

MiR-24-3p and various cancers: From a meta-analysis view

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Abstract

Background: A growing number of researches suggests that microRNAs (miRNAs) as oncogene or tumor suppressor genes play a fundamental role in various kinds of cancers. Among them, miR-24-3p, as a star molecule, is widely studied. However, the prognostic value of miR-24-3p is unclear and controversial. We conducted this meta-analysis to evaluate the prognostic value of miR-24-3p in a variety of cancers by integrated existing articles from four databases

Methods: PubMed, Embase, Web of Science, and Cochrane Library (last update in March 2020) were searched for approach literature. Hazard ratios (HRs) and odds ratios (ORs) were used to evaluate the association between miR-24-3p expression levels and prognostic value or clinicopathological characteristics, respectively.

Results: A total of 15 studies from 14 literature were finally qualified and concluded in the present meta-analysis. A significantly worse overall survival was observed in higher expression of miR-24-3p cancer group for OS(Overall survival) of log rank tests and cox multivariate regression by fixed effects model. Also, we found a significant correlation between elevated miR-24-3p levels to RFS (Recurrence-free survival) and DFS(Disease free survival). In addition, the pooled odds ratios (ORs) showed that evaluated miR-24-3p was also associated with the larger tumor size ($\geq 5\text{cm}$) and advanced TNM stage (II and III).

Conclusion: Built on the above findings, elevated expression levels of miR-24-3p may serve as a promising biomarker used to predict the worse prognosis of cancer patients.

1. Background

MicroRNAs (miRNAs), a kind of endogenous non-coding RNAs of 18–22 nucleotides in length, negatively regulate target genes expression at post-transcriptional level [1–3]. As either oncogenes or anti-oncogenes, they are found to play vital roles in a wide range of fundamental biological processes, such as proliferation[4, 5], differentiation[6, 7], apoptosis[8, 9], cell cycle[10–13], metastasis[14–16], stress response[17–19], metabolic[20–22] and etc. Owing to its detectability and stability in tissues, marrow or blood, a growing number of studies suggests that miRNAs can serve as promising biomarkers for the prognosis of carcinomas[23].

MiR-24-3p (used name was miR-24), a master regulator from the gene cluster of miR-23a–24-27a, has been identified as an onco or oncosuppressor-miR and its expression is closely associated with cancer occurrence and development by recent studies[24, 25]. Previous studies showed that miR-24-3p was highly expressed in many carcinomas[26, 27]. In addition, the evaluated expression of miR-24-3p was also found to be associated with cancer prognosis and tumor clinicopathological features but there were some opposite consequences[28, 29]. Up to now, a number of studies have been investigate this molecule in many kinds of cancer, but most individual study have their own limits, for example, small sample size or obtaining controversial results, and so on.

Accordingly, to explore the clinical prognostic value of miR-24-3p in various cancers, we performed this systematic review and meta-analysis to give a better understanding.

2. Methods

2.1. Literature search strategy

In this meta-analysis, the statement was used to as a guideline[30]. We performed a literature search using the online databases including PubMed, Embase, Web of science (WOS) and Cochrane library from inception to March 2020. The terms “miR-24 OR microRNA-24 OR miRNA-24 OR miR24” and “cancer OR tumor OR neoplasm OR carcinoma OR malignancy” were used to determine the correlative literature.

2.2. Inclusion and exclusion criteria

The inclusion criteria were: (1) studies were published in English; (2) miR-24-3p was investigated in carcinomas; (3) studies were identified the correlation between miR-24-3p expression levels and the prognosis of cancer patients; and (4) studies were provided hazard ratio (HR) and its corresponding 95% confidence intervals (CIs) or sufficient data which can further to assess its HR. The exclusion criteria: (1) studies were published in non-English; (2) studies were case report, abstracts, reviews, letters or meta-analysis; (3) studies were not relevant to the prognostic of cancer patients or the prognosis data originated from TCGA; or (4) studies did not offer sufficient data to calculate the HRs and 95% CI.

2.3. Data extraction

Built on the above criteria, all included studies were managed separately by two investigators (H Wang and CY Chen) and any disagreement were further to examined by a third author (KK Ding). The following characteristics were collected: the first author's name, year of publication, nationality, cancer type, specimen, method of detection, sample size, type of miRNA, outcome, tumor stage, lymph node metastasis, cut-off value, follow-up time, HR and its corresponding 95% CI. Moreover, the clinicopathological parameter data were also collected from qualified articles. For studies which not provide HR and 95% CI, the data were extracted from the Kaplan-Meier curves via Engauge Digitizer version 4.1[31]. The Newcastle-Ottawa Scale (NOS) was used to assess the quality of the pooled studies. High quality required an NOS score ≥ 5 .

2.4. Statistical analysis

The present meta-analysis was assessed by Stata SE12.0 software, RevMan5.2 software and Engauge Digitizer 4.1 software. Pooled HRs with their CIs were applied to describe the correlation between the expression of miR-24-3p and relevant survival outcome (OS, DFS, RFS), and the relation between miR-24-3p and relevant clinicopathologic features were also described by pooled odds ratios (ORs) and their CIs. The heterogeneity was evaluated by I^2 statistics and Q

tests. $P < 0.05$ and / or $I^2 > 50\%$ were defined as significant heterogeneity and random effects model was further to used. In addition, sensitivity analysis was used to evaluate the contribution of each study to the pooled HR and we could further to estimate the stability of the consequence. Finally, we evaluate the potential publication bias by funnel plot, Begg's test and Egger's test. $P < 0.05$ was known as obvious publication bias[32].

3. Results

As is shown in Fig. 1, 1099 literatures were obtained from online databases PubMed, Embase, Web of science (WOS) and Cochrane library. After removing the duplicates, abstract, review, case report, meta-analysis, studies which were not written in English and unrelated researches, 86 articles were subsequently full-text review. Among these, 71 articles were further to removed according to these criteria: studies not on patient ($n = 2$), studies without survival data ($n = 45$), survival data from TCGA ($n = 5$), multiple miRNAs ($n = 1$), or insufficient data ($n = 19$). Eventually, a total of 15 studies from 14 articles were included. (Liu et al., 2014; Meng et al., 2017; Gao et al., 2015; Organista-NAVA et al., 2015; Zhao et al., 2015; Wang et al., 2016; Mori et al., 2016; Kerimis et al., 2017; Dong et al., 2018; Su et al., 2018; Pan et al., 2018; Zhou et al., 2018; Liu et al., 2018; Yan et al., 2019) (Table 1). The overall sample size is 1518 patients coming from 25 to 247 which from 4 countries. Among these studies, several types of cancer include lung cancer ($n = 3$)[33–35], hepatocellular carcinoma ($n = 2$)[36, 37], colorectal cancer ($n = 2$)[29, 38], nasopharyngeal carcinoma ($n = 2$)[39, 40], osteosarcoma ($n = 1$)[41], ALL ($n = 1$)[42], AML ($n = 1$)[42], advanced gastric cancer ($n = 1$)[28], esophageal cancer ($n = 1$)[43], head and neck squamous cell carcinoma ($n = 1$)[44]. As for OS, RFS and DFS, there were seven studies directly provide HRs and its 95% CI [28, 29, 35–39]. In addition, the remaining eight studies only provided Kaplan-Meier curves[28, 33, 34, 40–44]. All studies measured the miR-24-3p expression level by qRT-PCR (quantitative real-time polymerase chain reaction).

