20 27 29 33 36 37 56 77} 10 (1156 patients),^{17 19 20 27-29 33 35 56 77} and 7 (872 patients) studies^{17 19 20 27 29 33 56} that contributed data for meta-analysis on 1-year, 3-year, and 5-year PFS rates, respectively (online supplemental figure 3). All the analyses indicated the superiority of CIK/DC-CIK therapy over non-CIK/DC-CIK therapy. The observed 1-year PFS rate was 86.5% in the intervention arm and 68.1% in the control with the pooled RR of 0.43 (95% CI: 0.33 to 0.55). Heterogeneity among the studies was low (I²=0%, p=0.48). The 3-year PFS rate was 47.8% in the

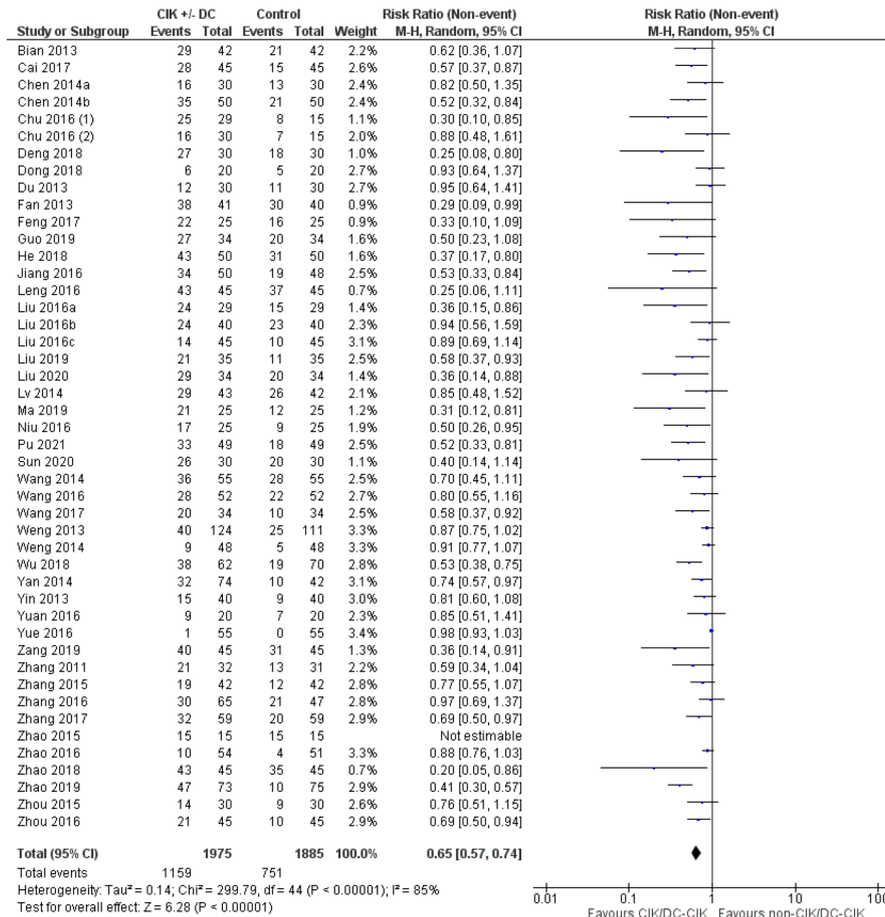


Figure 3 CIK/DC-CIK therapy versus non-CIK/DC-CIK therapy for overall response rate (ORR). Forty-five studies involving 3,860 patients contributed data to ORR analysis. *Study Chu 2016 appears twice in the figure as it contained 3 treatment arms and data were entered separately for CIK + chemotherapy versus chemotherapy (Chu 2016 (1)) and CIK versus chemotherapy (Chu 2016 (2)) by splitting the chemotherapy group into 2 subgroups, one for each CIK + chemotherapy and CIK treatment. CIK, cytokine-induced killer cell; DC, dendritic cell.

intervention arm and 30.5% in the control arm. The pooled RR was 0.76 (95% CI: 0.66 to 0.87) and heterogeneity among the studies was moderate ($I^2=53\%$, $p=0.02$). At 5 years, the PFS rate was 46.0% in the intervention arm and 25.9% in the control arm. The pooled RR was 0.71 (95% CI: 0.59 to 0.87) and heterogeneity among the studies was high ($I^2=68\%$, $p=0.005$).

