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Anti-vascular endothelial growth factor for diabetic macular oedema: a network meta-analysis (Review)

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# Table of Contents

- **Header** ................................................. 1
- **Abstract** ............................................... 1
- **Plain Language Summary** .......................... 2
- **Summary of Findings for the Main Comparison** 4
- **Background** ............................................. 6
- **Objectives** .............................................. 7
- **Methods** .................................................. 7
- **Results** ................................................. 10
  - Figure 1 .................................................. 11
  - Figure 2 .................................................. 14
  - Figure 3 .................................................. 16
  - Figure 4 .................................................. 17
  - Figure 5 .................................................. 18
  - Figure 6 .................................................. 19
  - Figure 7 .................................................. 20
  - Figure 8 .................................................. 21
  - Figure 9 .................................................. 21
  - Figure 10 .................................................. 22
  - Figure 11 .................................................. 23
  - Figure 12 .................................................. 25
- **Additional Summary of Findings** ............... 25
- **Discussion** ............................................. 30
- **Authors’ Conclusions** ............................... 31
- **Acknowledgements** .................................. 31
- **References** ............................................. 32
- **Characteristics of Studies** ......................... 39
- **Data and Analyses** .................................. 110
- **Additional Tables** ................................ 110
- **Feedback** ............................................... 118
- **What’s New** ............................................ 119
- **History** .................................................. 119
- **Contributions of Authors** ......................... 120
- **Declarations of Interest** ............................ 121
- **Sources of Support** .................................. 121
- **Differences Between Protocol and Review** ..... 121
- **Index Terms** ............................................ 122
Anti-vascular endothelial growth factor for diabetic macular oedema: a network meta-analysis

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ABSTRACT

Background

Diabetic macular oedema (DMO) is a common complication of diabetic retinopathy. Antiangiogenic therapy with anti-vascular endothelial growth factor (anti-VEGF) modalities can reduce oedema and thereby improve vision and prevent further visual loss. These drugs have replaced laser photocoagulation as the standard of care for people with DMO.

Objectives

The 2014 update of this review found high-quality evidence of benefit with antiangiogenic therapy with anti-VEGF modalities, compared to laser photocoagulation, for the treatment of DMO. The objective of this updated review is to compare the effectiveness and safety of the different anti-VEGF drugs in preserving and improving vision and quality of life using network meta-analysis methods.

Search methods

We searched various electronic databases on 26 April 2017.

Selection criteria

We included randomised controlled trials (RCTs) that compared any anti-angiogenic drug with an anti-VEGF mechanism of action versus another anti-VEGF drug, another treatment, sham or no treatment in people with DMO.

Data collection and analysis

We used standard Cochrane methods for pair-wise meta-analysis and we augmented this evidence using network meta-analysis methods. We focused on the relative efficacy and safety of the three most commonly used drugs as interventions of direct interest for practice: aflibercept and ranibizumab, used on-label; and off-label bevacizumab.

We collected data on three efficacy outcomes (gain of 15 or more Early Treatment Diabetic Retinopathy Study (ETDRS) letters; mean change in best-corrected visual acuity (BCVA); mean change in central retinal thickness (CRT)), three safety outcomes (all severe systemic adverse events (SSAEs); all-cause death; arterial thromboembolic events) and quality of life.

We used Stata ‘network’ meta-analysis package for all analyses. We investigated the risk of bias of mixed comparisons based on the variance contribution of each study, having assigned an overall risk of bias to each study.
Main results

Twenty-four studies included 6007 participants with DMO and moderate vision loss, of which two studies randomised 265 eyes of 230 participants and one was a cross-over study on 56 participants (62 eyes) that was treated as a parallel-arm trial. Data were collected on drugs of direct interest from three studies on aflibercept (975 eyes), eight studies on bevacizumab (515 eyes), and 14 studies on ranibizumab (1518 eyes). As treatments of indirect interest or legacy treatment we included three studies on pegaptanib (541 eyes), five studies on ranibizumab plus prompt laser (557 eyes), one study on ranibizumab plus deferred laser (188 eyes), 13 studies on laser photocoagulation (936 eyes) and six studies on sham treatment (793 eyes).

Aflibercept, bevacizumab and ranibizumab were all more effective than laser for improving vision by 3 or more lines after one year (high-certainty evidence). Approximately one in 10 people improve vision with laser, and about three in 10 people improve with anti-VEGF treatment: risk ratio (RR) versus laser 3.66 (95% confidence interval (CI) 2.79 to 4.79) for aflibercept; RR 2.47 (95% CI 1.81 to 3.37) for bevacizumab; RR 2.76 (95% CI 2.12 to 3.59) for ranibizumab. On average there was no change in visual acuity (VA) with laser after one year, compared with a gain of 1 or 2 lines with anti-VEGF treatment; laser versus aflibercept mean difference (MD) −0.20 (95% CI −0.22 to −0.17) logMAR; versus bevacizumab MD −0.12 (95% CI −0.15 to −0.09) logMAR; versus ranibizumab MD −0.12 (95% CI −0.14 to −0.10) logMAR. The certainty of the evidence was high for the comparison of aflibercept and ranibizumab with laser and moderate for bevacizumab comparison with laser due to inconsistency between the indirect and direct evidence.

People receiving ranibizumab were less likely to gain 3 or more lines of VA at one year compared with aflibercept: RR 0.75 (95% CI 0.60 to 0.94), moderate-certainty evidence. For every 1000 people treated with aflibercept, 92 fewer would gain 3 or more lines of VA at one year if treated with ranibizumab (22 to 148 fewer). On average people receiving ranibizumab had worse VA at one year (MD 0.08 logMAR units, 95% CI 0.05 to 0.11), moderate-certainty evidence; and higher CRT (MD 39 µm, 95% CI 2 µm to 76 µm, low-certainty evidence). Ranibizumab and bevacizumab were comparable with respect to aflibercept and did not differ in terms of VA: RR of gain of 3 or more lines of VA at one year 1.11 (95% CI 0.87 to 1.43), moderate-certainty evidence, and difference in change in VA was 0.00 (95% CI −0.02 to 0.03) logMAR, moderate-certainty evidence. CRT reduction favoured ranibizumab by −29 µm (95% CI −58 µm to −1 µm, low-certainty evidence). There was no evidence of overall statistical inconsistency in our analyses.

The previous version of this review found moderate-certainty evidence of good safety of antiangiogenic drugs versus control. This update used data at the longest available follow-up (one or two years) and found that aflibercept, ranibizumab and bevacizumab do not differ regarding systemic serious adverse events (SSAEs) (moderate- or high-certainty evidence). However, risk of bias was variable, loop inconsistency could be found and estimates were not precise enough on relative safety regarding less frequent events such as arterial thromboembolic events or death (low- or very low-certainty evidence).

Two-year data were available and reported in only four RCTs in this review. Most industry-sponsored studies were open-label after one year. One large publicly-funded study compared the three drugs at two years and found no difference.

Authors’ conclusions

Anti-VEGF drugs are effective at improving vision in people with DMO with three to four in every 10 people likely to experience an improvement of 3 or more lines VA at one year. There is moderate-certainty evidence that aflibercept confers some advantage over ranibizumab and bevacizumab in people with DMO at one year in visual and anatomic terms. Relative effects among anti-VEGF drugs at two years are less well known, since most studies were short term. Evidence from RCTs may not apply to real-world practice, where people in need of antiangiogenic treatment are often under-treated and under-monitored.

We found no signals of differences in overall safety between the three antiangiogenic drugs that are currently available to treat DMO, but our estimates are imprecise for cardiovascular events and death.

Plain Language Summary

Anti-vascular endothelial growth factor (anti-VEGF) drugs for diabetic macular oedema

What is the aim of this review?

The aim of this Cochrane Review was to find out which is the best type of anti-VEGF drug for diabetic macular oedema (DMO). Cochrane researchers collected and analysed all relevant studies to answer this question and found 24 studies.

Key messages

Anti-vascular endothelial growth factor for diabetic macular oedema: a network meta-analysis (Review)

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Anti-VEGF drugs given by injection into the eye improve vision in people with diabetic macular oedema as compared to no average improvement with laser photocoagulation. One of these drugs, aflibercept, probably works slightly better after one year. There did not appear to be important harms from any of these drugs.

**What was studied in the review?**

The light-sensitive tissue at the back of the eye is known as the retina. The central area of the retina is called the macula. People with diabetes can develop problems in the retina, known as retinopathy. Some people with diabetic retinopathy can also develop oedema (swelling or thickening) at the macula. DMO is a common complication of diabetic retinopathy and can lead to visual loss.

One type of treatment for DMO is anti-VEGF. This drug is given by means of an injection into the eye. It can reduce the swelling at the back of the eye and prevent visual loss. There are three main types of anti-VEGF drugs in use: aflibercept (Eyelea™), bevacizumab (Avastin) and ranibizumab (Lucentis™). Only aflibercept and ranibizumab have received marketing authorisation for the treatment of DMO. All three drugs are used to prevent visual loss and improve vision. They do this by slowing down the growth of new blood vessels and thereby reducing the swelling at the back of the eye. They may have adverse effects, particularly related to effects on blood vessels in the rest of the body. These effects may include stroke and heart attack.

**What are the main results of the review?**

Cochrane researchers found 24 relevant studies. Fourteen of these studies were industry-sponsored studies from USA, Europe or Asia. Ten studies were independent of industry funding and were from USA, Europe, Middle East and South America.

These studies investigated ranibizumab, bevacizumab and aflibercept. These anti-VEGF drugs were compared with no treatment, placebo treatment, laser treatment, or each other. The drugs were given every month, every two months, as needed or ‘treat and extend’, which means that the time period between treatments is extended if the condition has stabilised. Decisions about re-treating were based on visual acuity or by looking at the back of the eye.

The review reveals the following results.

- All three anti-VEGF drugs prevent visual loss and improve vision in people with DMO (high-certainty evidence).
- People receiving ranibizumab were probably slightly less likely to improve vision compared with aflibercept at one year after the start of treatment (moderate-certainty evidence). Approximately three in 10 people improve vision by 3 or more lines with ranibizumab and one in 10 additional people can achieve this with aflibercept.
- People receiving ranibizumab and bevacizumab probably have a similar visual outcome at one year after the start of treatment (moderate-certainty evidence).
- Aflibercept, ranibizumab and bevacizumab are similar for common and serious systemic harms (such as any disease leading to hospitalisation, disability or death) (moderate- or high-certainty evidence) but is less certain for arterial thromboembolic events (mainly stroke, myocardial infarction and vascular death) and death of any cause (very low-certainty evidence).