3.1. The association between miR-24-3p expression levels and the overall survival (OS)

Ten enrolled articles including eleven studies and 1212 patients were used to investigate the correlation between miR-24-3p expression levels and the OS by using log rank tests and presented the data of univariate. Generally, a significant correlation between miR-24-3p levels and OS (HR = 1.609, CI: 1.291–2.004, Figure. 2a). However, an obvious heterogeneity was also observed ($I^2 = 85.20\%$, $P = 0.000$, Table 2). Hence, the random effects model was followed in succession but the significance was disappeared (HR = 1.507, CI: 0.810–2.803, Table 2), indicating that the heterogeneity significantly influenced the consequence.

In order to explore the source of the heterogeneity, subgroup analyses were applied by factors including population (Asian(Chinese) and Non-Asian), sample size (≥ 100 and < 100), NOS scores (≥ 8 and < 8), specimen (tissue and non-tissue) tumor category 1 (solid tumor and non-solid tumor), tumor category 2 (digestive system and non-digestive system) and tumor (esophageal cancer, osteosarcoma, lung cancer, gastric cancer, colorectal cancer, ALL, AML and hepatocellular carcinoma). As a consequence, the heterogeneity was controlled successfully in six subgroups and all them have significant correlations: 1). The subgroup of non-Asian (HR = 2.615, CI: 1.668–4.099; $I^2 = 0.000\%$, $P = 0.693$). 2). The specimen derived from non-tissue (HR = 2.399, CI: 1.659–3.470; $I^2 = 0.000\%$, $P = 0.949$). 3) The sample size greater than or equal to 100 (HR = 2.779, CI: 2.051–3.766; $I^2 = 0.000\%$, $P = 0.873$). 4) The patients of hematologic tumor (HR = 2.425, CI: 1.491–3.944; $I^2 = 0.000\%$, $P = 0.751$). 5). The patients of hepatocellular carcinoma (HR = 2.607, CI: 1.756–3.871; $I^2 = 0.000\%$, $P = 0.637$). and 6). The patients of lung cancer (HR = 3.274, CI: 1.422–7.539, $I^2 = 0.000\%$, $P = 0.698$). In addition, significant correlations are also observed in the study of NOS score less than 8 by random effects model, which were consistent with the significance of the results by fixed effects model (Table 2). Moreover, significant correlations were observed between miR-24-3p expression levels and OS in the studies with the population derived from Asian(Chinese) (HR = 1.381, CI: 1.219–2.004), solid tumor (HR = 1.448, CI: 1.131–1.852), digestive system (HR = 1.705, CI: 1.291–2.253) and non-digestive system (HR = 1.461, CI: 1.021–2.090) by fixed effects model, while there were no significances identified in these groups when the random effects model was applied (Table 2). For patients of Colorectal cancer, the prognostic value of miR-24-3p expression levels to the OS was completely different (Kerimis D et al.[38] HR = 4.070, CI: 1.25–13.19; Gao Y et al.[29] HR = 0.285, CI: 0.139–0.584). Due to insufficient data, the consequence was lack of efficiency and the heterogeneity was also significant ($I^2 = 93.00\%$, $P = 0.000$). Therefore, more relevant studies are required to perform the analysis. Built on the above consequences, meta regression was further used, but there was no meaningful contribution identified to impact on the heterogeneity (Table 2). Subsequently, the sensitivity analysis was performed, but there was also no positive consequence (Figure. 2c). Next, funnel plots, Begg's test and Egger's test were implemented to assess the potential publication bias and two studies as the outliers were identified eventually (Figure. 2d) (Liu et al.[41] and Gao et al.[29]). After removing them, dramatically decline of the heterogeneity was observed ($I^2 = 34.60\%$, $P = 0.141$) in the overall analysis, and the significance of the prognostic effects of miR-24-3p expression was still obvious (Figure. 2b).

3.2. The independent role of miR-24-3p expression levels as a prognostic indicator

Five studies containing 775 patients implemented the cox multivariate regression to assess the prognostic value of miR-24-3p expression levels in carcinoma patients by adjusting other factors. The significant correlation of miR-24-3p expression levels to the OS (HR = 2.384, CI: 1.813–3.134) was observed by fixed effects model. However, the heterogeneity was relatively obvious ($I^2 = 82.30\%$, $P = 0.000$, Table 3) and the significance was vanished by random effects model (HR = 1.994, CI: 0.991–4.015). Homoplasticly, Subgroup analyses were applied to reduce the heterogeneity. As a result, the homogeneity was reached within the studies of sample size greater than or equal to 100 ($I^2 = 0.000\%$, $P = 0.861$), NOS less than 8 ($I^2 = 45.50\%$, $P = 0.176$) and the patients of hepatocellular carcinoma ($I^2 = 45.50\%$, $P = 0.176$). And the significant association was identified between miR-24-3p expression levels and OS with the sample size greater than 100 (HR = 3.369, CI: 2.414–4.701), NOS less than 8 (HR = 3.041, CI: 2.150–4.300) and the patients of hepatocellular carcinoma (HR = 3.041, CI: 2.150–4.300). In addition, the significant correlations were identified between miR-24-3p expression levels to the OS in the population from Asian(Chinese) (HR = 2.373, CI: 1.813–3.134), the specimen derived from tissue (HR = 2.448, CI: 1.804–3.323) and NOS larger than or equal 8 by fixed effects model, which become to no significance within those subgroups by random effects model (Table 3). For patients of Colorectal cancer (HR = 0.752, CI: 0.212–0.978), the prognostic value of miR-24-3p expression levels to the OS was also opposite (Kerimis D et al.[38] HR = 2.60, CI: 0.780–8.660; Gao Y et al.[29] HR = 0.456, CI: 0.394–1.434). Thus, more pertinent studies are required to perform the analysis. Similarly, there was no noteworthy contribution identified to greatly influence the

variation of HR by meta regression (Table 3). But the sensitivity analysis suggested that Gao et al.[29] has significant impact on the result (Figure. 4c). The heterogeneity was vanishing ($I^2 = 0.000\%$, $P = 0.591$, Figure. 4b) by removing this outlier and the correlation of miR-24-3p expression levels to the OS was also significant (HR = 3.039, CI: 2.268–4.074, Figure. 4b). Finally, funnel plots, Begg's test ($P = 0.734$) and Egger's test ($P = 0.460$) indicated that there was no bias. But, the number of enrolled studies was few, more data are needed to reinforce this result.