Overall response rate

The ORR was 58.7% in the intervention (CIK/DC-CIK) and 39.8% in the control (non-CIK/DC-CIK) arm for 3860 patients from 45 studies^{25 26 31 35 40 42-47 50-54 59-66 68-76 78-83 85-89 91} (figure 3). The pooled RR was 0.65 (95% CI: 0.57 to 0.74), and heterogeneity among the studies was high ($I^2=85\%$, $p<0.00001$).

Toxicity

Toxicity during the study intervention was reported by 31 studies with the majority of the data being provided in a descriptive manner. Two studies^{18 85} compared the rate of any adverse events between the treatment arms, and 11 studies^{30 35 39 40 42 54 59 60 68 70 87 91} reported adverse events of interest for each arm. Many of the described side effects

were thought to be related to chemotherapy administered together with CIK/DC-CIK therapy, including bone marrow suppression, nausea, vomiting, neuropathy, diarrhea, and liver dysfunction. Meta-analysis undertaken indicated equivalent adverse event rate from CIK/DC-CIK and non-CIK/DC-CIK therapy (HR=0.59, 95% CI: 0.16 to 2.25) with the pooled adverse event rate of 53.5% and 68.3%, respectively (online supplemental figure 4). Heterogeneity was high between the studies ($I^2=80\%$, $p=0.02$). Fever was the most frequently reported adverse event associated with CIK/DC-CIK infusion, affecting 6.7% to 29.9% of patients receiving CIK/DC-CIK therapy. Fever, in general, spontaneously resolved or only required symptomatic management.

Subgroup analyses

Potential sources of heterogeneity were explored by performing subgroup analysis on OS and PFS by study design (randomized vs non-randomized study design), disease stage (Stage I-III vs Stage IV), CIK therapy type (CIK vs DC-CIK therapy), or timing of CIK/DC-CIK therapy administration (concurrent vs sequential with coadministered anticancer therapy).

Randomized studies versus non-randomized studies

Of the 25 studies which provided OS HRs, 8 studies^{30 32 34 35 63 66 67 72} involving 991 patients were prospective randomized studies and 17 studies^{17-20 27 29 33 37 41 56 58 62 74 76 77 82 84} involving 2252

patients were either prospective non-randomized or retrospective studies. An OS benefit of CIK/DC-CIK therapy was demonstrated for both randomized studies (HR=0.57; 95% CI: 0.50 to 0.66) and non-randomized studies (HR=0.59, 95% CI: 0.51 to 0.67) (figure 4A). A test for