**How up to date is this review?**

Cochrane researchers searched for studies that had been published up to 26 April 2017.
### Summary of Findings for the Main Comparison

#### Antiangiogenic therapy versus control

**Patient or population:** people with diabetic macular oedema  
**Settings:** ophthalmology clinics  
**Interventions:** laser photoagulation, aflibercept, bevacizumab, ranibizumab

<table>
<thead>
<tr>
<th>Outcomes</th>
<th>Assumed risk*</th>
<th>Corresponding risk and relative risk* (95% CI), mixed evidence</th>
<th>Certainty of evidence and reason for downgrading</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gain 3+ lines of visual acuity at 1 year</td>
<td>100 per 1000</td>
<td>366 per 1000 (279 to 479) RR: 3.66 (2.79 to 4.79)</td>
<td></td>
</tr>
<tr>
<td>Visual acuity change at 1 year Measured on the logMAR scale, range −0.3 to 1.3. Higher values represent worse visual acuity.</td>
<td>On average visual acuity improved by 0.01 logMAR units in the laser group between the start of treatment and 1 year (effectively no change)</td>
<td>Average change in visual acuity was −0.20 (−0.22 to −0.17) logMAR units better with aflibercept compared with laser photoagulation</td>
<td>Average change in visual acuity was −0.12 (−0.15 to −0.09) logMAR units better with bevacizumab compared with laser photoagulation</td>
</tr>
<tr>
<td>Central retinal thickness µm (CRT) change at 1 year The aim of treatment is to reduce central retinal thickness so thinner is better.</td>
<td>On average CRT changed by −64 µm in the laser group between the start of treatment and 1 year (became thinner)</td>
<td>Average change in CRT was −114 (−147 to −81) µm more (thinner) with aflibercept compared with laser photoagulation</td>
<td>Average change in CRT was −46 (−78 to −14) µm more (thinner) with bevacizumab compared with laser photoagulation</td>
</tr>
<tr>
<td>Quality of life: NEI-VFQ composite score at 6 to 12 months An improvement by 5 units is clinically significant.</td>
<td>On average the composite score improved by +2 units in the laser group between the start of treatment and 6 to 12 months</td>
<td></td>
<td>Average change in composite score was 5.14 (2.96 to 7.32) with ranibizumab compared with laser photoagulation</td>
</tr>
<tr>
<td>Event</td>
<td>Control</td>
<td>Study</td>
<td>RR (95% CI)</td>
</tr>
<tr>
<td>-----------------------------------------------</td>
<td>---------</td>
<td>-------</td>
<td>-----------------</td>
</tr>
<tr>
<td><strong>All serious systemic adverse events at 1 to 2 years</strong></td>
<td>200 per 1000</td>
<td>196 per 1000 (166 to 232)</td>
<td>0.98 (0.83 to 1.16)</td>
</tr>
<tr>
<td></td>
<td>186 per 1000 (146 to 238)</td>
<td>194 per 1000 (160 to 234)</td>
<td>0.97 (0.80 to 1.17)</td>
</tr>
<tr>
<td><strong>Arterial thromboembolic events at 1 to 2 years</strong></td>
<td>45 per 1000</td>
<td>38 per 1000 (16 to 94)</td>
<td>0.88 (0.37 to 2.13)</td>
</tr>
<tr>
<td></td>
<td>41 per 1000 (15 to 117)</td>
<td>48 per 1000 (23 to 101)</td>
<td>1.09 (0.52 to 2.29)</td>
</tr>
<tr>
<td><strong>Death at 1 to 2 years</strong></td>
<td>20 per 1000 (7 to 61)</td>
<td>20 per 1000 (9 to 114)</td>
<td>1.01 (0.34 to 3.03) <strong>a</strong></td>
</tr>
<tr>
<td></td>
<td>32 per 1000 (9 to 114)</td>
<td>18 per 1000 (8 to 40)</td>
<td>1.61 (0.45 to 5.69) <strong>a</strong></td>
</tr>
</tbody>
</table>

The **assumed risk** in the laser group was estimated as the row sum of the events divided by the row sum of the participants (eyes) for dichotomous variables, and as the (unweighted) median change of visual acuity or central retina thickness.

The **corresponding risk** (and its 95% confidence interval) is based on the assumed risk in the comparison group and the **relative effect** of the intervention (and its 95% CI).

**a** The risk ratio was estimated from mixed (direct and indirect) comparisons.

GRADE Working Group grades of evidence

**High-certainty:** further research is very unlikely to change our confidence in the estimate of effect.

**Moderate-certainty:** further research is likely to have an important impact on our confidence in the estimate of effect and may change the estimate.

**Low-certainty:** further research is very likely to have an important impact on our confidence in the estimate of effect and is likely to change the estimate.

**Very low-certainty:** we are very uncertain about the estimate.
BACKGROUND

Description of the condition

Diabetic retinopathy (DR) is the most frequent and severe ocular complication of diabetes mellitus (DM) and the leading cause of blindness in the working age population in developed countries (Frank 2004; Klein 1984; Tranos 2004).

Diabetic macular oedema (DMO) is the swelling of the retina resulting from the exudation and accumulation of extracellular fluid and proteins in the macula (Ciulla 2003), due to the breakdown of the blood-retina barrier with an increase in vascular permeability (Antcliff 1999). About a third of people with diabetes have DR and one in 10 is affected by DMO (Yau 2012). The prevalence of DMO increases with diabetes duration, haemoglobin A1c, and blood pressure levels and is higher in people with type 1 compared with type 2 diabetes (Yau 2012).

Intraretinal fluid accumulation results in significant reduction in visual acuity that may be reversible in the short term, but prolonged oedema can cause irreversible damage resulting in permanent visual loss. Blurred vision represents the most common clinical symptom of DMO. Other symptoms can include metamorphopsia (distortion of visual image), floaters, changes in contrast sensitivity, photophobia (visual intolerance to light), changes in colour vision and scotomas (a localised defect of the visual field). During the last decades, the clinical gold standard to detect macular oedema has been fundus examination with contact lens, but non-contact lenses can also be used for this purpose with good sensitivity. Optical coherence tomography (OCT) has progressively been used as an objective and reproducible tool to measure retinal thickness and has been suggested to be the new gold standard for diagnosing DMO (Olson 2013; Ontario HTA 2009). The most severe form of DMO is CSMO, which was defined by the Early Treatment Diabetic Retinopathy Study (ETDRS) as: retinal oedema within 500 µm of the centre of the fovea; hard exudates within 500 µm of the centre of the fovea, if associated with adjacent retinal thickening (which may be outside the 500 µm limit); and one disc area of retinal oedema (1500 µm) or larger, any part of which is within one disc diameter of the centre of the fovea (ETDRS 1985). Since its introduction, OCT was found to be in good agreement with the clinical gold standard (slit-lamp examination with a contact lens) for detecting the presence of macular oedema and was found to be potentially more sensitive in cases of mild foveal thickening (Brown 2004). A simple OCT-based classification of DMO is often used as centre-involving or non-centre-involving DMO (Browning 2008).

Description of the intervention

Anti-angiogenic therapy has been believed a standard of care for treatment of DMO and has largely replaced laser photocoagulation (Jampol 2014), than which it was proven to be more effective (Virgili 2014). Anti-vascular endothelial growth factor (anti-VEGF) treatments inhibit VEGF angiogenic activity, binding to VEGF protein and thus preventing its receptor activation or interaction. These drugs were originally hypothesised as an alternative adjunctive treatment for DMO (Cunningham 2005), following evidence that VEGF-A plays a key role in the occurrence of increased vascular permeability in ocular diseases such as DMO (Aiello 2005).

Grid or focal laser photocoagulation could not be used in all patients with DMO; thus, either laser or sham procedures were current practice comparators in initial studies on the efficacy of antiangiogenic drugs for DMO (Macugen 2005; RESOLVE 2010; RESTORE 2011; Soheilian 2007), and only recently have directly comparative RCTs been conducted (DRCRnet 2015).

Safety of intravitreal antiangiogenic therapy is acceptable; endophthalmitis, the major adverse event (< 1/1000 injections) is related to the surgical injection procedure, rather than the drug itself. These drugs were shown not to increase systemic adverse events such as arterial thromboembolic events, but differences between drugs are not well known (Virgili 2014).

Another therapeutic option for DMO treatment is represented by steroids, administered as intravitreal injections or sustained release implants in order to obtain high local concentrations, maximising their anti-inflammatory, angiostatic and anti-permeability effects while minimising systemic toxicity (Ciulla 2004; Haller 2010; Kuppermann 2010). However, intravitreal steroids may cause cataract and ocular hypertension and the visual outcome is dependent on the lens status or the need for cataract surgery after about one year (Haller 2010; Campochiaro 2010). Currently, some investigators think intravitreal steroids are preferred in patients with anti-VEGF resistant and chronic DMO (Hussain 2010), as an alternative to switching between anti-VEGF drugs. This is also consistent with the EU label of the only approved dexamethasone intravitreal implant in Europe: “Ozurdex is indicated for the treatment of adult patients with visual impairment due to diabetic macular oedema (DME) who are pseudophakic or who are considered insufficiently responsive to, or unsuitable for non-corticosteroid therapy” (accessed on EMA on 4 December 2016). For ranibizumab, the EU label prescribes a 0.5 mg dosage, and that “treatment is initiated with one injection per month until maximum visual acuity is achieved and/or there are no signs of disease activity i.e. no change in visual acuity and in other signs and symptoms of the disease under continued treatment. In patients with wet AMD, DME and RVO, initially, three or more consecutive, monthly injections may be needed. Thereafter, monitoring and treatment intervals should be determined by the physician and should be based on disease activity, as assessed by visual acuity and/or anatomical parameters” (accessed on EMA on 4 December 2016). In the USA, ranibizumab "0.3 mg is recommended to be administered by intravitreal injection once a month (approximately 28 days)” (accessed on FDA on 4 December 2016).
Aflibercept has been approved in the USA, as accessed on FDA on 4 December 2016, and “the recommended dose for EYLEA is 2 mg (0.05 mL) administered by intravitreal injection every 4 weeks (monthly) for the first 5 injections followed by 2 mg (0.05 mL) via intravitreal injection once every 8 weeks (2 months)”. The EU label is similar (accessed on EMA on 4 December 2016).

Bevacizumab is widely used off-label although its use has been questioned based on regulatory or safety issues (Banfi 2013), but is still key for treating chorioretinal vascular disease in low- and middle-income countries thanks to its low cost (Stewart 2016).

**How the intervention might work**

VEGF plays a key role in the occurrence of increased vascular permeability in ocular diseases such as DMO (Aiello 2005). Anti-VEGF agents inhibit VEGF angiogenic activity, binding to VEGF protein thus preventing its receptor activation and interaction.