3.3. The correlation of miR-24-3p expression levels to the RFS /DFS

Except OS as a prognostic indicator, RFS and DFS are also be accepted as an evaluation criterion. Here, four studies reported RFS including 393 patients applied log rank tests, while only one also utilized cox multivariate regression. After pooling the HR, we observed a significant association between miR-24-3p expression levels to the RFS of log rank tests (HR = 2.315, CI: 1.491–3.594, figure. 5a) by fixed effects model. However, the heterogeneities were quite obvious ($I^2 = 66.70\%$, $P = 0.290$, Table 4). The random effects model was further implemented but the significance was disappeared (HR = 1.814, CI: 0.741–4.440), indicating that the heterogeneity influenced the consequences significantly. Furthermore, owing to limited number of statistics from cox multivariate regression, the sensitivity analysis and publication bias were only applied to analysis with data extracted from log rank tests. The sensitivity analysis result indicated that no studies had significant influence on the consequent (Figure. 5c). However, the investigation of potential publication bias identified an outlier (Figure. 5d, Wang S et al.[40]). After deleting this study, the heterogeneity was obvious declined ($I^2 = 45.30\%$, $P = 0.161$) and the significance of correlation between miR-24-3p expression levels and the RFS was not altered (HR = 2.575, CI: 1.642–4.029, Figure. 5b). Due to the limit included studies, more data are needed in order to enhance the result. In addition, there were only two studies containing 226 patients revealed the DFS statistics and almost no heterogeneity in both log rank tests and cox multivariate regression ($I^2 = 3.600\%$, $P = 0.309$, $I^2 = 0.000\%$, $P = 0.330$, respectively, Table 4) by used a fixed effects model. We also observed significant strong correlation between miR-24-3p expression levels to the DFS of both log rank tests (HR = 2.361, CI: 1.390–4.012) and cox regression tests (HR = 2.313, CI: 1.315–4.067) by fixed effects model.

3.4. Correlations between miR-24-3p levels and clinicopathological features among various carcinomas

Six studies containing 536 patients investigated the correlation of miR-24-3p expression levels to different clinical characteristics. As showed in Table 5, miR-24-3p expression levels were significant correlation with tumor size (OR = 1.655, CI: 1.124–2.437) by the fixed effects model with lesser heterogeneity ($I^2 = 37.50\%$, $P = 0.184$). In addition, there were no significance identified in the correlation between age (OR = 0.684, CI: 0.357–1.310), gender (OR = 1.286, CI: 0.758–2.107), lymph node metastasis (OR = 1.591, CI: 0.758–3.339) or TNM stage (OR = 1.437, CI: 0.959–2.154) with the expression levels of miR-24-3p. There were no heterogeneity in the analysis of age ($I^2 = 0.000\%$, $P = 0.525$) and gender ($I^2 = 0.000\%$, $P = 0.842$), but the heterogeneity of lymph node metastasis and TNM stage were obviously ($I^2 = 70.90\%$, $P = 0.064$; $I^2 = 85.50\%$, $P = 0.000$, respectively). In order to decrease the heterogeneity, sensitivity analysis and publication bias were further investigated to each of them. As a result, an outlier was found (Liu et al.[36]) in the TNM stage. After removing the outlier, the heterogeneity was dramatically decreased from 85.50–0.000% and the associations between high miR-24-3p expression levels to advanced TNM stage were significant (OR = 2.328, CI: 1.490–3.637). (Figure. 6). Moreover, there was no potential publication bias about TNM stage by funnel plot, Begg's test ($P = 0.086$) and Egger's test ($P = 0.734$). For the analyze of lymph node metastasis, there were only two studies and have obvious opposite result (Pan et al.[33], OR = 2.974, CI: 1.101–8.037; Zhou et al.[35], OR = 0.725, CI: 0.238–2.208). Due to insufficient data, the consequence would be lack of efficiency and the reasons of heterogeneity were unacceptable. Thus, more pertinent studies are required to perform the analysis.

4. Discussion

It is of great importance to explore prognostic biomarkers with the patients of carcinoma as specific biomarkers can further help to directly stratify patients and effectively guide clinical decision-making. MiR-24-3p, as an oncogene or tumor suppressor, plays a key role in the occurrence, progression and metastasis of human carcinoma was realized by more and more researchers gradually[24, 25]. Quan et al. [45] had made a meta-analysis to research the correlation between miR-23a/24-2/27a cluster with human cancers, but they only had limited data to draw a conclusion that high expression levels of miR-23a/24-2/27a indicated a worse prognosis and no further analyzing the correlation between miR-24-3p expression levels to the clinicopathological characteristics. Subsequently, more and more studies which focusing on the miR-24-3p expression levels with cancer progression, metastasis and prognosis of patients were carried on[4–7, 14–16]. Thus, the exact role of miR-24-3p on the clinical prognosis of patients in various human carcinomas still need to further investigate. In this meta-analysis, total 15 studies including 1518 people were recruited. Among them, ten studies containing 1212 patients provided the statistics of the OS by log rank tests. By the pooling strategy, we know that the elevated miR-24-3p expression levels were linked to worse prognosis of cancer patients. Subsequently, several approaches were put in place to investigate the heterogeneity. First, subgroup analyses were conducted to identify the potential sources of heterogeneity. We found that the heterogeneity was achieved within the non-Asian population, non-tissue, the studies of sample size greater than or equal 100, hematologic tumor, hepatocellular carcinoma and lung cancer. But it was not controlled in other groups, such as the Asian population, studies of sample sizes less than to 100, solid tumor and so on. Second, the sensitivity analysis was used, but no outlier was identified to impact on the pooled results significantly. Third, two studies were identified as outliers by publication bias evaluation (Liu et al.[41] and Gao et al.[29]). After retrieving the data of outliers, we found that the specimen recruited in them were all from Asian (Chinese), tissue and the sample size less than 100 which all have strong heterogeneity. Besides, Liu et al.[41] was the sole study that focused on the Osteosarcoma and Gao et al.[29] had an opposite conclusion with Kerimis et al.[38] who also investigated the miR-24-3p expression levels to colorectal cancer. After removing those two outliers, greatly declined of the heterogeneity was observed. Built on the mentioned above, these two studies could be the major sources of heterogeneity. However more relevant data are needed to further investigate because of the limit number of studies. There were five studies including 775 patients obtained the data of HRs by cox multivariate regression. Cox multivariate regression has been known as an effective approach because it can evaluate the contribution of each factor independently by adjusting other factors[46]. Thus, the consequences by cox multivariate regression are always considered as independent effects of each factor on the clinical outcome. As a result, we found that the significance was inconsistent among different effects model. This phenomenon suggested that the heterogeneity was relatively obvious and the consequences were instable. Through the subgroup analyses, we found that the heterogeneity was declined in hepatocellular carcinoma and achieved in