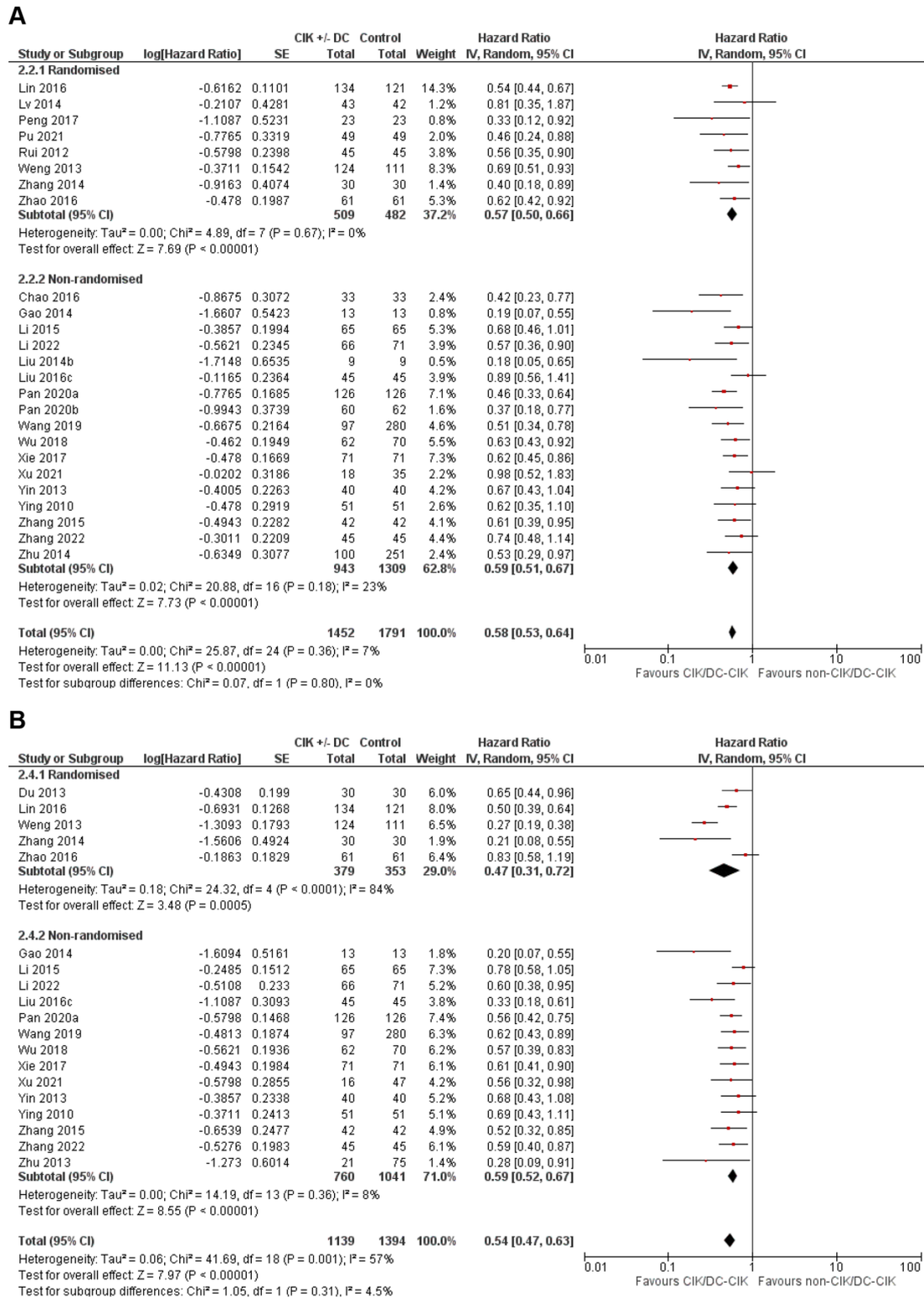


Figure 4 Subgroup analysis by study design for (A) overall survival (OS) and (B) progression-free survival (PFS). Twenty-five studies involving 3,243 patients and nineteen studies involving 2,533 patients contributed data to OS and PFS analysis respectively.

subgroup difference did not reach statistical significance ($I^2=0\%$, $p=0.80$). For PFS subgroup analysis, 732 patients from five randomized studies^{26 30 34 35 72} and 1801 patients from 14 non-randomized studies^{18–20 27 29 33 35 36 62 74 76 77 82 84} were analyzed. A benefit from CIK/DC-CIK therapy was again shown for both prospective randomized (HR=0.47, 95% CI: 0.31 to 0.72) and non-randomized studies (HR=0.59, 95% CI: 0.47 to 0.63) (figure 4B). A test for subgroup differences was not statistically significant ($I^2=4.5\%$, $p=0.31$).