**Why it is important to do this review**

DMO results in a significant burden of low vision and blindness, thus the extent of the existing evidence base for the effectiveness and safety of these agents needs to be assessed and updated. There is a continuing clinical need to establish evidence-based recommendations regarding anti-VEGF agents.

**OBJECTIVES**

The 2014 update of this review found high-quality evidence of benefit with antiangiogenic therapy with anti-VEGF modalities, compared to laser photoagulation, for the treatment of DMO. As was concluded in the previous version (Virgili 2014), the objective of this updated review is to compare the effectiveness and safety of the different anti-VEGF drugs in preserving and improving vision and quality of life using network meta-analysis methods.

**METHODS**

**Criteria for considering studies for this review**

**Types of studies**

We included randomised controlled trials (RCTs).

**Types of participants**

People with DMO for whom anti-VEGF treatment is indicated. We expected to include most of the studies also included in Virgili 2014.

**Types of interventions**

Any antiangiogenic drug with anti-VEGF modalities compared with another drug with anti-VEGF modalities, laser treatment, sham treatment or no treatment. The reasons for selecting treatments of direct and indirect treatment have been discussed in the Description of the intervention section. As explained above, we remark that steroids may be compared with anti-VEGF drugs but this needs a different approach, specifically patient subgroups and timing, and their inclusion could lead to violation of similarity in a review aiming to compare different anti-VEGF drugs such as this.

Regarding drug dose and monitoring/retreatment regimen, in efficacy analyses we included schemes that are either on-label or commonly used in clinical practice, such as the PRN regimen, as presented in the Description of the intervention section. Particularly, both 0.3 mg and 0.5 mg ranibizumab dose are included as available in studies. These two ranibizumab doses were merged into one group in our NMA since studies suggest no difference between them when used monthly (Heier 2016). Regarding aflibercept, we selected the bi-monthly retreatment regimen since this is the approved label in the USA. We used all available data regardless of safety and dose for safety analyses as previously done in Moja 2014.

**Types of outcome measures**

**Primary outcomes**

Best-corrected visual acuity (BCVA) expressed as the proportion of participants with at least 15 ETDRS letters (3 ETDRS lines or 0.3 logMAR) of improvement in BCVA from baseline to 12 months.

**Secondary outcomes**

- Mean change in BCVA from baseline to 12 months, measured using ETDRS charts.
- Mean change in central retinal thickness (CRT), from baseline to 12 months, measured using optical coherence tomography (OCT).
- Mean change in quality of life from baseline to 12 months, measured using a validated instrument.

Measurements at varying lengths of follow-up were pooled at annual intervals, plus or minus six months, the primary analysis being that at 12 months. The time point closer to 12 months, or the
latest time point in the window frame in the case of symmetry, was chosen where multiple time points were available.

**Adverse events**
The following adverse events were considered.
- All-cause mortality.
- Arterial thromboembolic events (ATC 1994).
- Systemic serious adverse events (SSAEs).

Adverse events were analysed at the longest available follow-up time (Moja 2014).

**Search methods for identification of studies**

**Electronic searches**
The Cochrane Eyes and Vision Information Specialist conducted systematic searches in the following databases for randomised controlled trials and controlled clinical trials. There were no language or publication year restrictions. The date of the search was 26 April 2017.
- Cochrane Central Register of Controlled Trials (CENTRAL; 2017, Issue 3) (which contains the Cochrane Eyes and Vision Trials Register) in the Cochrane Library (searched 26 April 2017) (Appendix 1);
- MEDLINE Ovid (1946 to 26 April 2017) (Appendix 2);
- Embase Ovid (1980 to 26 April 2017) (Appendix 3);
- LILACS (Latin American and Caribbean Health Science Information database (1982 to 26 April 2017) (Appendix 4);
- ISRCTN registry (www.isrctn.com/editAdvancedSearch; searched 26 April 2017) (Appendix 5);
- US National Institutes of Health Ongoing Trials Register ClinicalTrials.gov (www.clinicaltrials.gov; searched 26 April 2017) (Appendix 6);

**Searching other resources**

We handsearched the reference lists of the included trials for other possible trials. We accessed the Novartis Clinical Trials database (www.novocrd.com/crdWebApp/cclinicaltrialrepository/public/main.jsp) on 28 May 2014 and checked all trials indexed under the headings: Ophthalmic Disorders and ranibizumab.

**Data collection and analysis**

**Selection of studies**

Two review authors independently selected the studies for inclusion. The titles and abstracts of all reports identified by the electronic searches and handsearching were examined by the review authors. We classified the abstracts as (a) definitely include, (b) unsure and (c) definitely exclude. We obtained and re-assessed full-text copies of those classified as either (a) definitely include or (b) unsure. Having reviewed the full-text copies, we classified the studies as (1) included, (2) awaiting assessment and (3) excluded. Studies identified by both review authors as ‘excluded’ were excluded and documented in the review. Studies identified as ‘included’ were included and assessed for methodological quality. The review authors were unmasked to the report authors, institutions and trial results during this assessment. Disagreements between the two review authors were resolved by a third review author.

**Data extraction and management**

Two review authors independently extracted the data for the primary and secondary outcomes onto paper data extraction forms developed by the Cochrane Eyes and Vision Group. A pilot test of this form was carried out using a small number of studies. We resolved discrepancies by discussion. One review author entered all data into Review Manager 5 (Review Manager 5 2014). The entered data were checked by a second author. In case standard deviations were not available in the publication, and could not be obtained from the authors, these were imputed from standard deviations of other studies with the same comparison.

**Assessment of risk of bias in included studies**

Two review authors independently assessed the included trials for bias according to the methods described in Chapter 8 of the Cochrane Handbook for Systematic Reviews of Interventions (Higgins 2011b). The following parameters were assessed: sequence generation; allocation concealment; masking (blinding) of participants, personnel and outcome assessors; incomplete outcome data; selective outcome reporting. We evaluated these parameters for each outcome measure or class of outcome measure. We classified each parameter as low risk of bias, high risk of bias or unclear. If the information available in the published trial reports was inadequate to assess methodological quality, we contacted the trial authors for clarification. We had planned that if they did not respond within six months we would assess the trial based on the available information. However, in the latest update of this review we assessed the trial had the authors not responded within one month.

We followed Salanti 2014 to assess the risk of bias of mixed evidence (mixed evidence not defined previously).

1. Summary risk of bias for each trial: we considered all domains but gave more importance to allocation concealment and masking of outcome assessor.
2. Summary risk of bias for the mixed evidence: based on the percentage contribution of each direct comparison to each network estimate using the contribution plot (Chaimani 2013). We finally integrated the risk of bias of a given comparison with the assessment of transitivity, or similarity of the characteristics of the studies. We expected the transitivity assumption would hold as long as treatment comparisons were not related to:
   - acute versus chronic DMO, defined using the cut-off of three or more years of duration;
   - average severity of DMO using OCT CRT of 400 micrometres as a cut-off;
   - treatment regimen, such as monthly versus less than monthly and number of injections in the first year;
   - drug dose for ranibizumab, since this is commercially available in two doses (0.3 mg in the USA, 0.5 mg otherwise);
   - whether the trial was industry sponsored.

Assessment of reporting biases
To investigate small-study bias at the network level we employed the comparison-adjusted funnel plot, which is an adaptation of the funnel plot. We subtracted from each study-specific effect size the mean of meta-analysis of the study-specific comparison and plotted it against the study’s standard error (Chaimani 2013).

Data synthesis
Methods for direct treatment comparisons
If there was no substantial statistical heterogeneity, and if there was no clinical heterogeneity between the trials, we combined the results in a meta-analysis using a random-effects model. A fixed-effect model was used if the number of trials was three or less. In the case of substantial statistical heterogeneity (that is I² value more than 50%) or clinical heterogeneity, we combined the results in a meta-analysis using a random-effects model if the individual trial results were all consistent in the direction of the effect (that is the RR or MD and confidence intervals largely fall on one side of the null line); when the individual trial results were inconsistent in the direction of the effect, we did not combine study results but presented a narrative or tabulated summary of each study.

Methods for indirect and mixed comparisons
We performed network meta-analysis using the methodology of the multivariate meta-analysis model where different treatment comparisons are treated as different outcomes (Salanti 2012). For this analysis, we used the ‘network’ suite of commands available in STATA (StataCorp, 2011; Stata Statistical Software: Release 14. College Station, TX) (White 2015).

We presented mixed effects as RRs or MDs against laser photocoagulation as a single comparison. We prepared league tables presenting mixed comparisons in the inferior-left part and direct comparisons in the superior-right part of the table in order to allow for the inspection of both types of evidence. The same information was presented graphically. We also presented the contribution of direct and indirect evidence to mixed evidence using contribution plots (Chaimani 2013).

Assessment of statistical heterogeneity
In standard pairwise meta-analyses, we estimated heterogeneity variances for each direct comparison. We assessed statistically the presence of heterogeneity within each pairwise comparison using the I² statistic (Higgins 2011b). The I² statistic measures the percentage of variability that cannot be attributed to random error. In network meta-analysis, we assumed a common estimate for the heterogeneity variance across the different comparisons. The assessment of statistical heterogeneity in the entire network was based on the magnitude of the heterogeneity variance parameter (τ²) estimated from the network meta-analysis models.

Measurement of treatment effect
Data analysis followed the guidelines set out in Chapter 9 of the Cochrane Handbook for Systematic Reviews of Interventions (Deeks 2011). For dichotomous outcomes, we calculated a summary risk ratio (RR). For continuous outcome, we calculated the mean difference (MD). We planned to calculate a standardised mean difference (SMD) had different scales been used to measure the same continuous outcome.

We did not use ranking measures in this review, since our main interest was to compare only three drugs: aflibercept, bevacizumab and ranibizumab.

Unit of analysis issues
The unit of randomisation was the eye of individual participants. We included one cross-over study comparing ranibizumab and bevacizumab and treated this as a parallel arm study (Wiley 2016), which equals to assume a moderate (0.5) correlation within-person. However, relative drug safety is impossible to assess with a paired design.

We accepted studies presenting systemic adverse events as the unit of analyses, i.e. when an individual suffers from more than one severe adverse event in the study.

Dealing with missing data
Where data were missing due to dropping out of participants, we conducted a primary analysis based on participants with complete data (available case analysis). Following the guidance available in Chapter 16 of the Cochrane Handbook for Systematic Reviews of Interventions (Higgins 2011a), we considered that missing outcome data are missing at random if the reasons for loss to follow-up are documented and judged to be unrelated to outcome in both study arms.
Assessment of statistical inconsistency

Local approaches for evaluating inconsistency
To evaluate the presence of inconsistency locally, we used the node-splitting approach (Dias 2010). We assumed a common heterogeneity estimate within each loop.