the studies of sample size larger than or equal 100. In addition, the sensitivity analysis identified one outlier, Gao et al.[29] who has an opposite conclusion with Kerimis et al[38]. After removing this study, the heterogeneity had been significantly vanished. High miR-24-3p expression had a significantly worse survival and there was no publication bias. Thus, the power of miR-24-3p expression levels might serve as an independent prognostic indicator and we need more data to reinforce this conclusion. Also, we detected additional indexes such as RFS and DFS. MiR-24-3p expression levels were deemed to be significantly associated with DFS of statistics extracted from both log rank tests and cox regression analysis. For the RFS of cancer patients, only the fixed effects model revealed a significant correlation between miR-24-3p expression with this prognostic index and the heterogeneity was palpable. We identified an outlier (Wang et al.[40]) through publication bias evaluation. After removing this study, the heterogeneity was declined and the significance of association between miR-24-3p expression levels and the RFS was not altered.

As for the clinicopathological parameters, six studies including 536 patients had evaluated the association of miR-24-3p expression levels to the distinctive clinical parameters. The over-expression of miR-24-3p was found to be significantly related to larger tumor size by fixed effects model. Moreover, we found a significant heterogeneity between miR-24-3p expression levels to TNM stage. Applying sensitivity analyses, we identified one study (Liu et al.[36]) which had greatly impact on the result for the TNM stage. After removing this study, the heterogeneity completely disappeared, the association between miR-24-3p expression levels to the TNM stage was also significant. In addition, there were only two studies about the lymph node metastasis are enrolled and the conclusion might be not reliable. The analyzes of clinical features of a definite carcinoma should be normalized for the cut-off values, the feature categories and so on, to enrich the enrolled cases and characteristics for the meta-analysis.

As far as we know, this meta-analysis was the most comprehensive and systematic one to explore the correlation between the miR-24-3p expression levels with the prognosis of cancer patients in depth. Subgroup analysis, meta regression, sensitivity analysis and publication bias had been used to investigate the possible source of the heterogeneity to the greatest extent[47]. In spite of this, several flaws were hard to avoid in this meta-analysis. First, inevitable limitation from insufficient data in this analysis (only 15 studies with 1518 patients). Second, the cut-off values of the miR-24-3p expression levels were not exactly among those studies, thus, the accuracy of prognostic results may be influenced. Third, part of HRs was calculated from the survival curves which may cause some bias. Four, the number of recruited studies for DFS, RFS and clinicopathological features analyses were relatively insufficient. Taking above reasons into account, we need better designed and large sample size studies for further research before applying miR-24-3p as a prognostic biomarker of tumor in clinical applications.

5. Conclusions

The over expression of miR-24-3p was an underlying risk of poor prognosis in various human carcinomas, especially in hepatocellular carcinoma and lung cancer. As for other types of carcinomas, the results are not yet stable and more studies including normalized research conditions are required to further identify miR-24-3p prognostic values. In addition, high miR-24-3p expression levels were linked to the progression of cancers, developing more malignant behavior, such as larger tumor sizes and the advanced TNM stages.miR-24-3p expression levels could serve as a potential prognostic marker of human carcinoma.

Abbreviations

WOS

Web of science, OS:Overall survival, DFS:Disease-free survival, RFS:Recurrence-free survival, DMFS:Distant metastasis-free survival, NOS:Newcastle-Ottawa scale scores, U:univariate, M:multivariate, qRT-PCR:Quantitative Real-time PCR, 95%CI:95% confidence interval, Fixed:Fixed effects model, HR:hazard ratio, Random:Random pooling model, HCC:Hepatocellular carcinoma, CRC:Colorectal cancer, Advanced GC:Advanced Gastric cancer, ALL:Acute Lymphocytic Leukemia, AML:Acute Myelocytic Leukemia, NPC:Nasopharyngeal carcinoma, HNSCC:Head and neck squamous cell carcinoma, NSCLC:Non-small cell lung carcinoma.

Declarations

-Ethics approval and consent to participate:

All analyses were based on previous published studies,thus no ethical approval and patient consent are required.

-Consent for publication

All subjects participating in the image acquisition signed the consent form

-Availability of data and material

The authors declare that all data supporting the findings of this study are available within the article and the enrolled articles for meta-analysis.

The datasets generated and/or analysed during the current study are available in the PubMed, Embase, Web of science (WOS) and Cochrane library repository.

PubMed: <https://pubmed.ncbi.nlm.nih.gov/>

Embase: <https://www.embase.com/login>

Web of science: <http://apps.webofknowledge.com>

Cochrane library: <https://www.cochranelibrary.com/>

-Competing interests

The authors declare that they have no competing interests

-Funding

Not applicable

-Authors' contributions

He Wang and Chunyang Chen contributed equally to this work and shall share first authorship. Study design and basic data extraction: He Wang, Chunyang Chen, Weijie Zhang, Jianquan Hou. Data extraction from the Kaplan–Meier curves: He Wang, Chunyang Chen and Weijie Zhang. Third party evaluation: Keke Ding. Manuscript composition: He Wang and Chunyang Chen. All authors read and approved the final manuscript.

