A systematic review and meta-analysis to compare the efficacy of conbercept with ranibizumab in patients with macular edema secondary to retinal vein occlusion

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Abstract

Background: The objective of this review and meta-analysis is to investigate the efficacy of conbercept and ranibizumab, combined with or without laser photocoagulation, in patients with macular edema secondary to retinal vein occlusion (RVO-ME).

Methods: Several databases have been used to identify relevant publications. After screening, a meta-analysis was conducted to compare conbercept and ranibizumab with the support of RevMan 5.3 (Cochrane Library Software, Oxford, UK).

Results: In this study, 9 randomized controlled trials and 6 retrospective trials were included with a total of 1180 patients. No significant difference was found in best corrected visual acuity (BCVA) or central macular thickness (CMT) in the baseline parameters [BCVA (weighted mean difference (WMD): –0.01; 95% confidence interval CI: –0.03 to 0.01; P = .17), CMT (WMD: 20.14; 95% CI: –26.70 to 66.97; P = .40)]. No significant differences were found in the improvements of BCVA and adverse events (AEs) between the 2 groups after injection of loading dosage [the 1st month BCVA (WMD: –0.01; 95% CI: –0.04 to 0.02; P = .54), the 3rd month BCVA (WMD: –0.02; 95% CI: –0.05 to 0.01; P = .23), the 6th month BCVA (WMD: –0.02; 95% CI: –0.05 to 0.01; P = .27), AEs (odds ratio: 0.84; 95% CI: 0.38 to 1.84; P = .66)]. However, there were significant differences between conbercept and ranibizumab treatment in terms of CMT [1st month CMT (WMD: –11.70; 95% CI: –19.71 to –3.68; P < .01), 3rd month CMT (WMD: –10.08; 95% CI: –15.62 to –4.53; P < .01), 6th month CMT (WMD: –15.83; 95% CI: –22.88 to –8.78; P < .01)] and the number of injections (WMD, –0.36; 95% CI: –0.68 to –0.40; P = .03).

Conclusion: The current pooled evidence suggested that both therapies of intravitreal conbercept and intravitreal ranibizumab with or without laser photocoagulation are effective in vision function in RVO-ME patients, and confirmed that conbercept has advantages over ranibizumab in terms of CMT and the number of injections for treating RVO-ME. In addition, conbercept has the statistically same visual gains and safety as ranibizumab in RVO-ME patients. Longer-term follow-up surveys on the safety and effectiveness of these 2 treatment regimens are required.

Abbreviations: AEs = adverse events, BCVA = best corrected visual acuity, CI = confidence interval, CMT = central macular thickness, RVO-ME = retinal vein occlusion macular edema, WMD = weighted mean difference.

Keywords: conbercept, ranibizumab, retinal vein occlusion

1. Introduction

Retinal vein occlusion (RVO) is the second most common vascular disease of the retina, with the first being diabetic retinopathy, in which various complications such as macular edema, may occur with a consequent visual impairment or visual loss.\textsuperscript{[1]} RVO also can lead to many serious complications, including retinal hemorrhage, macular edema, and neovascular glaucoma. Among these complications, macular edema secondary to RVO is the main cause of visual impairment and visual loss.\textsuperscript{[1]} RVO based on localization of venous occlusion is mainly divided into 3 types, including branch retinal vein occlusion (BRVO) that accounts for 80% of the RVO, hemi-retinal vein occlusion (HRVO) as well as central retinal vein occlusion (CRVO).\textsuperscript{[3]} According to the diagnostic criteria of American CRVO group,\textsuperscript{[5]} CRVO can be subdivided into 2 types on the basis of fluorescein fundus angiography (FFA):

1. perfused CRVO, characterized by retinal circulation stasis, blood capillary leakage, and capillary without perfusion area < 10 PD, still has blood perfusion. This type generally has a better prognosis;
(2) nonperfused CRVO is marked by retinal capillary without perfusion area > 10 PD.