Global approaches for evaluating inconsistency
To check the assumption of consistency in the entire network, we used the 'design-by-treatment' model using the 'network' command in STATA (White 2015). This method accounts for different sources of inconsistency that can occur when studies with different designs (two-arm trials versus three-arm trials) give different results as well as disagreement between direct and indirect evidence. Using this approach, we judged the presence of inconsistency from any source in the entire network based on a Chi² test.

'Summary of findings' table and GRADE assessment
We prepared one 'Summary of findings' table for each relevant comparison, including all seven outcomes in a table (GRADEpro 2014). As originally intended, the primary analysis was conducted at 12 months. Relevant comparisons were identified to answer the question of which antiangiogenic drug is most effective among on-label (aflibercept, ranibizumab) and off-label (bevacizumab) drugs that are currently available. Because most of the available evidence is around ranibizumab, we reported on the comparison of aflibercept and bevacizumab with ranibizumab. Analyses conducted at 24 months were presented textually because a network meta-analysis was not feasible.

We graded the certainty of the evidence for mixed estimates as explained above. We started from the premise that RCTs provide high-certainty evidence and downgraded for each GRADE parameter to get an overall certainty for each outcome as high, moderate, low or very low (Higgins 2014; Salanti 2014; Schünemann 2011). We estimated the absolute risk in the control group from the data in the included studies as the raw proportion for dichotomous outcomes and the median value for continuous outcomes.

Sensitivity and subgroup analyses
We had not planned sensitivity analyses but we decided post-hoc to conduct one excluding studies which were assessed as being at overall high or unclear risk of bias. Moreover, we acknowledge that DRCRnet 2015 emphasised that the differences in absolute benefit between aflibercept, bevacizumab and ranibizumab at one year were dependent on baseline visual acuity, but when we considered this post hoc subgroup analysis we did not find enough study data to conduct such meta-regression analyses.

RESULTS

Description of studies
See: Characteristics of included studies; Characteristics of excluded studies; Characteristics of ongoing studies and Characteristics of studies awaiting classification.

The previous version of this review included 18 trials. Update searches run in April 2017 yielded a further 1166 records (Figure 1). After 397 duplicates were removed, the Cochrane Information Specialist screened the remaining 769 records and removed 577 references that were not relevant to the scope of the review. We screened the remaining 192 references and obtained 19 full-text reports for further assessment. We identified eight reports of six new trials for inclusion in the review (DRCRnet 2015, Ishibashi 2014, Lopez-Galvez 2014, REVEAL 2015, Turkoglu 2015, Wiley 2016). A further four trials were deemed eligible but did not provide sufficient data for analysis (Chen 2016; Huang 2016; Jovanovic 2015; Fouda 2017). We have contacted these authors to ask for further information and will assess these studies if we receive additional data. We excluded one study (NCT02985619 (BEVATAAC)) and have identified six new ongoing studies and will assess these for inclusion in the review when data becomes available (NCT02194634;NCT02259088; NCT02348918; NCT02645734; NCT02699450; NCT02712008).
Figure 1. Study flow diagram.
Included studies

We included a total of 24 studies in this updated systematic review and network meta-analysis. BOLT 2010, DA VINCI 2011, Ishibashi 2014, Korobelnik 2014, Macugen 2005, Macugen 2011, READ2 2009, RELATION 2012, RESOLVE 2010, RESPOND 2013, RESTORE 2011, and RISE-RIDE were industry-sponsored, multicentre RCTs conducted in the USA or Europe, whereas REVEAL 2015 was industry-sponsored but conducted in Asia. Ahmadieh 2008, Azad 2012, Ekinici 2014, LUCIDATE 2014, Nepomuceno 2013, Soheilian 2007, and Turkoglu 2015 were independent studies conducted in Brazil, India, Iran, Turkey, and the UK, five of which included bevacizumab. DRCRnet 2010, DRCRnet 2015, Wiley 2016 were publicly-sponsored studies, mainly by the US National Eye Institute, and conducted in the USA or UK. DRCRnet 2015 was the only large parallel-arm study that compared all commercially available drugs (aflibercept, bevacizumab, ranibizumab) and was a large publicly-funded trial comparing aflibercept, bevacizumab and ranibizumab with monthly monitoring and treatment as needed (PRN). Wiley 2016 was a cross-over trial comparing the same three drugs. Lopez-Galvez 2014 was an open-label trial comparing ranibizumab with laser; it was conducted in Spain and results were available only in abstract form.

Only six trials maintained the randomisation scheme at two years’ follow-up (BOLT 2010; DRCRnet 2010; DRCRnet 2015; Macugen 2011; READ2 2009; RISE-RIDE). Two industry-sponsored trials used randomisation up to two years (Macugen 2011; RISE-RIDE), while three others obtained follow-up data but allowed anti-VEGF treatment in the control arm after one year (Korobelnik 2014; RESOLVE 2010; RESTORE 2011). We did not extract data on comparisons of antiangiogenic therapy with triamcinolone and other intravitreal steroids, which were study arms in Ahmadieh 2008, Azad 2012, DRCRnet 2010 and Soheilian 2007, for reasons presented above and also because this comparison is the subject of another Cochrane Review (Grover 2008). Standard deviations of change in CRT were imputed from other studies in REVEAL 2015.

Types of interventions

Eleven studies assessed ranibizumab (DRCRnet 2010; Lopez-Galvez 2014; LUCIDATE 2014; READ2 2009; RELATION 2012; RESOLVE 2010; RESPOND 2013; RESTORE 2011; REVEAL 2015; RISE-RIDE; Turkoglu 2015), six investigated bevacizumab (Ahmadieh 2008; Azad 2012; BOLT 2010; Ekinici 2014; Nepomuceno 2013; Soheilian 2007), two pegaptanib (Macugen 2005; Macugen 2011), and three aflibercept (DA VINCI 2011; and two studies conducted in the USA and Europe using the same protocol, which we will refer to as a single study (Korobelnik 2014). DRCRnet 2015 and Wiley 2016 were the only studies comparing ranibizumab, bevacizumab or aflibercept directly. The drug dose was the same in most studies (0.5 mg ranibizumab, 1.25 mg bevacizumab, 0.3 mg pegaptanib, 2 mg aflibercept) except for RESOLVE 2010 where dose adjustment was allowed for ranibizumab, and also RISE-RIDE, DRCRnet 2015 and Wiley 2016 where 0.3 mg ranibizumab was also delivered.

Anti-VEGF treatment regimens were monthly in RISE-RIDE, in one arm of Korobelnik 2014 and in Wiley 2016. Monthly, bimonthly and ‘as needed’ or pro re nata (PRN) regimens were adopted in four arms of DA VINCI 2011, and we selected PRN for efficacy data extraction because this is current practice with other anti-VEGF drugs. Ahmadieh 2008 was a short-term study which delivered only the first three injections. Most other studies adopted three initial injections followed by various maintenance regimens. Two studies on aflibercept, reported in Korobelnik 2014 (VISTA and VIVID), compared laser photocoagulation with both monthly injections (2q4) and a regimen of five initial monthly injections followed by bimonthly injections (2q8) followed by a ‘treat-and-extend’ regimen in year two.


Types of participants

Trials included participants with DMO diagnosed clinically, and often these trials used OCT for confirming macular centre involvement. Baseline visual acuity of participants was generally between 20/200 and 20/40. Most trials required a three- to six-month interval from previous central or peripheral laser, and a few small studies required that participants had not received previous antiangiogenic treatment.

Types of outcomes

The data structure of our efficacy and safety outcomes can be seen in Table 1 where the sum of cases for each outcome is shown.

Studies awaiting assessment

Several trials were included as ongoing in the previous version of this review. We checked the completion status...
on the study trial register and tried to contact the authors, but were not able to obtain additional information (NCT00387582; NCT00997191 (IBeTA); NCT01445899 (MATISSE); NCT01565148 (IDEAL)). Two Chinese trials (Chen 2016, 72 participants; Huang 2016, 78 participants) compared ranibizumab plus laser or, respectively, ranibizumab to grid laser. These trials provided baseline and final CRT data at six months as well as the proportion with visual improvement, but the improvement cut-off was unclear, as was the measurement tool. Jovanovic 2015 included 72 participants (120 eyes) randomised to either bevacizumab (one or more injections with or without macular laser photocoagulation depending on results after four to six weeks) or macular laser alone to treat DMO. However, results were not provided at desired fixed follow-up times by each randomisation group. Fouda 2017 included 42 participants (70 eyes) randomised to aflibercept or ranibizumab and treated with three initial injections and then PRN. The authors did not find any significant difference between the two drugs in terms of BCVA, but used decimal rather than logMAR visual acuity and we could not use these data in analyses (authors contacted). The authors reported no difference regarding CRT and a smaller but statistically significant number of injections with aflibercept versus ranibizumab.

**Excluded studies**
See ‘Characteristics of excluded studies’ table for the list of exclusions with reasons.

**Risk of bias in included studies**
See ‘Risk of bias in included studies’; Figure 2.
Figure 2. Methodological quality summary: review authors’ judgements about each methodological quality item for each included study.
**Allocation**

Sequence generation was judged at low risk of bias in 12 studies and was unclear in nine (Azad 2012; Ekinci 2014; Lopez-Galvez 2014; Ishibashi 2014; Korobelnik 2014; READ2 2009; RELATION 2012; RESPOND 2013; Turkoglu 2015). Method for allocation concealment was also unclear in these studies, as they were in Nepomuceno 2013. Allocation concealment was judged at high risk of bias in Ishibashi 2014.

**Blinding**

Masking of participants and outcome assessors was obtained in 14 and 12 trials respectively, and was unclear in seven and nine trials respectively. LUCIDATE 2014, READ2 2009 and RESPOND 2013 were unmasked.

**Incomplete outcome data**

Eleven trials were judged at low risk of attrition bias (Azad 2012; BOLT 2010; DAVINCI 2011; DRCRNet 2010; DRCRNet 2015; Korobelnik 2014; LUCIDATE 2014; Macugen 2005; Nepomuceno 2013; RESOLVE 2010; RESTORE 2011); and eight trials were judged at unclear risk of bias in which some participants were missing but reasons for missingness were not fully reported (Ahmadieh 2008; Ishibashi 2014; Macugen 2011; READ2 2009; RISE-RIDE; Soheilian 2007; Turkoglu 2015; Wiley 2016). Five trials were judged at high risk of attrition bias: Ekinci 2014 excluded 15 participants after randomisation due to ocular and systemic complications; Lopez-Galvez 2014 lost about 20% of participants in each arm and did not report the reasons; RELATION 2012, RESPOND 2013 and REVEAL 2015 lost many more participants in the laser arm than in the ranibizumab arms.