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Tables

Table1:

Note: The dashes mean no data

HCC: Hepatocellular carcinoma, CRC: Colorectal cancer, Advanced GC: Advanced Gastric cancer, ALL: Acute Lymphocytic Leukemia, AML: Acute Myelocytic Leukemia, NPC: Nasopharyngeal carcinoma, HNSCC: Head and neck squamous cell carcinoma, NSCLC: Non-small cell lung carcinoma, OS: Overall survival,

DFS: Disease-free survival, RFS: Recurrence-free survival, DMFS: Distant metastasis-free survival, NOS: Newcastle-Ottawa scale scores, U: univariate, M: multivariate, qRT-PCR: Quantitative Real-time PCR

Study(year)	Country	Malignancy	Sample type	Sample(F/M) size(high/low)	Assay	Survival	miRNA	HR (95%CI)	Follow-up	Cut-off value
Meng et al. (2014)	China	HCC	Blood	72(36/36)	qRT-PCR	OS/DFS	miR-24-3p	OS: 2.364(1.341–4.167) U(Reported) 2.141(1.158-3.960) M(Reported) DFS: 2.117(1.197–3.744) U(Reported) 2.055(1.114–3.792) M(Reported)	60	Median
Meng et al. (2014)	China	CRC	Tissue	95(48/47)	qRT-PCR	OS	miR-24-3p	OS: 0.285(0.139-0.584) U(Reported) 0.456(0.212-0.978) M(Reported)	60	Median
Kerimis et al. (2017)	Greece	CRC	Tissue	154(115/39)	qRT-PCR	OS/DFS	miR-24-3p	OS: 4.070(1.250–13.19) U(Reported) 2.600(0.780–8.660) M(Reported) DFS: 4.730(1.120–19.89) U(Reported) 4.510 (1.050–19.33) M(Reported)	120	0.40RQU
Dong et al. (2018)	China	Advanced GC	Tissue	247	qRT-PCR	OS	miR-24	OS: 2.945(1.344-4.575) U(Reported) 3.162 (1.324-4.544) M(Reported)	15	$\Delta\Delta Cq=-2$
Organista-NAVA et al. (2015)	Mexico	ALL	Marrow	111(50/61)	qRT-PCR	OS	miR-24	OS: 2.450(1.500-4.000) U(K-M Curve)	about120	8.22fold(75p)
Organista-NAVA et al. (2015)	Mexico	AML	Marrow	36(18/18)	qRT-PCR	OS	miR-25	OS: 1.320(0.030-58.25) U(K-M Curve)	about120	2.54fold(75p)
Su et al. (2018)	China	NPC	Tissue	120(60/60)	qRT-PCR	DMFS	miR-24	DMFS: 0.350(0.120-0.980) M(Reported)	about168	Median
Wang et al. (2016)	China	NPC	FFPE tissue	25(4/21)	qRT-PCR	RFS	miR-24-3p	RFS: 0.180(0.020-1.660) U(K-M Curve)	about117	-
Liu et al. (2014)	China	HCC	Tissue	207(116/91)	qRT-PCR	OS/RFS	miR-24	OS: 2.860(1.650-4.970) U(K-M Curve) 3.580(2.360-5.460) M(Reported) RFS: 4.280(2.150-8.520) U(K-M Curve) 4.750(2.660-8.470) M(Reported)	130 100	-
Zhou et al. (2018)	China	Lung cancer	Tissue	50(25/25)	qRT-PCR	OS	miR-24	OS: 2.392 (0.400-14.45) U(Reported)	80	Median
Mori et al. (2016)	Italy	HNSCC	Tissue	108(52/56)	qRT-PCR	RFS	miR-24	RFS: 1.770(1.040-3.800) U(K-M Curve)	about70	Median
Yan et al. (2019)	China	Esophageal cancer	Tissue	86(34/52)	qRT-PCR	OS	miR-24	OS: 0.540(0.210-1.350) U(K-M Curve)	20	-
Zhao et al.	China	NSCLC	Tissue	53(39/14)	qRT-	RFS	miR-	RFS: 1.740(0.390-	30	Median

(2015)					PCR		24-3p	7.200) U(K-M Curve)		
Liu et al. (2018)	China	Osteosarcoma	Tissue	84(42/42)	qRT-PCR	OS	miR-24	OS: 0.310(0.160-0.630)U(K-M Curve)	50	Median
Pan et al. (2018)	China	Lung cancer	Tissue	70(41/29)	qRT-PCR	OS	miR-24	OS: 3.570(1.390-9.150)U(K-M Curve)	60	-

Table2:

Note: 95%CI: 95% confidence interval, Fixed: Fixed effects model, HR: hazard ratio, Random: Random pooling model

	Subgroups	No. of studies	No. of patients	Pooled HR(95%CI)		Meta regression p-value	Heterogeneity	
				Fixed	Random		I ²	p-value
Overall	OS	11	1212	1.609(1.291-2.004)	1.507(0.810-2.803)		85.20%	0
Population	Asian (Chinese)	8	911	1.381(1.073-1.777)	1.271(0.585-2.762)	0.341	88.50%	0
	Non-Asian	3	301	2.615(1.668-4.099)	2.615(1.668-4.099)		0.00%	0.693
Specimen	Tissue	8	993	1.291(0.982-1.698)	1.336(0.574-3.108)	0.505	88.40%	0
	Non-Tissue	3	219	2.399(1.659-3.470)	2.399(1.659-3.470)		0.00%	0.949
Sample size	≥100	4	719	2.779(2.051-3.766)	2.779(2.051-3.766)	0.069	0.00%	0.873
	≤100	7	493	0.882(0.641-1.212)	0.951(0.376-2.404)		85.30%	0
NOS	≥8	6	700	1.078(0.771-1.507)	1.336(0.456-4.094)	0.688	89.60%	0
	≤8	5	512	2.177(1.627-2.913)	1.932(1.152-3.241)		59.80%	0.041
Tumor Category1	Solid tumor	9	1065	1.448(1.131-1.852)	1.427(0.687-2.960)	0.675	87.50%	0
	Hematologic tumor	2	147	2.425(1.491-3.944)	2.425(1.491-3.944)		0.00%	0.751
Tumor Category2	Digestive system	6	945	1.705(1.291-2.253)	1.520(0.676-3.420)	0.982	87.40%	0
	Non-Digestive system	5	267	1.461(1.021-2.090)	1.505(0.466-4.863)		85.50%	0.891
Tumor	Esophageal Cancer	1	86	0.540(0.210-1.350)	0.540(0.210-1.350)	—	—	—
	Osteosarcoma	1	84	0.310(0.160-0.630)	0.310(0.160-0.630)	—	—	—
	Gastric cancer	1	247	2.945(1.344-4.575)	2.945(1.344-4.575)	—	—	—
	ALL	1	111	2.450(1.500-4.000)	2.450(1.500-4.000)	—	—	—
	AML	1	36	1.320(0.030-58.25)	1.320(0.030-58.25)	—	—	—
	Hepatocellular carcinoma	2	279	2.607(1.756-3.871)	2.607(1.756-3.871)	—	0.00%	0.637
	Lung cancer	2	120	3.274(1.422-7.539)	3.274(1.422-7.539)	—	0.00%	0.698
	Colorectal cancer	2	249	0.585(0.317-1.080)	1.032(0.076-13.954)	—	93.00%	0