Clinical studies have shown\(^{[1]}\) that 34% of perfused CRVO patients can develop into nonperfused CRVO patients. A multicenter epidemiological study\(^{[4]}\) showed that the global prevalence of CRVO is as high as 0.8%. The incidence of CRVO in China is 0.1%\(^{[5]}\). Therefore, more than 1.3 million people in China suffer from visual impairment or loss caused by CRVO every year.\(^{[5]}\) BRVO and CRVO are of the most clinically relevant types of laser photocoagulation. Owing to the damages to the retinal pigment epithelium, further damaging some photoreceptors in the hypoxic area and reducing the retinal oxygen consumption.\(^{[6]}\) And lasers directly destroy abnormal neovascularization, reducing the leakage of neovascularization. At the same time, the photocoagulation spots produce adhesion, so that the edema of the retina is closer to the choroidal capillaries to obtain a richer blood supply, promoting the absorption of edema and the retinal recovery of the structure and function. Grid-pattern and scatter photocoagulation are 2 types of laser photocoagulation. Owing to the damages to the retina, and limited visual acuity improvements from a long-term perspective, either grid-pattern or scatter laser coagulation has been employed in combination with anti-angiogenic therapy for RVO-ME patients non-responsive to anti-angiogenic agent monotherapy.\(^{[7]}\) Under normal physiological conditions, the tight junctions between retinal capillaries, together with the drainage function of the retinal pigment epithelium (RPE), jointly prevent the accumulation of fluid under the neurosensory layer or the RPE layer. Vascular endothelial growth factor (VEGF) has been proven to play a key part in the abnormal pathophysiologic process.\(^{[8]}\) Various factors, including hypoxia and ischemia as well as other stimuli, could up-regulate VEGF expression, which was found to be increased in the fluids of RVO patients’ eyes.\(^{[9]}\) In addition, a higher intravitreal VEGF level was noted to correlate with a severer clinical manifestation.\(^{[10]}\) Therefore, several anti-angiogenic drugs, such as conbercept, bevacizumab and ranibizumab, have been widely applied in the therapy of diabetic macular edema.\(^{[11,12]}\) They have been reported to provide a fast and obvious decrease of the central macular thickness (CMT) and a marked improvement in best corrected visual acuity (BCVA) in the therapeutic process of macular edema. Ranibizumab (or Lucentis; Novartis, Basel, Switzerland), which is widely adopted in wet age-related macular degeneration patients, also applied in diabetic macular edema patients, is a high-affinity recombinant antigen-binding fragment. The 48 kDa drug binds to all kinds of receptors of VEGF-A\(^{[11,13]}\) Conbercept (also known for Lumitin, or KH903, Kang Hong Biotech Co, Ltd, Sichuan, China) is a recombinant anti-angiogenic fusion protein, structurally similar to aflibercept (Eylea, Regeneron Pharmaceuticals, Eastview, NY), another fusion protein binding to all isoforms of placental growth factor (PIGF), VEGF-A, and VEGF-B. The 143 kDa drug engineered in Chinese hamster ovary cells (CHOC) was a protein with a human cDNA sequence.\(^{[14,15]}\) Moreover, it has been confirmed that conbercept has a more powerful binding affinity for VEGF than that of ranibizumab, owing to the special Fab fragment with the attached fourth extracellular domain of VEGFR-2.\(^{[16,17]}\) It also has been reported that some patients with diabetic macular edema who are ineffective to bevacizumab or ranibizumab (regardless of the number of injections) are still responsive to the treatment with conbercept.\(^{[16]}\) Several meta-analyses have assessed the effectiveness of intravitreal ranibizumab (IVR) and intravitreal conbercept (IVC) for treating RVO-ME patients.\(^{[18–21]}\) However, they focus either on therapy methods, such as intravitreal anti-VEGF agents, laser photocoagulation, and intravitreal dexamethasone, or on the different dosages of ranibizumab for treating this condition. – treatment regimens (conbercept 0.5 mg, ranibizumab 0.5 mg) are often adopted during the course of administering anti-VEGF agents. It is still unknown whether the conbecept and ranibizumab groups have the same outcomes in a large sample size. Therefore, we performed this meta-analysis based on 9 randomized controlled trials (RCTs) and 6 retrospective trials (Given that conbercept is currently only used in China to treat RVO-ME, all studies came from China) to compare the effectiveness of ranibizumab 0.5 mg and conbercept 0.5 mg, with or without laser photocoagulation.