Stage I–III versus Stage IV

Four studies^{32 56 63 77} involving 363 patients with Stage I–III CRC and 12 studies^{20 28 30 33 35 37 58 66 67 74 76 82} involving 1595 Stage IV patients contributed data to the subgroup analysis on OS by the disease stage. HR for Stage I–III patients was 0.64 (95% CI: 0.48 to 0.85), while that for Stage IV patients was 0.57 (95% CI: 0.50 to 0.65) and the benefit of CIK/DC-CIK therapy was observed across all stages of CRC (online supplemental figure 5A). Test for subgroup differences failed to reach statistical significance ($I^2=0\%$, $p=0.48$), although the observed 95% CI was much narrower for Stage IV patients. For the subgroup analysis on PFS, four studies^{27 33 56 77} involving 321 patients with Stage I–III disease and eight studies^{20 26 28 30 35 74 76 82} involving 1045 Stage IV patients were analyzed. A benefit from CIK/DC-CIK therapy was demonstrated for both Stage I–III (HR=0.60, 95% CI: 0.40 to 0.88) and Stage IV disease (HR=0.59, 95% CI: 0.52 to 0.67) (online supplemental figure 5B). A test for subgroup difference was not statistically significant ($I^2=0\%$, $p=0.94$).

CIK therapy versus DC-CIK therapy

Ten studies^{17 18 20 28 29 32 34 35 56 63} (1391 patients) and 16 studies^{19 27 30 33 37 41 58 62 66 67 72 74 76 77 82 84} (1912 patients) which evaluated CIK and DC-CIK therapy, respectively, were assessed in the subgroup analysis on OS by the type of CIK therapy. HR for studies examining CIK therapy was 0.57 (95% CI: 0.47 to 0.69), while that for studies examining DC-CIK therapy was 0.61 (95% CI: 0.54 to 0.69) (figure 5A). Both types of CIK therapy were found to benefit OS. A test for subgroup differences did not reach statistical significance ($I^2=0\%$, $p=0.58$). Subgroup analysis on PFS by CIK therapy type contained nine studies^{18 20 26 28 29 34–36 56} involving 1294 patients, where the intervention arm contained CIK therapy, and 11 studies^{19 27 30 33 62 72 74 76 77 82 84} involving 1299 patients, where the intervention arm contained DC-CIK therapy. PFS benefit was demonstrated for both CIK-examining (HR=0.63, 95% CI: 0.53 to 0.74) and DC-CIK-examining studies (HR=0.50, 95% CI: 0.41 to 0.61) (figure 5B). A test for subgroup differences met statistical significance ($I^2=66.5\%$, $p=0.08$) with improved HR seen for DC-CIK, although HRs for the two subgroups overlapped each other, suggesting that the advantage of DC-CIK over CIK therapy alone may not be clinically meaningful.

Concurrent CIK/DC-CIK therapy versus sequential CIK/DC-CIK therapy

Subgroup analysis was performed comparing studies where CIK/DC-CIK therapy was administered either concurrently or sequentially with the non-CIK/DC-CIK therapy. For OS analysis, 16 studies^{19 20 28 33 35 37 41 56 58 62 63 67 72 74 82 84} involving 2000 patients with concurrent administration and 8 studies^{17 27 29 30 32 34 66 77} involving 846 patients with sequential administration were considered (figure 6A). CIK/DC-CIK therapy administered in either manner improved OS; the HR was 0.63 (95% CI: 0.56 to 0.71) for concurrent administration and 0.59 (95% CI: 0.53 to 0.65) for sequential administration. A test for subgroup differences reached statistical significance ($I^2=76.3\%$, $p=0.04$) with lower HR being observed for sequential administration, although 95% CIs of the two subgroups overlapped each other. Subgroup analysis on PFS was similarly in favor of CIK/DC-CIK therapy for both concurrent (HR=0.56, 95% CI: 0.46 to 0.67) and sequential administration (HR=0.54, 95% CI: 0.46 to 0.63) (figure 6B). Twelve studies^{19 20 26 28 33 35 56 62 72 74 82 84} involving 1460 patients who had concurrent administration and five studies^{27 29 30 34 77} involving 580 patients who had sequential administration were evaluated, and a test for subgroup differences did not meet statistical significance ($I^2=0\%$, $p=0.43$).