Table3:

Note: 95%CI: 95% confidence interval, Fixed: Fixed effects model, HR: hazard ratio, Random: Random pooling model

Subgroups		No. of studies	No. of patients	Pooled HR(95%CI)		Meta regression p-value	Heterogeneity	
				Fixed	Random		I ²	p-value
Overall	OS	5	775	2.384(1.813-3.134)	1.994(0.991-4.015)		82.30%	0
Population	Asian (Chinese)	4	621	2.373(1.792-3.142)	1.897(0.848-4.242)	0.801	86.70%	0
	Non-Asian	1	154	2.600(0.780-8.660)	2.600(0.780-8.660)		—	—
Specimen	Tissue	4	703	2.448(1.804-3.323)	1.940(0.765-4.9221)	0.932	86.60%	0
	Blood	1	72	2.141(1.158-3.96)	2.141(1.158-3.960)		—	—
Sample size	≥100	3	608	3.369(2.414-4.701)	3.369(2.414-4.701)	0.157	0.00%	0.861
	<100	2	167	1.166(0.722-1.883)	1.005(0.221-4.575)		89.50%	0.002
NOS	≥8	3	496	1.593(1.020-2.488)	1.530(0.402-5.818)	0.502	87.20%	0
	<8	2	279	3.041(2.150-4.300)	2.914(1.778-4.774)		45.50%	0.176
Tumor	Colorectal cancer	2	249	0.752(0.394-1.434)	1.021(0.186-5.594)	—	82.50%	0.017
	Hepatocellular carcinoma	2	279	3.041(2.150-4.300)	2.914(1.778-4.774)	—	45.50%	0.176

Table4:

Note: 95%CI: 95% confidence interval, Fixed: Fixed effects model, HR: hazard ratio, Random: Random pooling model

		No. of studies	No. of patients	Pooled HR(95%CI)		Heterogeneity	
				Fixed	Random	I ²	p-value
RFS	Univariate	4	393	2.315(1.491-3.594)	1.814(0.741-4.440)	66.70%	0.029
DFS	Univariate	2	226	2.361(1.390-4.012)	2.386(1.362-4.180)	3.60%	0.309
	Multivariate	2	226	2.313(1.315-4.067)	2.313(1.315-4.067)	0.00%	0.33

Table5:

Note: 95%CI: 95% confidence interval, Fixed: Fixed effects model, HR: hazard ratio, Random: Random pooling model

Clinicopathological parameters	No. of studies	No. of patients	Pooled OR (95%CI)		Heterogeneity	
			Fixed	Random	I ²	p-value
Gender	5	329	1.286(0.758-2.107)	1.286(0.758-2.107)	0.00%	0.842
Age	2	156	0.684(0.357-1.310)	0.684(0.357-1.310)	0.00%	0.525
Tumor size	4	433	1.655(1.124-2.437)	1.605(0.985-2.689)	37.50%	0.184
Lymph node metastasis	2	120	1.591(0.758-3.339)	1.503(0.377-5.989)	70.90%	0.064
TNM stage	5	486	1.437(0.959-2.154)	1.419(0.469-4.291)	85.50%	0