**Selective reporting**

Table 1 shows the reporting of all outcomes across 24 trials. Reporting was almost complete for mean VA change at one year (21 studies, 4489 complete cases). Mean CRT change was available in 16 studies (3491 cases). Gain of 3 or more VA lines was reported at one year in 17 studies (4031 cases). SSAEs at one or two years were reported from 18 studies (4229 cases). ATC thromboembolic events were reported in 15 (3718 cases) and death in 17 (4455 cases).

**Other potential sources of bias**

The baseline visual acuity was not balanced in Soheilian 2007; the visual acuity was around 20/100 in the bevacizumab and bevacizumab-triamcinolone arms and 20/70 in the laser arm, suggesting that milder CSMO was included in the laser arm. The trial investigators adjusted for baseline values in the analyses, which also took into account the within-participant correlation (150 eyes of 129 participants, 16% of participants with both eyes in the analyses). However, we could not take within-participant correlation into account when analysing dichotomous visual acuity.

Three studies included both eyes of some participants in analyses: Ahmadieh 2008 14 out of 101 participants; Nepomuceno 2013 15 out of 48 participants; Wiley 2016 6 out of 56 participants. RELATION 2012 was terminated early when ranibizumab was approved for DMO in Germany. Early termination was unlikely to be associated with treatment effect.

**Effects of interventions**

See: Summary of findings for the main comparison

**Antiangiogenic therapy versus control**

Summary of findings 2 Ranibizumab versus aflibercept for diabetic macular oedema;

**Summary of findings 3 Ranibizumab versus bevacizumab for diabetic macular oedema**

**Antiangiogenic drugs versus laser photocoagulation or control: efficacy and safety**

See: Summary of findings for the main comparison presents the evidence on the comparison of each drug with laser photocoagulation (efficacy at one year) or control (laser photocoagulation or sham at the longest available follow-up of one or two years).

**Efficacy at one year**

As found in the previous version of this review based on direct meta-analyses (Virgili 2014), there was high-certainty of evidence of benefit for aflibercept, bevacizumab and ranibizumab compared to laser photocoagulation at one year. Specifically, aflibercept, bevacizumab and ranibizumab were all more effective than laser for improving vision by 3 or more lines after one year, since about one in 10 people improve vision with laser, and about three in 10 people improve with anti-VEGF treatment: risk ratio (RR) versus laser was 3.66 (95% CI 2.79 to 4.79) for aflibercept; 2.47 (95% CI 1.81 to 3.37) for bevacizumab; and 2.76 (95% CI 2.12 to 3.59) for ranibizumab. Regarding change of mean BCVA, on average there was no change with laser after one year, compared with a gain of 1 or 2 lines with anti-VEGF treatment: laser versus aflibercept mean difference (MD) −0.20 (95% CI −0.22 to −0.17) logMAR; versus bevacizumab −0.12 (95% CI −0.15 to −0.09) logMAR; versus ranibizumab −0.12 (95% CI −0.14 to −0.10) logMAR (negative logMAR in favour of anti-VEGF group). The certainty of evidence was moderate for bevacizumab versus laser regarding mean BCVA change due to inconsistency of direct and indirect evidence.
Safety at the longest available follow-up

This network meta-analysis confirms that aflibercept, bevacizumab and ranibizumab do not increase the risk of all SSAEs compared to laser photocoagulation or sham at one year. We considered this evidence of high-certainty. (Summary of findings for the main comparison). Of notice, SSAEs are a generic indicator of harm, mostly including hospitalisation or death for any cause and unrelated to antiangiogenic effect.

Regarding ‘Antiplatelet Trialists Collaboration arterial thromboembolic events’ and all-cause death, no statistically significant difference was found between any anti-VEGF drug and control, but the certainty of the evidence was generally low due to imprecision (large 95% CIs).

Quality of life

Only RESTORE 2011, RESPOND 2013 and Turkoglu 2015 presented quality of life data for ranibizumab versus laser photocoagulation at six to 12 months (3 studies, 412 participants). Ranibizumab improved NEI-VFQ composite score by 5.14 units (95% CI 2.96 to 7.32) compared to laser (Summary of findings for the main comparison). The certainty of the evidence was moderate due to risk of bias issues (RESPOND 2013 was unmasked and Turkoglu 2015 was unclear for most items).

Macugen 2011 obtained QOL data at two years and we did not include these data since pegaptanib was not of direct interest and sham, rather than laser, was the comparator. RISE-RIDE obtained QOL data at two years and was not included since sham, rather than laser, was the control group.

Ranibizumab versus aflibercept and bevacizumab

Efficacy at one year

Table 2 presents the number of studies (participants/eyes) in all treatment arms of the network for the efficacy outcomes at one year. Figure 3 presents the corresponding networks’ structure. As seen, more data was available for ranibizumab, alone or combined with laser, with respect to aflibercept and bevacizumab. Figure 4, Figure 5 and Figure 6 present forest plots with effects for each study, estimates from direct pairwise meta-analysis and mixed estimate from the network meta-analysis. Summary of findings 2 and Summary of findings 3 present comparisons of ranibizumab versus aflibercept and bevacizumab.
Figure 4. All direct and mixed comparisons: gain of 3 or more lines of visual acuity at 1 year.
Figure 5. All direct and mixed comparisons: mean change in visual acuity at 1 year
Comparing the available drugs as monotherapy, all efficacy outcomes significantly favoured aflibercept over ranibizumab and bevacizumab (Table 3; Table 4; Table 5). Compared with ranibizumab and bevacizumab, aflibercept increased the chances of gaining 3 or more lines (17 studies, 4031 eyes) by about 30%, since the RR for gain was 0.75 (95% CI 0.60 to 0.94) and 0.68 (95% CI 0.53 to 0.86) versus ranibizumab and bevacizumab, respectively. The corresponding figures for mean BCVA change (21 studies, 2689 eyes) were a difference of 0.08 (95% CI 0.05 to 0.11) logMAR and 0.08 (95% CI 0.05 to 0.11) logMAR and were 38.90 (95% CI 2.27 to 75.52) micron and 68.32 (95% CI 28.69 to 107.96) micron for CRT change (16 studies, 3491 eyes), all favouring aflibercept.

Ranibizumab and bevacizumab did not differ in term of functional outcomes: RR of gain 1.11 (95% CI 0.87 to 1.43) and difference in mean VA change 0.00 (95% CI −0.02 to 0.03) logMAR. However, CRT reduction favoured ranibizumab by −29.4 (95% CI −58.2 to −0.70) micron.

There was no evidence of overall statistical inconsistency in our efficacy analyses (Table 3; Table 4; Table 5). We found evidence of statistical inconsistency in one comparison (bevacizumab versus laser) for mean BCVA change and in the loop connecting ranibizumab, ranibizumab plus prompt laser and laser for mean CRT change, where two direct meta-analyses also showed high heterogeneity in the same loop.

Mean risk of bias was low for mixed and direct comparisons among aflibercept, bevacizumab and ranibizumab for all efficacy outcomes, except for the comparison between bevacizumab and ranibizumab regarding mean BCVA change and mean CRT change, which were judged at unclear risk of bias (Figure 7).
We had not pre-planned any subgroup analyses and were unable to obtain data to carry out post hoc subgroup analyses by baseline BCVA. DCRNet 2015, the only large study comparing the three drugs, found that aflibercept was superior to bevacizumab and ranibizumab for participants with lower vision (69 ETDRS letter or less or approximately 20/50 or 0.4 logMAR or worse), whereas differences between the three drugs were unimportant for participants with better vision.

**Efficacy at two years**

Three publicly funded studies (BOLT 2010; DCRNet 2010; DCRNet 2015) and two industry-sponsored studies (Macugen 2011; RISE-RIDE) provided data at two years. There was only one study for each comparison, making data unsuitable for a network meta-analysis. Only DCRNet 2015 (complete cases: aflibercept n = 201, bevacizumab n = 185, ranibizumab n = 191) compared different antiangiogenic drugs, and found no VA differences between ranibizumab 0.3 mg and aflibercept (gain 3+ VA lines, RR 0.94, 95% CI 0.73 to 1.22; difference in mean VA change 0.01, 95% CI −0.04 to 0.06). Ranibizumab and bevacizumab did not differ in terms of gain of 3 or more VA lines (RR 0.94, 95% CI 0.72 to 1.24) but the difference in mean VA change favoured ranibizumab (mean difference −0.05, 95% CI −0.09 to 0.00), although it was not precisely estimated. Although effects on CRT favoured aflibercept over ranibizumab and ranibizumab over bevacizumab, none was statistically significant: mean difference −22 micron (95% CI −50 to 6 micron) and −23 micron (95% CI −52 to 6 micron) respectively.

We were unable to obtain data allowing subgroup analyses by baseline BCVA. DCRNet 2015 found that such subgroup differences were attenuated at two years.

**Safety at the longest available follow-up**

Table 6 presents the number of studies (participants/eyes) in the network for safety outcomes at the longest available follow-up, and Figure 8 presents the corresponding network structure. Figure 9, Figure 10 and Figure 11 present forest plots for each study as well as their direct meta-analysis and mixed estimates from the network meta-analysis.
Figure 8. Network structure for safety outcomes at 1 year

Figure 9. All direct and mixed comparisons: serious systemic adverse events at the longest available follow-up (1 or 2 years)
Figure 10. All direct and mixed comparisons: arterial thromboembolic events at the longest available follow-up (1 or 2 years)
Two-year data were available and reported in only four RCTs in this review. Most industry-sponsored studies were open-label after one year. Differently from efficacy analyses at one year, our safety analyses included data from RISE-RIDE on ranibizumab with monthly treatments up to two years, as well as data from the monthly treatment arm (2q4) of Korobelnik 2014, which became PRN in the second year. Though no analysis suggested a difference among drugs for any safety outcome, only estimates for SSAEs (18 studies, 4229 eyes) reached sufficient precision to exclude very large differences among drugs. Overall, no difference was detected in mixed evidence estimates for any drug compared to laser or sham. Moreover, RR 95% CI width excluded differences of 20% to 30% or more between aflibercept, bevacizumab and ranibizumab, while estimates for pegaptanib were less precise (Table 7). No overall (P = 0.86) or loop-specific inconsistency was detected.

Fifteen studies (3718 eyes) contributed to this analysis on ‘Antiplatelet Trialists Collaboration arterial thromboembolic events’ (Table 8). No difference was detected in mixed evidence estimates for any drug compared to laser or sham or between drugs, but estimates were very imprecise. No overall inconsistency was detected (P = 0.19), but direct evidence from DRCRnet 2015 showed increased risk for ranibizumab compared to aflibercept (RR 2.26, 95% CI 1.15 to 4.23) which was larger and inconsistent with indirect evidence (P = 0.002), resulting in mixed evidence showing no difference (RR 1.25, 95% CI 0.50 to 3.05).