2. Methods

2.1. Literature search

Relevant studies in human subjects were spotted by searching WanFang data, the China National Knowledge Infrastructure (CNKI), OVID, Springer, WOS, PubMed, EMBASE, Cochrane Library, and clinicaltrials.gov up to March 20, 2019. The search was performed, with the following Medical Subject Heading terms: “retinal vein occlusion or macular edema,” and “ranibizumab or Lucentis,” and “conbercept or KH903 or Lumitin,” with a filter restricting the results to only clinical trials. Language restrictions were limited within English and Chinese in the retrieval.

2.2. Inclusion and exclusion criteria

When papers met the following inclusion standards, the publications were selected:

1. study design – RCTs or retrospective trials;
2. intervention – IVR vs IVC treatment, or IVR plus laser photocoagulation vs IVC plus laser photocoagulation treatment;
3. population – RVO-ME patients;
4. duration – at least 3 months’ follow-up;
5. outcome variables – assessing at least 1 of the outcomes mentioned below.

Papers were excluded if

1. they were animal experimental studies, meeting abstracts, case reports, review articles, letters, or editorials; or
2. they were duplicated in those databases.

2.3. Outcome measures

The variables we have adopted were:

1. the BCVA presented in LogMAR, indicating functional improvement;
2. the CMT, measured with the aid of optical coherence tomography.
3. the number of AEs; and
4. the number of injections during the follow-up periods.

2.4. Data extraction
The methodological quality of the papers was separately evaluated by 2 reviewers, who also collected and sorted the data in an optimal procedure. All disagreements about data extraction were solved by discussion until the 2 reviewers reached a consensus. The following basic data were extracted from each study and filled out in a pre-set form comprising first author, publication year, study design, type of diagnosis, treatment regimen, period of follow-up, dosage, sample size, inclusion and exclusion criteria, and other records.

2.5. Quality assessment
In accordance with the Cochrane Handbook for Systematic Reviews of Interventions, the risk-of-bias tool was used to assess papers quality. Papers with factors that are not in line with the principles would have been marked with a high risk of bias. Six basic facets affecting the quality of RCTs were evaluated: completeness of outcome reporting, management of incomplete outcome data, personnel and outcome assessors, patient blinding, allocation concealment, as well as sequence generation.

2.6. Statistical analysis
This systematic review was conducted according to the Preferred Reporting Protocols for Systematic Reviews and Meta-Analyses (PRISMA) statement. The odds ratios (ORs) and weighted mean differences (WMDs) were adopted to compare dichotomous and continuous variables respectively. Chi-squared test was used to assess the statistical heterogeneity between studies, and I² statistic was adopted to evaluate the quantity of heterogeneity. A condition, I² > 50% and P < .05 at the same time, was regarded as heterogeneity. If there was insterstudy heterogeneity shown by evidence, a random-effects model was applied or else we adopted the fixed-effects model. All results were recorded with 95% confidence intervals (CI). We performed the subgroup analyses to separately estimate the effects of the adoption of laser photocoagulation at baseline and the last follow-up. We created standard funnel plots to visualize their symmetry and further to test the potential publication bias.

All analyses were based on published studies. Thus, no ethical approval and informed consent were required.

3. Results
3.1. Selection of qualified studies
This study selection procedure is presented in Figure 1. A total of 167 possibly related papers were noted by our extensive literature retrieval, 89 of which were dropped out after reviewing titles and abstracts, and 61 were excluded due to duplication or low quality. Of the remaining 17, 2 were excluded for being unable to extract outcomes of interest. Thus, 9 RCTs and 6 retrospective trials were included in the final meta-analysis.

3.2. Study characteristics
The basic characteristics of the 15 trials, including randomized controlled trials and retrospective trials are presented in Table 1. Nine RCTs and 6 retrospective trials, were enrolled in this study. All the trials were conducted in China. Among these trials, 4 trials were carried out to compare conbercept plus laser photocoagulation versus ranibizumab plus laser photocoagulation in patients with RVO. And 10 compared conbercept with ranibizumab, without the combination of laser photocoagulation. The sample size ranged from 30 to 384.