Figures

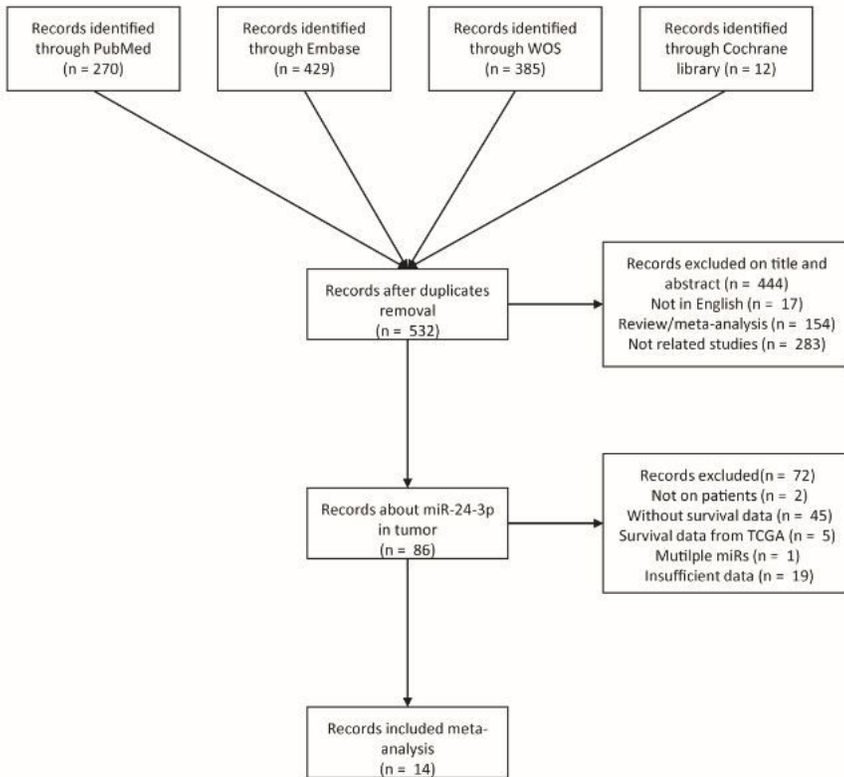


Figure 1

The flow chart of the meta-analysis

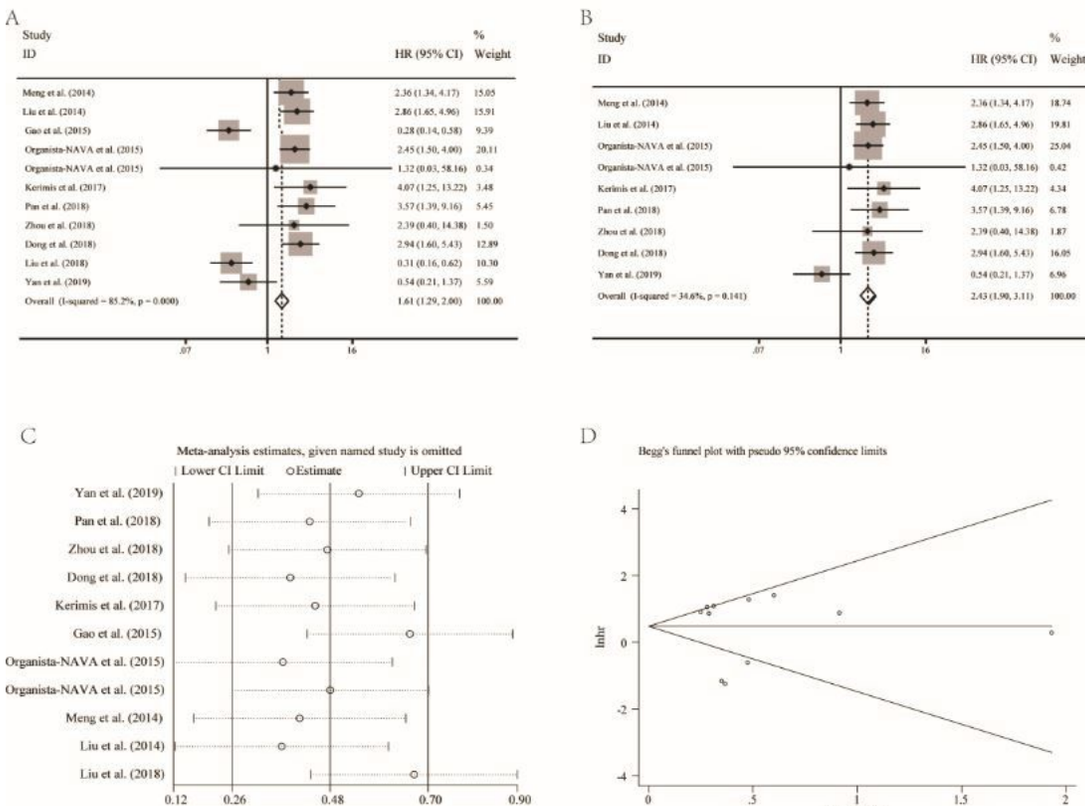


Figure 2

Association between miR-24-3p expression levels and (A) overall survival and (B) overall survival without the outliers as well as corresponding (C) sensitivity analysis and (D) publication bias evaluation

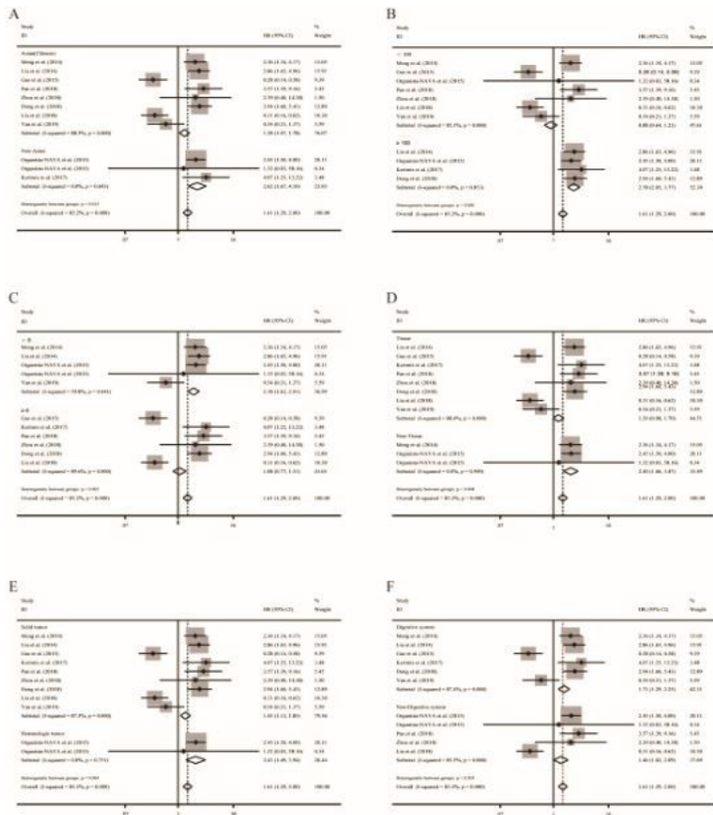


Figure 3 Subgroup analyses of (A) population (Asian and Non-Asian), (B) sample sizes (<100 and ≥100), (C) NOS scores (<8 and ≥8), (D) specimen (Tissues and Non-Tissue), (E) tumor category1 (Solid tumor and Hematologic tumor), (F) tumor category2 (Digestive system and Non-Digestive system) for overall survival

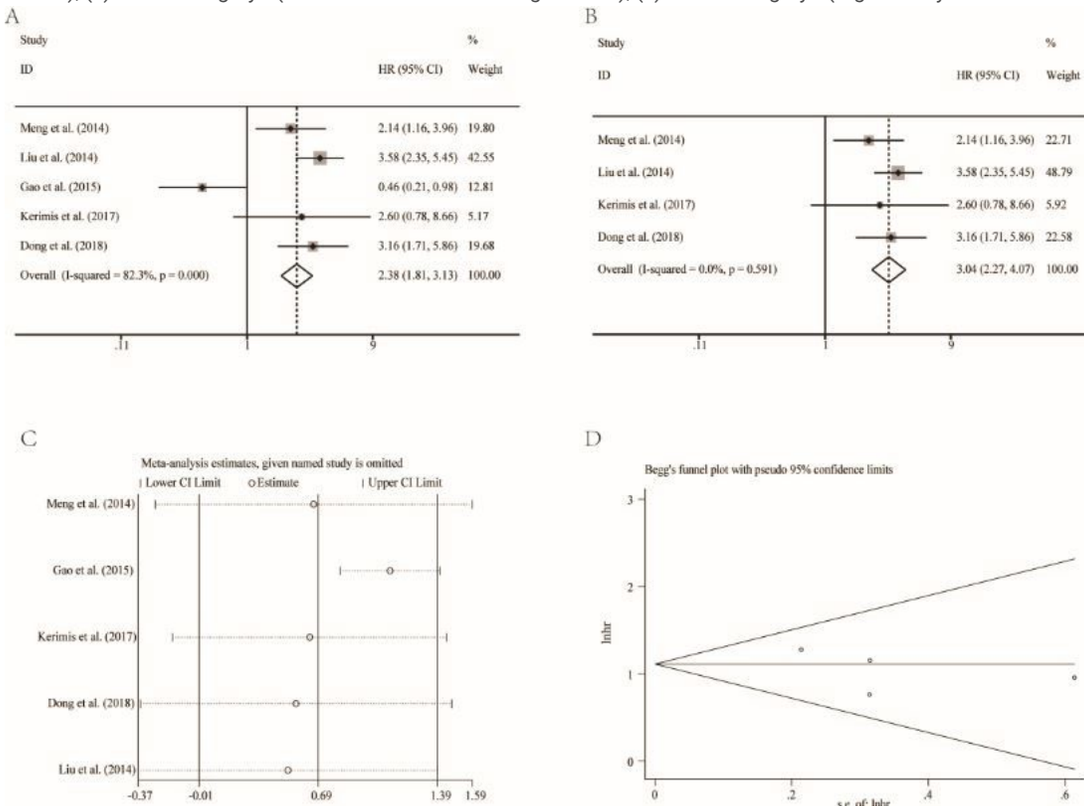


Figure 4 The independent role of miR-24-3p as a prognostic indicator for (A) overall survival, (B) overall survival without outliers, and (C) sensitivity analysis, (D) publication bias evaluation