Seventeen studies (4455 eyes) contributed to the analysis of ‘all-cause mortality’ (Table 9). No difference was detected for direct, indirect and mixed evidence estimates for any drug compared to laser or sham or between drugs, but estimates were imprecise. Mean risk of bias was low for mixed and direct comparisons between aflibercept and ranibizumab and unclear for bevacizumab versus ranibizumab for SSAEs. Regarding ATC arterial thromboembolic events and all-cause death, risk of bias was low for aflibercept versus ranibizumab but it was unclear or high for bevacizumab versus ranibizumab (Figure 7).

**Quality of the evidence**

See above for the discussion of risk of bias of mixed evidence in pairwise comparisons of interest.
Statistical heterogeneity between studies

When a direct meta-analysis was possible, no heterogeneity of effects was found for the following outcomes: gain of 3 or more BCVA lines, mean BCVA change, SSAEs, ATC arterial thrombembolic events, death. As reported above, there was high heterogeneity in the meta-analysis of change in CRT for the comparisons of ranibizumab with laser ($I^2 = 91\%$) and ranibizumab plus deferred laser versus laser ($I^2 = 80\%$), but not in other two meta-analyses in this network.

Estimates of between-study standard deviation $\tau$ in the network meta-analyses suggested little heterogeneity for dichotomous outcomes, except for ATC arterial thrombembolic events when it was moderate ($\tau = 0.51$). Values for BCVA change and CRT change were $8^{-10} logMAR$ and 27 micron, respectively. These values mean that heterogeneity was negligible for VA change, but was compatible with a predictive intervals width increased by at least 100 micron for the CRT change.

Similarity between studies

Table 10 shows baseline characteristics (BCVA, CRT) and the number of injections across study and treatment arms. Overall, most studies included participants with mean BCVA about 20/60 and CRT between 400 and 500 micron, which we believe sufficiently homogeneous. The number of injections was high compared with current practice (seven to 10 in year one), except for few small studies delivering a low number of injections. No heterogeneity was suspected between studies using 0.3 versus 0.5 mg ranibizumab. In the safety analyses, we included two studies with monthly injections - one arm of Korobelnik 2014 for aflibercept and RISE-RIDE for ranibizumab - and, again, no heterogeneity seemed to arise from lower intensity regimens in other studies. Regarding sponsorship, there were fewer industry-sponsored studies on bevacizumab, but these studies were also smaller than other studies, and the impact of such differences cannot be assessed. Finally, we did not consider the OCT model used in each study as a source of heterogeneity in CRT change since this was balanced between the arms of each study.

Selective reporting

Comparison adjusted funnel plots showed some asymmetry for the outcome 'gain of 3 or more VA lines', but no specific direct comparison seemed to be affected (Figure 12). Instead, asymmetric observations to the left of the non-significance area were seen for the outcome 'mean CRT change at one year' regarding the comparison 'ranibizumab versus laser', which also suffered high heterogeneity.
Overall quality of evidence for the main comparisons between drugs

Summary of findings 2 and Summary of findings 3 show summary data as well as the overall quality of evidence for the comparisons of interest in this review, based on the data reported above.

Sensitivity analyses on studies at low risk of bias

We conducted analyses of efficacy outcomes after excluding 10 studies at unclear or high risk of bias. These analyses confirmed the findings with all studies.

- Gain of 3 or more BCVA lines: ranibizumab versus aflibercept RR 0.74 (95% CI 0.59 to 0.93); bevacizumab versus ranibizumab RR 0.89 (95% CI 0.69 to 1.15); no overall or comparison-specific inconsistency.

- Mean change in BCVA: ranibizumab versus aflibercept 0.08 (95% CI 0.05 to 0.12) logMAR; bevacizumab versus ranibizumab 0.0 (95% CI 0.2 to −0.03); no overall inconsistency (P = 0.130), some inconsistency for the comparisons of bevacizumab versus laser (P = 0.046) and ranibizumab versus laser (P = 0.043), same direction of effects.

- Mean change in CRT: ranibizumab versus aflibercept 84 (95% CI 43 to 125) micron; bevacizumab versus ranibizumab 16 (95% CI −35 to 67) micron; overall inconsistency was detected (P = 0.012), which was due to the loop ranibizumab plus prompt laser versus ranibizumab plus deferred laser versus laser (P < 0.001 for all comparisons).

Therefore, sensitivity analyses confirmed the results of main analyses.
### ADDITIONAL SUMMARY OF FINDINGS

**Ranibizumab versus aflibercept for diabetic macular oedema**

**Patient or population:** people with diabetic macular oedema  
**Settings:** ophthalmology clinics  
**Interventions:** aflibercept, ranibizumab

<table>
<thead>
<tr>
<th>Outcomes</th>
<th>Illustrative comparative risks* (95% CI)</th>
<th>Relative effect (95% CI), mixed evidence**</th>
<th>Certainty of the evidence (GRADE)</th>
<th>Reason for downgrading certainty of evidence</th>
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<tr>
<td><strong>Assumed risk</strong></td>
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<td>Aflibercept</td>
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<td>Ranibizumab</td>
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<td><strong>Gain 3+ lines of visual acuity at 1 year</strong></td>
<td>370 per 1000</td>
<td>278 per 1000 (222 to 348)</td>
<td>RR: 0.75 (0.60 to 0.94)</td>
<td>☭✭✭ moderate</td>
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</table>
| **Visual acuity change at 1 year**  
*Measured on the log-MAR scale, range —1.3 to 1.3. Higher values represent worse visual acuity.* | On average visual acuity improved by —0.23 logMAR units in the aflibercept group between the start of treatment and 1 year | Average change in visual acuity was 0.08 (0.05 to 0.11) logMAR units worse with ranibizumab compared with aflibercept | ☭✭✭ moderate | —1 for imprecision as confidence intervals include both clinically important and clinically unimportant effects |
| **Central retinal thickness μm (CRT) change at 1 year**  
*The aim of treatment is to reduce central macular thickness so thinner is better.* | On average CRT changed by —181 μm in the aflibercept group between the start of treatment and 1 year (became thinner) | Average change in CRT was 39 (2 to 76) μm more (thicker) with ranibizumab compared with aflibercept | ☭✭ low | —1 for high heterogeneity in two direct comparisons and large predictive intervals —1 for imprecision |
| **Quality of life at 1 year** | No data available. | | | |

---

*Illustrative comparative risks are illustrative and may not reflect risks in your specific population.  
**Relative effect and certainty of evidence were obtained from the network meta-analysis.  
GRADE: Grading of Recommendations, Assessment, Development, and Evaluation.
<table>
<thead>
<tr>
<th>Adverse Event</th>
<th>Event Rate</th>
<th>Event Rate</th>
<th>RR</th>
<th>GRADE</th>
<th>Notes</th>
</tr>
</thead>
<tbody>
<tr>
<td>All serious systemic adverse events at 1 to 2 years</td>
<td>345 per 1000</td>
<td>338 per 1000</td>
<td>0.98</td>
<td>⊕⊕⊕⊕</td>
<td>high</td>
</tr>
<tr>
<td>Arterial thromboembolic events at 1 to 2 years</td>
<td>60 per 1000</td>
<td>74 per 1000</td>
<td>1.24</td>
<td>⊗</td>
<td>very low</td>
</tr>
<tr>
<td>Death at 1 to 2 years</td>
<td>30 per 1000</td>
<td>35 per 1000</td>
<td>1.16</td>
<td>⊗</td>
<td>very low</td>
</tr>
</tbody>
</table>

The **assumed risk** in the aflibercept group was estimated as the row sum of the events divided by the row sum of the participants (eyes) for dichotomous variables, and as the (unweighted) median change of visual acuity or central retina thickness.

The **corresponding risk** (and its 95% confidence interval) is based on the assumed risk in the comparison group and the **relative effect** of the intervention (and its 95% CI).

**CI:** Confidence interval; **RR:** Risk ratio.

GRADE Working Group grades of evidence

**High-certainty:** further research is very unlikely to change our confidence in the estimate of effect.

**Moderate-certainty:** further research is likely to have an important impact on our confidence in the estimate of effect and may change the estimate.

**Low-certainty:** further research is very likely to have an important impact on our confidence in the estimate of effect and is likely to change the estimate.

**Very low-certainty:** we are very uncertain about the estimate.
**Ranibizumab versus bevacizumab for diabetic macular oedema**

**Patient or population:** people with diabetic macular oedema  
**Settings:** ophthalmology clinics  
**Interventions:** bevacizumab, ranibizumab

<table>
<thead>
<tr>
<th>Outcomes</th>
<th>Illustrative comparative risks* (95% CI)</th>
<th>Relative effect (95% CI), mixed evidence**</th>
<th>Certainty of the evidence (GRADE)</th>
<th>Reason for downgrading certainty of evidence</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Assumed risk</strong></td>
<td><strong>Corresponding risk</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Bevacizumab</td>
<td>Ranibizumab</td>
<td>RR 1.11</td>
<td>(0.87 to 1.43)</td>
<td>⚫⚫⚫</td>
</tr>
<tr>
<td><strong>Gain 3+ lines of visual acuity at 1 year</strong></td>
<td>300 per 1000</td>
<td>333 per 1000 (261 to 429)</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Visual acuity change at 1 year</strong></td>
<td>On average visual acuity improved by −0.19 logMAR units in the bevacizumab group between the start of treatment and 1 year</td>
<td>Average change in visual acuity was 0.00 (−0.02 to 0.03) logMAR units (same) with ranibizumab compared with bevacizumab</td>
<td>⚫⚫⚫</td>
<td>moderate</td>
</tr>
<tr>
<td>Measured on the logMAR scale, range −1.3 to 1.3. Higher values represent worse visual acuity.</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Central retinal thickness (CRT) change at 1 year</strong></td>
<td>On average CRT changed by −98 μm in the bevacizumab group between the start of treatment and 1 year (became thinner)</td>
<td>Average change in CRT was −29 (−58 to −1) μm more (thinner) with ranibizumab compared with bevacizumab</td>
<td>⚫⚫</td>
<td>low</td>
</tr>
<tr>
<td>The aim of treatment is to reduce central macular thickness so thinner is better.</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Quality of life at 1 year</strong></td>
<td>No data available</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>All serious systemic adverse events at 1 to 2 years</strong></td>
<td>240 per 1000</td>
<td>250 per 1000 (202 to 307)</td>
<td>RR 1.04</td>
<td>(0.84 to 1.28)</td>
</tr>
<tr>
<td><strong>Arterial thromboembolic events at 1 to 2 years</strong></td>
<td>60 per 1000</td>
<td>70 per 1000 (26 to 189)</td>
<td>RR 1.17</td>
<td>(0.43 to 3.13)</td>
</tr>
</tbody>
</table>
The assumed risk in the bevacizumab group was estimated as the row sum of the events divided by the row sum of the participants (eyes) for dichotomous variables, and as the (unweighted) median change of visual acuity or central retina thickness.