3.3. BCVA
Among all studies, only 1 study reported BCVA in ETDRS letters, and we transformed ETDRS into LogMAR. The BCVAs of the follow-up in the first (n = 844), third (n = 1032) and sixth (n = 541) month were presented. The pooled results revealed that no significant differences were found in BCVA before treatment (n = 1078) (WMD: −0.01; 95% CI: −0.03 to 0.01; P = .17) and after treatment (the 1st month BCVA (WMD: −0.01; 95% CI: −0.04 to 0.02; P = .54), the 3rd month BCVA (WMD: −0.02; 95% CI: −0.05 to 0.01; P = .23), the 6th month BCVA (WMD: −0.02; 95% CI: −0.05 to 0.01; P = .27]) between the conbercept and ranibizumab group. A similar outcome was observed between the IVC cohort and the IVR cohort, when we have the studies subgrouped according to whether adopting laser photocoagulation or not (Fig. 2). No substantial statistical heterogeneity was observed across all studies.

3.4. CMT
CMT is viewed as one of the most relevant prognostic factors for ME levels. Hence it was also evaluated in the current study. All the CMTs were recorded with the aid of optical coherence tomography pictures from the first time to the last time of the follow-up period in both the IVC cohort and the IVR cohort. The CMTs were collected in the first (n = 946), third (n = 1134), and sixth (n = 643) month during the follow-up. There was no statistically significant difference in CMT at baseline (n = 1180) when comparing the conbercept group with ranibizumab group (WMD: 20.14; 95% CI: −26.70 to 66.97; P = .40). However, CMT significantly differed between the conbercept group and ranibizumab group after treatment whether or not with the combination of laser photocoagulation (1st month CMT (WMD: −11.70; 95% CI: −19.71 to −3.68; P < .01), 3rd month CMT (WMD: −10.08; 95% CI: −15.62 to −4.53; P < .01), 6th month CMT (WMD: −15.83; 95% CI: −22.88 to −8.78; P < .01), RVO-ME patients applied with monthly injections of conbercept had a more obvious reduction of CMT from the initial thickness in comparison with RVO-ME patients applied with ranibizumab. Subgroup analyses were also conducted to compare the IVC group to the IVR group. All these analyses exhibited statistically significant differences showing the advantage of the IVC group over the IVR group (Fig. 3)

3.5. Number of intravitreal injections
Given that 1+PRN regimen and 3+PRN regimen were performed, we also compared the number of injections. Seven trials (n = 704) reported intravitreal injection numbers. There was a significant heterogeneity (I² = 81%) in this analysis, so a random-effects model was used. Pooled results indicated that (WMD, −0.36; 95% CI: −0.68 to −0.04; P = .03) there was a significant difference. In other words, conbercept groups experienced fewer injections, compared to ranibizumab groups (Fig. 4).
Records identified through authenticated database searching (n = 167)

Records after duplicates removed (n = 106)

Records screened (n = 17)

Records excluded after titles and abstracts screening (n = 89)

Full-text articles assessed for eligibility (n = 15)

Full-text articles excluded for no extractable data (n = 2)

Studies included in the final meta-analysis (n = 15)

Figure 1. Flow diagram of study selection and identification.

Table 1: Characteristics of included studies.

| Name and published yr | Collection period | Number of eyes | RVO type | Protocol | Design | Follow-up period (mo) |
|-----------------------|-------------------|----------------|----------|----------|--------|-----------------------|
| Bai S[24]             | 2017              | 40             | BRVO     | 1 + PRN + L | RCT    | 6                     |
| Chen B[23]            | 2019              | 40             | BRVO     | 1 + PRN   | RCT    | 3                     |
| Chen L[27]            | 2019              | 32             | CRVO + BRVO | 1 + PRN   | Retro  | 6                     |
| Chen T[25]            | 2018              | 179            | CRVO + BRVO | 1 + PRN + L | RCT    | 3                     |
| Chen X[26]            | 2018              | 50             | CRVO     | 3         | RCT    | 6                     |
| Huang Y[26]           | 2018              | 44             | CRVO     | 3         | Retro  | 6                     |
| Li F[36]              | 2017              | 18             | BRVO     | 1 + PRN   | RCT    | 6                     |
| Li Ting[31]           | 2017              | 23             | CRVO     | 3 + PRN   | Retro  | 6                     |
| Lian H[30]            | 2016              | 23             | CRVO     | 3 + PRN   | Retro  | 6                     |
| Lin L[37]             | 2018              | 11             | BRVO     | 1 + PRN   | RCT    | 3                     |
| Wang DX[34]           | 2016              | 20             | CRVO     | 3 + PRN + L | RCT    | 6                     |
| Yan H[31]             | 2018              | 16             | BRVO     | 1         | Retro  | 3                     |
| Zhang YY[32]          | 2018              | 30             | CRVO + BRVO | 1 + PRN   | RCT    | 6                     |
| Zhang ZQ[33]          | 2017              | 24             | BRVO     | 1 + PRN + L | RCT    | 6                     |
| Zhao XL[33]           | 2019              | 22             | BRVO     | 1 + PRN   | Retro  | 6                     |