DISCUSSION

Chemotherapy with/without biological therapy remains the standard treatment for patients with CRC with the high-risk resected disease, and the majority of those with advanced disease. This therapeutic approach is associated with limited survival benefit, unlike immunotherapy, which has demonstrated long-term survival outcome in some solid tumors owing to its mechanism of action.^{92 93} New therapeutic approaches which involve modulation of the immune system may provide new treatment options for a broader range of patients with CRC and improve their survival outcome. Autologous adoptive immunotherapy such as CIK therapy represents a highly personalized cancer treatment. While it remains a non-standard treatment option for solid cancers, there are a growing number of clinical trials examining such immunotherapy.⁹⁴

Our study demonstrated that providing CIK or DC-CIK therapy to patients with CRC improved OS, PFS, and ORR compared to standard treatment. The upper 95% CI of pooled HRs for 5-year OS rate and 3-year and 5-year PFS rates exceeded 0.85, a commonly applied cut-off to delineate no effect from an important effect, raising the possibility that the observed benefit for these endpoints may not be precise. However, for all the other endpoints, the observed HRs favoring CIK/DC-CIK therapy appeared robust. The OS and PFS benefit of CIK/DC-CIK therapy persisted when prospective randomized studies alone were examined in the subgroup analysis, with no subgroup differences being identified compared with non-randomized studies. While the number of