The corresponding risk (and its 95% confidence interval) is based on the assumed risk in the comparison group and the relative effect of the intervention (and its 95% CI).

** The risk ratio was estimated from mixed (direct and indirect) comparisons.

CI: Confidence interval; RR: Risk ratio.

### GRADE Working Group grades of evidence

- **High-certainty**: further research is very unlikely to change our confidence in the estimate of effect.
- **Moderate-certainty**: further research is likely to have an important impact on our confidence in the estimate of effect and may change the estimate.
- **Low-certainty**: further research is very likely to have an important impact on our confidence in the estimate of effect and is likely to change the estimate.
- **Very low-certainty**: we are very uncertain about the estimate.
**DISCUSSION**

**Summary of main results**

This review and network meta-analysis confirms the findings of the previous version which found high-certainty evidence that antiangiogenic therapy provides benefit over laser treatment in people with DMO at one year and concluded that further studies should compare different drugs. This update found four studies with direct comparisons between drugs and augmented this evidence with indirect comparisons in a network meta-analysis based on 24 included studies. At one year, all efficacy outcomes significantly favoured aflibercept over ranibizumab and bevacizumab. Aflibercept increased the chances of gaining 3 or more BCVA lines by about 30%, and conferred an advantage of between half and 1 BCVA line over the other drugs (moderate-certainty of evidence). Ranibizumab and bevacizumab did not differ in terms of functional outcomes (moderate-certainty of evidence), but ranibizumab was more effective in terms of CRT reduction (low-certainty of evidence). There was no evidence of statistical inconsistency in our analyses, except for the comparison between ranibizumab and bevacizumab for mean BCVA change, but the differences between direct and indirect evidence were clinically irrelevant.

The BCVA difference between aflibercept versus ranibizumab or bevacizumab was largely below the threshold of 1 ETDRS line (five letters or 0.1 logMAR) that was used for non-inferiority in trials on DMO (OZDRY 2015, PLACID 2013) and AMD (CATT 2011), suggesting this difference was not clinically relevant. The previous version of this review found moderate-certainty evidence of good safety of antiangiogenic drugs, including aflibercept, bevacizumab, ranibizumab and pegaptanib, versus control. This update found high-certainty evidence that aflibercept, ranibizumab and bevacizumab do not differ regarding SAEs, excluding RR differences between drugs by more than 25%. However, estimates were imprecise on well-defined hard events such as death or arterial thromboembolic events (very low-certainty evidence).

Two-year data on direct comparisons were available only for one large multicentre publicly funded study showing smaller differences between aflibercept, bevacizumab and ranibizumab as compared to one year (DRCRnet 2015).

**Overall completeness and applicability of evidence**

This review confirms and enhances the findings of DRCRnet 2015; that aflibercept confers some advantage over bevacizumab and ranibizumab at one year. Two-year data were available and reported in only four RCTs in this review. Most industry sponsored studies were open-label after one year. Thus, long-term outcomes have to be inferred from observational trials.

We could not investigate subgroup effects in this review due to lack of subgroup data. DRCRnet 2015 found the relative benefit with aflibercept versus ranibizumab and bevacizumab is larger when visual acuity is lower than about 20/50, and modest above this level. Moreover, we could not investigate the potential effect of differences in dose and regimen. Nonetheless, our review includes studies with a broad range of characteristics, but without major differences in populations. Particularly, 0.3 mg ranibizumab was used in the direct comparison with aflibercept and bevacizumab in DRCRnet 2015; yet indirect evidence, mostly based on 0.5 mg ranibizumab, was consistent. Regarding treatment frequency: the only study on monthly 0.3 mg ranibizumab was RISE-RIDE for indirect evidence, but this could not be included since one-year data were not available. We excluded the monthly aflibercept treatment arms of Korobelnik 2014 for efficacy outcomes at one year since this is not the registered label.

We would like to remark that this evidence is obtained in clinical trials with high treatment and monitoring standards. A pragmatic RCT would be needed to assess the real-world effectiveness of anti-VEGF treatment for DMO, since it could be dependent on the adequacy of monitoring treatment response, which is also sensitive to resource constraints, as found for AMD (Pagliarini 2014). Moreover, evidence on safety from non-randomised, real-world data was not included in our review. As found for AMD, real-world studies suggest that people with DMO may differ from those in RCTs (Ziemssen 2017). However there is more compelling real-world evidence that patients are under-treated and have less favourable outcomes than in RCTs for AMD (Chong 2016) compared to DMO (Jiang 2015; Patrao 2016).

**Quality of the evidence**

The quality of evidence has been presented above with reasons for downgrading shown in Summary of findings 2 and Summary of findings 3. Overall, inconsistency was not an issue in our network meta-analyses. We also think that transitivity and generalisability, or indirectness according to GRADE (Schünemann 2011), were not a problem since studies included a broad range of people with DMO that resembles those in clinical practice. Minor funnel plot asymmetry was detected only for CRT change and did not involve the treatments of direct interest in this review.

The tight monitoring of participants in RCTs differs from clinical practice, where a lower number of intravitreal injections and under-treatment are common. As for age-related macular degeneration, this may overestimate benefit with anti-VEGF treatment. However, effect differences among drugs may be less biased if similar regimens are compared in RCTs, although this remains presumptive.

**Potential biases in the review process**
Bevacizumab is an off-label drug for treating DMO in most countries. Because small RCTs using bevacizumab may have been conducted but not published because no difference was found, we could have missed small unpublished studies.

Agreements and disagreements with other studies or reviews

Although we did not systematically search for other reviews on anti-VEGF treatments for DMO, the previous version of this review, which focused on anti-VEGF drugs effects compared to control, reported on other network meta-analyses which were inconclusive (Ford 2012; MEDCAC 2012; Regnier 2014); and included a much smaller set of studies, as did Korobelnik 2015 more recently. Zhang 2016 conducted a network meta-analysis of 21 studies of anti-VEGF drugs versus any control, including studies on intravitreal steroids, such as dexamethasone and triamcinolone, and one retrospective comparative, non-randomised study (Arevalo 2013). The authors concluded that aflibercept was superior to other drugs at 12 months: they found a difference of about 0.04 logMAR (2 ETDRS letters) between aflibercept and ranibizumab as well as between ranibizumab and bevacizumab, but these did not reach statistical significance, as did differences in retinal thickness. Differently from Zhang 2016, we included a larger number of studies on anti-VEGF drugs, but not intravitreal steroids since their benefit profile, as well as their local and systemic harm profile, only partly overlap with that of anti-VEGF drugs and similarity of studies regarding study design and target population would be less likely achieved. Currently, some investigators think intravitreal steroids are preferred in people with anti-VEGF resistant and chronic DMO (Hussain 2015), as an alternative to switching between anti-VEGF drugs. We suggest that a network meta-analysis including both anti-VEGF drugs and steroids is of interest, but a different approach should be used, specifically regarding heterogeneity of effects by time horizon and participants’ subgroups.

The European Society of Retina Specialists have recently published guidelines on the management of DMO, which cover a broad spectrum of clinical questions ranging from imaging interpretation to diabetes management (EURETINA 2017). These guidelines rely on individual study results, particularly those of DRCRnet 2015 which compared aflibercept, bevacizumab and ranibizumab directly regarding anti-VEGF drug choice. They concluded that “aflibercept is the drug of choice in DME eyes with baseline BCVA below 69 letters, as it shows superiority to bevacizumab over 2 years and over ranibizumab in the first year of treatment” and that “all three medications are equivalent in improving vision in eyes with a baseline BCVA letter score of 69 or more”. Our review does not provide additional evidence regarding the effect of baseline vision on visual outcome, since few studies maintained randomisation beyond one year and subgroup data were not available to conduct a network meta-analysis. Regarding the difference between aflibercept and ranibizumab at one year, EURETINA 2017 stated that “it remains unclear to which extent the slower effect of ranibizumab seen in Protocol T [DRCRnet 2015] compared to aflibercept can be accounted to the lower dose (0.3 mg) of ranibizumab used in this study”. Our review found that the difference between aflibercept and ranibizumab was consistent with indirect evidence based on studies that mostly used ranibizumab 0.5 mg, suggesting no dose effect as previously suggested (Heier 2016).

Authors’ conclusions

Implications for practice

There is moderate-certainty evidence that aflibercept confers some advantage in improving visual function over ranibizumab and bevacizumab in people with DMO at one year. An anatomic benefit was found with ranibizumab over bevacizumab (low-certainty evidence), but there was little difference on functional outcomes (low- and moderate-certainty evidence). Relative effects among anti-VEGF drugs at two years are less well known, since most studies did not maintain randomisation after one year or were short term. A single large publicly-funded trial found no differences in visual outcomes among these drugs at two years. Evidence from RCTs may not apply to real-world practice, where people in need of antiangiogenic treatment are often under-treated and under-monitored.

We found no signals of differences in safety between the three antiangiogenic drugs that are currently available to treat DMO, particularly for a summary outcome measure such as the sum of all SSAEs (high- or moderate-certainty evidence). However, our estimates were imprecise regarding arterial thromboembolic events and all-cause death (very low-certainty evidence).

Implications for research

Further studies should be directed to effectiveness in real-world use and focus on monitoring and treatment regimens. A network meta-analysis including steroids for DMO is needed, which should take into account different harms as well as account for differences in populations (e.g. regarding pseudophakic patients and chronic DMO). A network meta-analysis of the relative safety of different antiangiogenic drugs could be conducted including people with different diseases.

Acknowledgements

Cochrane Eyes and Vision (CEV) created and executed the search strategies. We thank Maria Diener-West, Catey Bunce, Tianjing Li...
and Paolo Lanzetta for their comments on the review and Anupa Shah for her assistance throughout the review process.

Wen Xing and Catey Bunce provided visual acuity data on the BOLT 2010 study.

Dr Oliver Zeitz and Dr Christopher James (Bayer HealthCare) provided additional data regarding DA VINCI 2011.

Meagan Huggins provided clarification regarding the randomisation process in DRCRnet 2010.

Dr Oliver Comyn provided data on LUCIDATE 2014.

Dr Kana Inoue and Dr Norio Watanabe extracted data from Ishibashi 2014.