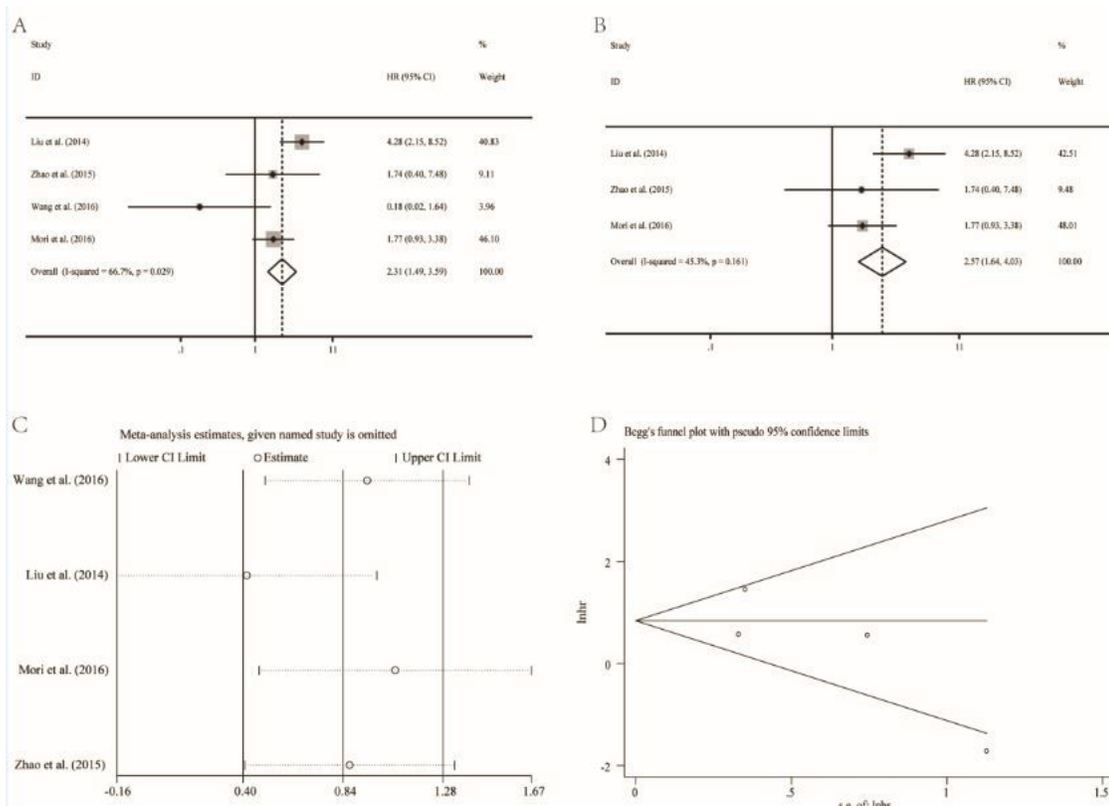


Figure 5 Association between miR-24-3p expression levels and (A) recurrence-free survival and (B) recurrence-free survival without the outliers as well as corresponding (C) sensitivity analysis and (D) publication bias evaluation

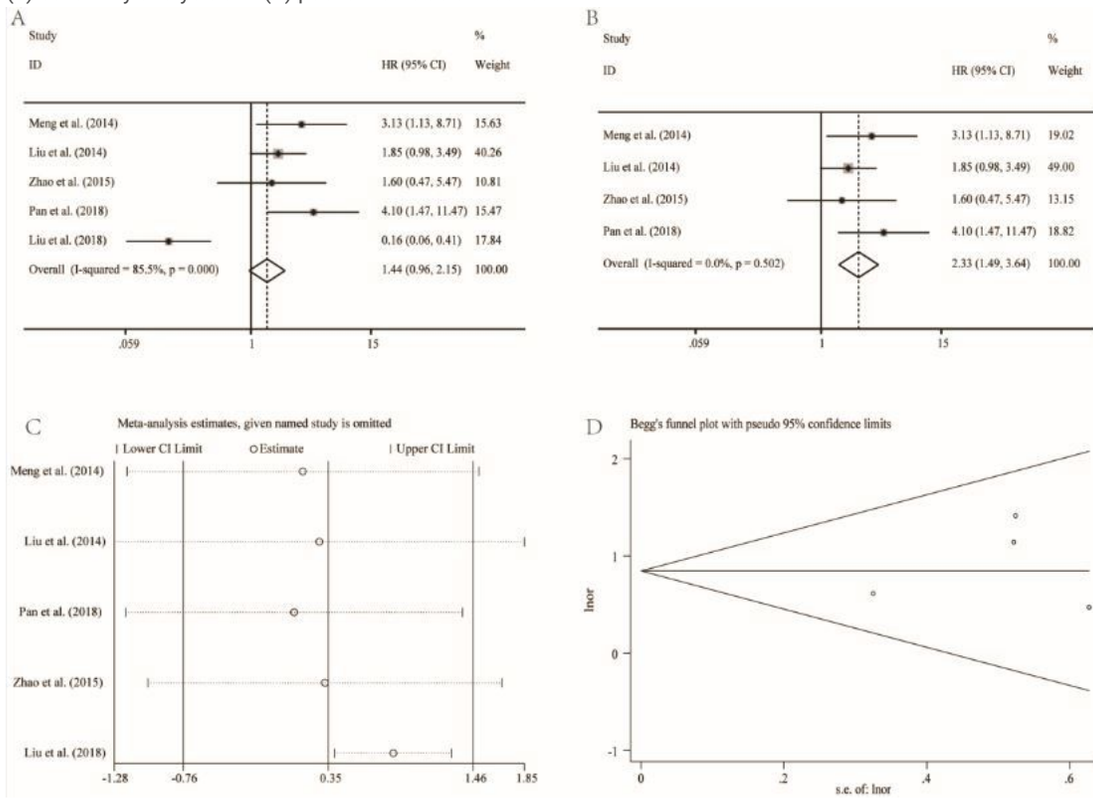


Figure 6 Association between miR-24-3p expression level and TNM stages of cancer patients, (A) overall pooling result, (B) pooling result without the outliers, and (C) sensitivity analysis, (D) publication bias evaluation

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