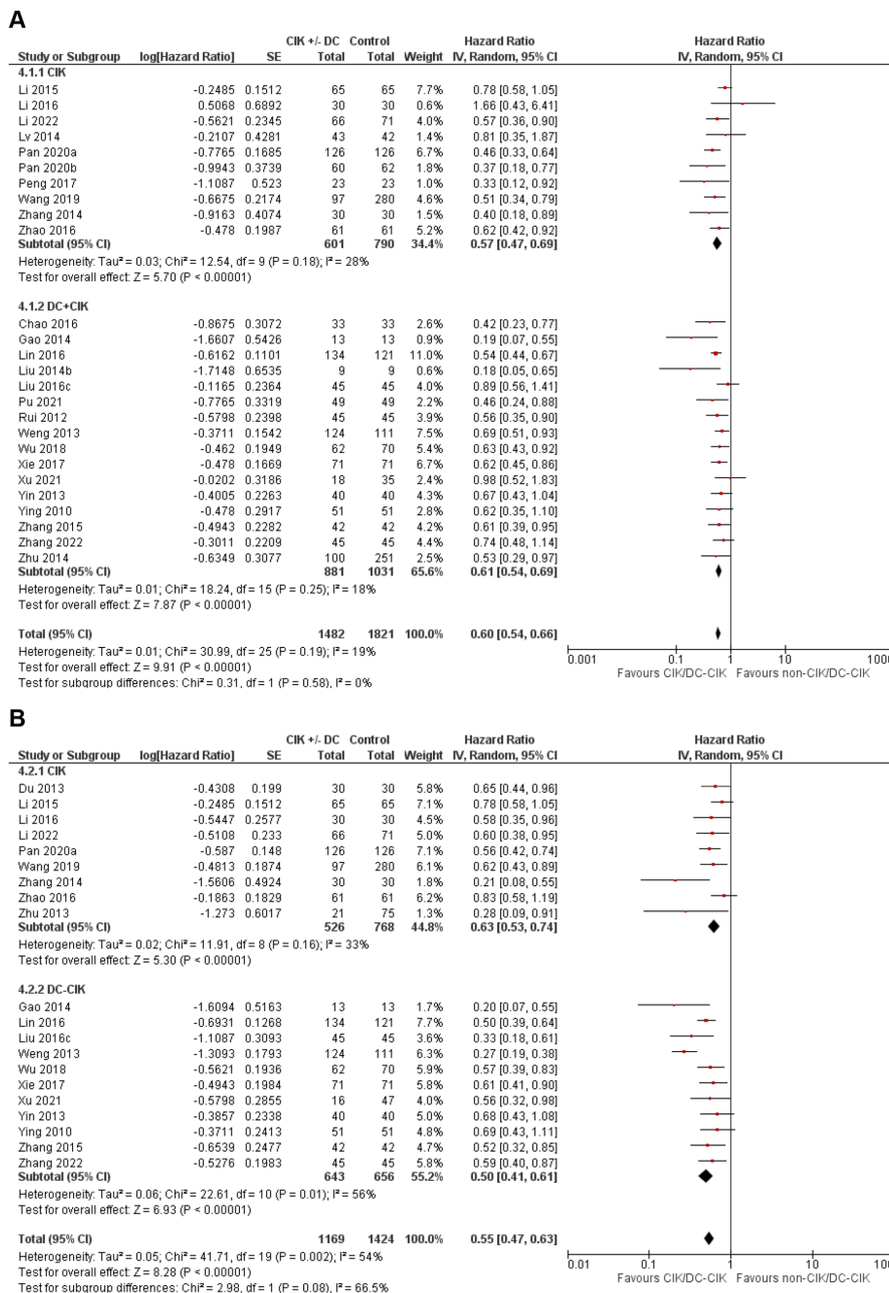


Figure 5 Subgroup analysis by CIK therapy type (with or without DC therapy) for (A) overall survival (OS) and (B) progression-free survival (PFS). Twenty-six studies involving 3,303 patients and twenty studies involving 2,593 patients contributed data to OS and PFS analysis respectively. CIK, cytokine-induced killer cell; DC, dendritic cell.

randomized studies assessed was small, HRs and associated 95% CIs reported by each study, especially for the OS endpoint, were all comparable, indicating consistency in the results and so strengthening the overall finding.

Subgroup analysis by CRC disease stage indicated a lack of differences for both OS and PFS. However, the observed 95% CIs associated with the pooled HRs were persistently narrower for Stage IV patients compared with Stage I–III patients, with the upper limits of 95% CIs for Stage I–III patients exceeding 0.85 for both endpoints. Together with the uncertainties around the best way to incorporate CIK/DC-CIK therapy into the established 3–6 months of monoadjuvant or doublet-adjuvant chemotherapy,

depending on the disease stage and accompanying other prognostic factors, our study highlights that patients with Stage IV disease may be a more suitable target to evaluate CIK/DC-CIK therapy application, at least initially. The immunosuppressive effect of cancer surgery, including T cell and NK cell dysfunction and expansion of myeloid-derived suppressor cells and regulatory T cells in the postoperative period has been described previously,⁹⁵ although how this affects the antitumor activity of CIK/DC-CIK therapy is not known.

Subgroup analysis based on combining DC therapy with CIK therapy revealed statistically significant subgroup differences in favor of DC-CIK over CIK therapy for PFS,