IVC = intravitreal conbercept; IVR = intravitreal ranibizumab; CRVO = central retinal vein occlusion; BRVO = branch retinal vein occlusion; PRN = pro re nata; L = laser photocoagulation; RCT = random controlled trial; Retro = retrospective trial
3.6. AEs

AEs are worthy of paying attention, so we also analyzed the data to compare the number of AEs between the IVC cohort and the IVR cohort. Eight studies (n=848) reported AEs, including floaters in front of the eye, pain at the injection site, increased corneal edema, intraocular pressure, subconjunctival hemorrhage, anterior chamber inflammation, etc. The rest studies either did not report AEs or no adverse event occurred. No serious adverse event (vitreous hemorrhage, retinal proliferation, retinal tears, transient cerebral ischemia, cerebral hemorrhage) was reported in all the studies. Fixed-effects model analysis was...
adopted, owing to the heterogeneity test results ($P = .85, I^2 = 0\%$).
No statistical difference was found (Fig. 5) in the AEs between these 2 regimens (IVC 0.5mg and IVR 0.5mg) (OR: 0.74; 95% CI: 0.44 to 1.24; $P = .25$).

### 3.7. Publication bias and sensitivity analysis

In this study, the BCVAs of meta-analysis and the results of CMT were analyzed sensitively by the method of eliminating documents one by one. The results showed that the findings of meta-analysis before and after exclusion were consistent, indicating that the combined results were stable and reliable. Relatively symmetrical funnel plots showing the distribution indicated there was little potential publication bias, in spite of a trial of small sample size enrolled in this analysis (Fig. 6).

### 4. Discussion

Cone cells are relatively concentrated in the macula, which is responsible for the most sensitive part of central vision. Long-
term macular edema leads to a large number of apoptosis of cone cells and irreversible damages to visual function.\[^{[38]}\] Occlusive retinal veins result in ischemia and hypoxia at the same time, both of which are the stimuli of the release of VEGF.\[^{[39]}\] Then the destruction of tight junctions between capillaries by the increased level of VEGF increases vascular permeability, promoting fluid exudation and accumulation between the Henle fibers in the outer plexiform layer.\[^{[40]}\] Ranibizumab, a powerful-affinity recombinant segment, has the ability to block all corresponding receptors of VEGF-A, and has been confirmed to be a good choice in the therapy of some neovascular diseases of retina including RVO.\[^{[41]}\] Two large-size studies, BRAVO and HORIZON, respectively, have tested the effectiveness and potential AEs of ranibizumab for the treatment of BRVO-induced ME.\[^{[42,43]}\] Conbercept is now only available in China. It is a novel anti-angiogenic drug with a similar structure to aflibercept, sharing the same Fc segment of human immunoglobulin G(IgG) fused with VEGFR-1 and VEGFR-2. What makes these 2 agents different is that conbercept includes an additional VEGFR-2 binding domain, which possibly increases the association rate of VEGF and extends its periods of half-life in the human vitreous. Conbercept, in the treatment process, also has presented to be well tolerated and displayed its powerful effectiveness in other ME diseases.\[^{[41]}\]