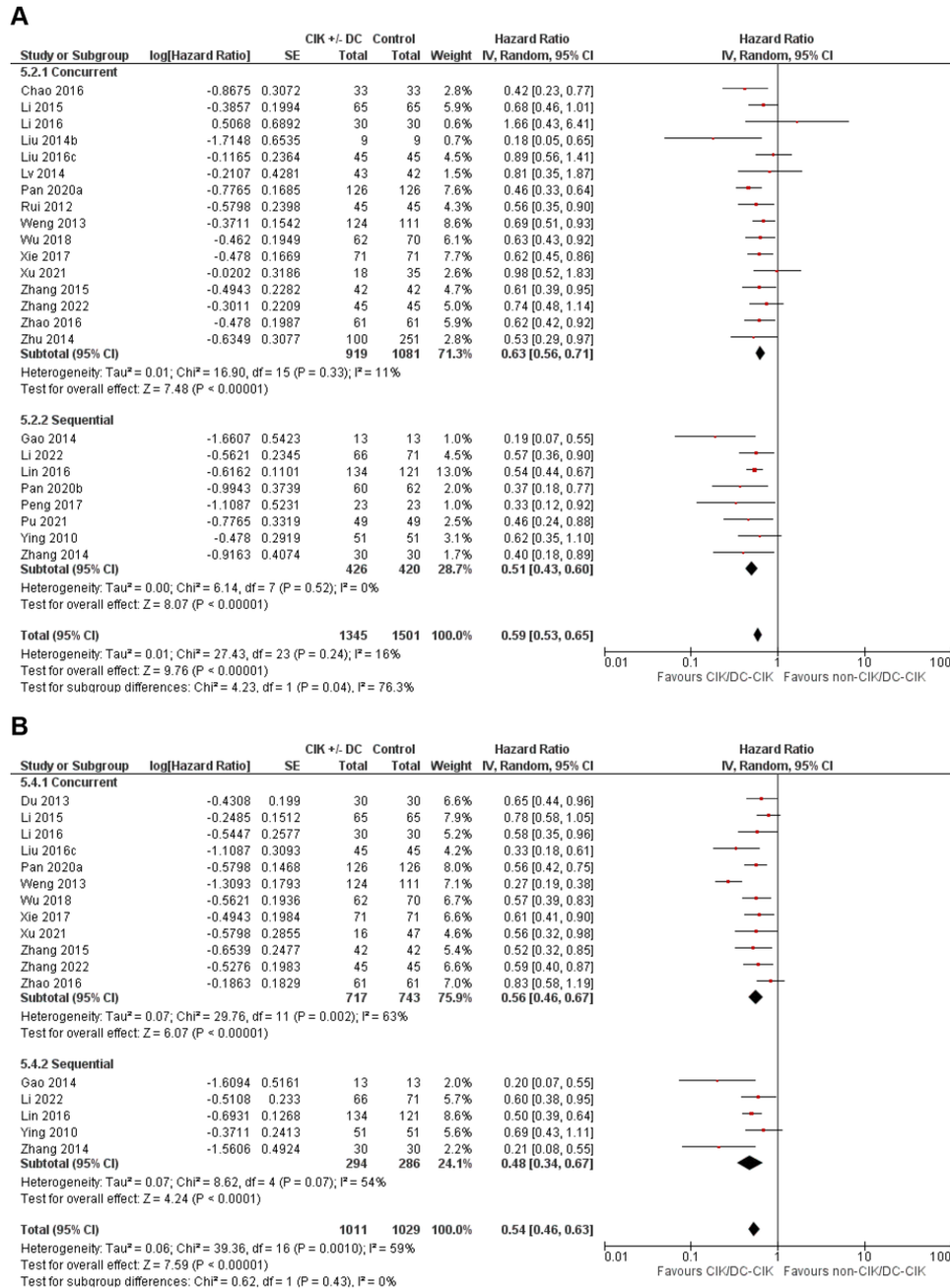


Figure 6 Subgroup analysis by CIK/DC-CIK therapy administration timing for (A) overall survival (OS) and (B) progression-free survival (PFS). Twenty-four studies involving 2,846 patients and seventeen studies involving 2,040 patients contributed data to OS and PFS analysis respectively. CIK, cytokine-induced killer cell; DC, dendritic cell.

but not for OS. DCs are major antigen-presenting cells and the essential link between the innate and adaptive immune systems.⁹⁶ Coculturing of CIK cells with DCs results in increased CIK cytolytic function, including cytotoxic activity against a tumor cell line resistant to CIK cells cultured in the absence of DCs.¹⁴ This review observed more patients who received DC-CIK therapy than CIK therapy; however, the results suggest that the addition of DC therapy to CIK therapy does not have a strong clinical benefit, as only statistical significance was observed for PFS and not for OS. This result points to the need for future clinical trials investigating the benefit of including DC therapy in CIK therapy, and whether other

combinations such as immune checkpoint inhibitors or CAR-T incorporation with CIK therapy may be of better value for patients with CRC.

Subgroup differences were similarly detected for OS for concurrent versus sequential administration of CIK/DC-CIK. Subgroup analyses for both PFS by CIK therapy type and OS by CIK therapy administration timing had similar HRs with highly overlapping 95% CIs, making it unclear whether the differences are clinically meaningful. The timing of CIK/DC-CIK delivery for patients with CRC may not be critical and could be selected based on logistical issues.