In the present meta-analysis, we have reviewed the literature on treatment of RVO-ME regarding the effectiveness and side effects, which compared ranibizumab and conbercept. The analyzed results of these trials exhibited that conbercept and ranibizumab were feasible to treat RVO-ME. Furthermore, conbercept and ranibizumab were comparably effective, and both IVR and IVC led to marked visual improvements, demonstrated by the LogMAR, and noticeable reductions of CMT. The regimens to treat ME secondary to RVO mainly consist of 3+ pro re nata (PRN) and 1+PRN protocol. Intravitreal injection of either ranibizumab or bevacizumab, through PRN strategies, as the MARVEL study exhibited, has shown to be effective in BCVA improvement and CMT reduction.\[^{[44]}\] Miwa confirmed that 3+PRN and 1+PRN strategies experienced similar visual outcomes at the last time of the follow-up,\[^{[45]}\] which were similar to ours. However, conbercept group experienced fewer injections compared to ranibizumab group. A certain amount of VEGF is necessary to maintain a relatively stable balance of the choroidal and retinal microenvironment. Studies revealed that treatment with anti-VEGF agents could trigger the apoptosis of multiple types of cells in the retina, such as retinal ganglion cells (RGCs), and those within the inner nuclear layer such as amacrine cells as well as bipolar cells.\[^{[46,47]}\] Another study indicated that repeated intravitreal injections may lead to unexpected retinal degeneration and atrophy in the macula, bringing about ultimate visual function impairment.\[^{[48]}\] Therefore, the treatment with PRN strategies reduced the number of injections unnecessary and unfavorable side effects. In addition, a systemic review\[^{[49]}\] also indicated that prolonged high-level exposure to anti-VEGF...
harbored the potential for cerebrovascular accidents and increased risks for death. A latest meta-analysis showed that 0.5 mg of IVR regimen and 0.3 mg of that had the similar results with regard to BCVA and effectiveness as well as AEs, and that the number of IVC 0.5 mg (larger dose) monthly injection was obviously correlated with the reduction of CMT. Some researchers are concerned about the serious AEs of arterial thromboembolism with anti-VEGF treatment, but the current meta-analysis suggested that there was no such an adverse event that happened during the follow-up periods within the 2 groups. Our data revealed that intravitreal injection of conbercept or ranibizumab is safe for patients to treat RVO-ME. In an attempt to achieve reliable outcomes, we conducted subgroup analyses according to whether or not adopting laser photocoagulation, the results of which indicated that all subgroups did not substantially change the merged results. The majority of comparisons in the current meta-analysis showed no heterogeneity.

This analysis has numbers of advantages. Firstly, a relatively large sample size has been acquired. Secondly, this current meta-analysis meets rigorous inclusion and exclusion standards. In addition, reviewers have rigorously obeyed the statement of PRISMA and the rules of Cochrane Handbook for Systematic Reviews of Interventions through the whole document retrieval, literature filtration, quality assessment as well as data analysis, which makes the conclusion drawn by us more valid and reliable. Although this current study contains a complete record of the presently available trials on the effectiveness of IVC and IVR for treating RVO-induced macular edema, we also have to underline some limitations. The first disadvantage of this systematic review and meta-analysis is the enrolled trials with relatively short follow-up periods. Longer follow-up surveys after the first injection would have provided more accurate clinical findings with certainty in treatment recommendations. Another possible shortcoming presented in this study is that the enrolled studies consisted of 2 types of treating regimens, which may bring about heterogeneity, and compromise the final outcomes. However, subgroup analyses in this study have shown that the IVC with or without laser photocoagulation and IVR with or without laser photocoagulation subgroups presented statistically similar outcomes and did not change the pooled outcomes. Moreover, only a few trials revealed the effective rate, and they had different standards to define effective rate, which made it hard to analyze sufficient data for meaningful outcomes. Fourth, we have failed to subgroup-analysis the outcomes according to RVO type, because some trials did not declare the RVO type or combined CRVO and BRVO in 1 study. Another limitation of this study is that the analysis of the number of injections was on the basis of data collected from trials with diverse treating regimens and follow-up periods. At last, we were unable to access the data from unpublished results and unextractable papers, so publication bias cannot be completely excluded.
5. Conclusions

The current pooled evidence suggested that both therapies of intravitreal conbercept and intravitreal ranibizumab with or without laser photocoagulation are effective in improving visual function in RVO-ME patients, and indicated that conbercept shows advantages over ranibizumab in terms of CMT and number of injections for the treatment of RVO-ME. In addition, conbercept has the statistically same visual gains and safety as ranibizumab in RVO-ME patients. More long-term follow-up surveys on the efficacy and side effects of these 2 treatment strategies are required.

Author contributions

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