There have been two previous publications that systematically reviewed the literature for CIK/DC-CIK therapy in CRC.^{16,97} In 2010, Zhang and Schmidt-Wolf, in cooperation with Stanford University, established the International Registry on CIK Cells (IRCC) to evaluate clinical trials of CIK therapy.^{97,98} The registry identifies both prospective and retrospective clinical trials involving CIK therapy for cancer treatment from PubMed, Web of Science Core Collection, WHO International Clinical Trials Registry Platform, ClinicalTrials.gov as well as proceedings of the American Society of Clinical Oncology and European Cancer Conference Annual Scientific Meetings. In addition, the IRCC incorporates clinical trials submitted by individual researchers for inclusion.⁹⁹ In 2020, the registry recorded 106 clinical trials, of which only 6 examined CIK therapy in patients with CRC.⁹⁷ This contrasts with the 29 trials including 2610 patients with CRC reported in the published systematic review and meta-analysis in 2017 by Zhang *et al*, which purely compared the clinical benefit of CIK therapy plus chemotherapy to CIK therapy in patients with CRC with advanced disease.¹⁶ They also used two Chinese databases, CNKI and Wanfang Data, in addition to the English databases Cochrane Library, Embase, and PubMed. The majority of the studies were published in Chinese similar to our findings.

To date, China has taken the lead in research of adoptive immunotherapy including CIK therapy.^{15,100} Therefore, the inclusion of articles published in Chinese was necessary to comprehensively review the currently available literature examining the clinical efficacy of CIK therapy in CRC. Additionally, the current work included clinical trials which compared CIK therapy with non-CIK treatment not limited to chemotherapy, to increase the number of trials assessed. Consequently, the review considered 70 studies involving 6743 patients and is the largest systemic review on CIK/DC-CIK therapy in CRC. It meta-analyzed OS and PFS, the two most important clinical endpoints in assessing the efficacy of any cancer therapy. Endpoints covered by Zhang¹⁶ were limited to OS and DFS rates as well as ORR. The CRC population covered by this review is also broader having included patients at all stages.

This study has a number of limitations. The heterogeneity observed in the clinical study design requires caution when interpreting results. There are general guidelines for the production of CIK therapy. The CIK therapy product is generated from PBMCs cultured for 21–28 days in the presence of anti-CD3 stimulation and the cytokines interferon-gamma and interleukin-2. Prior to transfusion, the therapy product is expected to have minimum percentage of NK-like T cells.¹⁰¹ While having basic production guidelines makes reproducing this therapy achievable, we observed heterogeneity in the culture systems used to generate these cells, including the media, concentration of stimuli and cytokines used, and intervals of cytokine addition in culture. Characterization of the cell therapy product prior to transfusion to meet the guidelines was normally not provided.

Clinical parameters such as anticancer treatment history, demographics, and number of treatment cycles were also observed to be heterogeneous among the studies analyzed. These variables could contribute to the heterogeneity observed in our analysis that was not rectified by our subgroup analyses. As the studies identified were all undertaken in China, clinical trials in non-Chinese ethnicity are needed to confirm its efficacy outside of Chinese patients. Finally, the possibility of publication bias was raised as only a handful of studies reported negative outcomes of CIK/DC-CIK therapy for the efficacy endpoints assessed.

Despite these limitations, our data strongly support that complementing conventional treatment regimens with CIK/DC-CIK therapy in patients with CRC provides clinical benefits. By highlighting the parameters that contribute to the heterogeneity in the study designs, we suggest that standardization of these will lead to greater adoption of CIK therapy worldwide.

CONCLUSION

CIK therapy in combination with standard treatments, in particular chemotherapy, provides clinical benefits for patients with CRC. The benefit existed whether the included studies were prospective and randomized or not, strengthening the finding. CIK therapy was well tolerated, with fever being the most common adverse event. While DC therapy is commonly combined with CIK therapy for patients with CRC, our study suggests that this may not provide extra benefit. The findings support further evaluation of the clinical utility of CIK therapy in CRC.

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Competing interests None declared.

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Data availability statement All data relevant to the study are included in the article or uploaded as supplementary information.

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