**REFERENCES**

References to studies included in this review

Ahmadieh 2008 [published data only]


Azad 2012 [published data only]


BOLT 2010 [published data only]


DA VINCI 2011 [published data only]


DRCRnet 2010 [published data only]


Anti-vascular endothelial growth factor for diabetic macular oedema: a network meta-analysis (Review)


Macugen 2011 [published data only]


Nepomuceno 2013 [published data only]

READ2 2009 [published data only]


RELATION 2012 [unpublished data only]

NCT01131585. A 12-month, two-armed, randomized, double-masked, multicenter, phase IIIb study assessing the efficacy and safety of laser photocoagulation as adjunctive to ranibizumab intravitreal injections vs. laser photocoagulation monotherapy in patients with visual impairment due to diabetic macular edema followed by a 12 month follow up period. clinicaltrials.gov/show/NCT01131585 (first received 25 May 2010).

RESOLVE 2010 [published data only]
CRFB002DD2201. A randomized, double-masked, multicenter, phase II study assessing the safety and

Anti-vascular endothelial growth factor for diabetic macular oedema: a network meta-analysis (Review)

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RESPOND 2013 [unpublished data only]

RESTORE 2011 [published data only]


REVEAL 2015 [published data only]

RISE-RIDE [published data only]


Soheilian 2007 [published data only]


Turkoglu 2015 [published data only]

Wiley 2016 [published data only]
References to studies excluded from this review
Ahmadieh 2013  [published data only]

CRFB002DFR08 (LUDIC)  [unpublished data only]

CRFB002DGB14 (RELIGHT)  [unpublished data only]

DRCRnet 2007  [published data only]

DRCRnet 2011  [published data only]

DRCRnet 2012  [published data only]

Faghihi 2008  [published data only]

NCT02985619 (BEVATAAC)  [published data only]
NCT02985619. Bevacizumab or triamcinolone for persistent diabetic macular edema (BEVATAAC) [Randomized trial evaluating bevacizumab or triamcinolone for persistent diabetic macular edema]. clinicaltrials.gov/ct2/show/NCT02985619 (first received 1 December 2016).

Paccola 2008  [published data only]

Solaiman 2010  [published data only]

Zehetner 2013  [published data only]

References to studies awaiting assessment
Chen 2016  [published data only]

Fouda 2017  [published data only]

Huáng 2016  [published data only]

Jovanović 2015  [published data only]
Anti-vascular endothelial growth factor for diabetic macular oedema: a network meta-analysis (Review)

References to ongoing studies

ChiCTR-TRC-12002417  [unpublished data only]

NCT01635790 (BRDME)  [published data only]

NCT02194634  [published data only]

NCT02259088  [published data only]
NCT02259088. A 12-month, randomized, efficacy and safety study of 0.5 mg ranibizumab vs laser in Chinese DME patients [A 12–month, randomized, double–masked, multicenter, laser–controlled phase III study assessing the efficacy and safety of 0.5 mg ranibizumab dosed PRN in subjects with visual impairment due to diabetic macular edema in Chinese patients]. clinicaltrials.gov/ct2/show/NCT02259088 (first received 3 October 2014).

NCT02348918  [published data only]
NCT02348918. A phase 2 randomized, controlled, double–masked, multicenter clinical trial designed to evaluate the safety and exploratory efficacy of Luminate® (ALG-1001) as compared to Avastin® and focal laser photoagulation in the treatment of diabetic macular edema. clinicaltrials.gov/ct2/show/NCT02348918 (first received 12 January 2015).

NCT02645734  [published data only]

NCT02699450  [published data only]

NCT02712008  [published data only]

Additional references

Aiello 2005

Antcliff 1999

Arevalo 2013
Arevalo JF, Lasave AF, Wu L, Diaz-Llopis M, Gallego-Pinzon R, Alezzandrini AA, et al. Intravitreal bevacizumab plus grid laser photoagulation or intravitreal bevacizumab or grid laser photoagulation for diffuse diabetic macular...

ATC 1994

Banfi 2013

Borm 2009

Brown 2004

Browning 2008

Campochiaro 2010

CATT 2011

Chaimani 2013

Chong 2016

Ciulla 2003

Ciulla 2004

Cunningham 2005

Deeks 2011

 Dias 2010

ETDRS 1985

EURETINA 2017

Ford 2012

Frank 2004

Glanville 2006

GRADEpro 2014 [Computer program]

Grover 2008
Anti-vascular endothelial growth factor for diabetic macular oedema: a network meta-analysis (Review)

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Guyatt 2011

Haller 2010

Heier 2016

Higgins 2011a

Higgins 2011b

Higgins 2014

Hussain 2015

Jampol 2014

Jiang 2015

Klein 1984

Korobelnik 2015

Kuppermann 2010

MEDCAC 2012

Moja 2014

Olson 2013

Ontario HTA 2009

OZDRY 2015

Pagliarini 2014

Patrao 2016
PLACID 2013

Regnier 2014

Review Manager 5 2014 [Computer program]

Salanti 2012

Salanti 2014

Schnüemann 2011

Stewart 2016

Tranos 2004

White 2015

Yau 2012

Zhang 2016

Ziemssen 2017

References to other published versions of this review

Parravano 2008

Parravano 2009

Virgili 2012

Virgili 2014

* Indicates the major publication for the study
## Characteristics of included studies  
**[ordered by study ID]**

### Ahmadieh 2008

<table>
<thead>
<tr>
<th>Methods</th>
<th>Parallel group RCT</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>People were randomly allocated to treatment but in bilateral cases eyes were randomly allocated to treatment</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Participants</th>
<th>Country: Iran</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number of people randomised: 101 (115 eyes)</td>
<td></td>
</tr>
<tr>
<td>Average age: 60 years (range 39 to 74)</td>
<td></td>
</tr>
<tr>
<td>Sex: 51% women</td>
<td></td>
</tr>
<tr>
<td>Inclusion criteria:</td>
<td></td>
</tr>
<tr>
<td>- CSMO unresponsive to previous macular laser photocoagulation (with the last session being more than 3 months prior)</td>
<td></td>
</tr>
<tr>
<td>Exclusion criteria:</td>
<td></td>
</tr>
<tr>
<td>- VA ≥ 20/40</td>
<td></td>
</tr>
<tr>
<td>- history of cataract surgery within the past 6 months</td>
<td></td>
</tr>
<tr>
<td>- prior intraocular injection or vitrectomy</td>
<td></td>
</tr>
<tr>
<td>- glaucoma or ocular hypertension</td>
<td></td>
</tr>
<tr>
<td>- PDR with high-risk characteristics</td>
<td></td>
</tr>
<tr>
<td>- vitreous haemorrhage</td>
<td></td>
</tr>
<tr>
<td>- significant media opacity</td>
<td></td>
</tr>
<tr>
<td>- presence of traction on the macula</td>
<td></td>
</tr>
<tr>
<td>- monocular</td>
<td></td>
</tr>
<tr>
<td>- pregnancy</td>
<td></td>
</tr>
<tr>
<td>- serum creatinine level ≥ 3 mg/100ml</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Interventions</th>
<th>Intervention:</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>- bevacizumab (1.25 mg) n = ? (41 eyes)</td>
</tr>
<tr>
<td>Comparator:</td>
<td>- sham injection n = ? (37 eyes)</td>
</tr>
</tbody>
</table>

"Three consecutive injections were performed at 6-week intervals. Injections were done under sterile conditions with topical anaesthesia and insertion of a lid speculum. For the IVB group, 1.25 mg (0.05 cc) bevacizumab (Avastin, made for F. Hoffmann-La Roche Ltd Basel, Switzerland by Genentech Inc., San Francisco, CA, USA) was injected intravitreally with a 30-gauge needle through the superotemporal quadrant." Page 485

"In the control group, a needleless syringe was pressed against the conjunctiva and sclera in each session." Page 485

There was another intervention arm that combined bevacizumab with triamcinolone acetonide, but this is not included in this review (n = 37 eyes)

<table>
<thead>
<tr>
<th>Outcomes</th>
<th>Primary outcome:</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>- change in CRT</td>
</tr>
</tbody>
</table>

"Central macular thickness was defined by the average thickness of a central macularregion 1,000 lm in diameter centered on the patient's foveola." Page 485

Secondary outcomes:
- change in BCVA (logMAR)
Ahmadieh 2008  (Continued)

| • intraocular pressure   |
| • cataract progression  |
| • intraocular inflammation |
| • any serious adverse event |

Follow-up: 18 and 24 weeks

Notes
Date study conducted: November 2005 to September 2006
Funding: not reported
Conflict of interest: “The authors have no proprietary interest in this study.”
Trial registration: NCT00370422

Risk of bias

<table>
<thead>
<tr>
<th>Bias</th>
<th>Authors' judgement</th>
<th>Support for judgement</th>
</tr>
</thead>
<tbody>
<tr>
<td>Random sequence generation (selection bias)</td>
<td>Low risk</td>
<td>&quot;Randomization was performed using a random block permutation method according to a computer-generated randomization list. The block lengths varied randomly. A random allocation sequence was performed by a biostatistician. Details of the series were unknown to the investigators.&quot; Page 485</td>
</tr>
<tr>
<td>Allocation concealment (selection bias)</td>
<td>Low risk</td>
<td>&quot;Randomization was performed using a random block permutation method according to a computer-generated randomization list. The block lengths varied randomly. A random allocation sequence was performed by a biostatistician. Details of the series were unknown to the investigators.&quot; Page 485</td>
</tr>
<tr>
<td>Blinding of participants and personnel (performance bias)</td>
<td>Low risk</td>
<td>&quot;Subjects were masked to the treatment modality. Visual acuity assessment and OCT were performed by optometrists who were masked to the groups.&quot; Page 485</td>
</tr>
<tr>
<td>Blinding of outcome assessment (detection bias)</td>
<td>Low risk</td>
<td>See above</td>
</tr>
<tr>
<td>Incomplete outcome data (attrition bias)</td>
<td>Unclear risk</td>
<td>No incomplete outcome data were reported, but number of participants at 24 weeks' follow-up was not specified</td>
</tr>
<tr>
<td>Selective reporting (reporting bias)</td>
<td>High risk</td>
<td>The study protocol is mentioned. However, dichotomous VA outcomes are not provided</td>
</tr>
</tbody>
</table>
### Ahmadieh 2008  (Continued)

<table>
<thead>
<tr>
<th>Other bias</th>
<th>Low risk</th>
<th>28 eyes of 14 participants (14%) with bilateral CSMO were included in the analysis</th>
</tr>
</thead>
<tbody>
<tr>
<td>Overall risk of bias</td>
<td>Low risk</td>
<td>Low risk of bias for most items</td>
</tr>
</tbody>
</table>

### Azad 2012

<table>
<thead>
<tr>
<th>Methods</th>
<th>Parallel group RCT One eye per person, unclear how eye selected</th>
</tr>
</thead>
</table>
| Participants | Country: India  Number of people randomised: 40 (40 eyes) Average age: 54 years Sex: 42% women Inclusion criteria:  
- diffuse DMO on FFA refractory to at least two prior sessions of macular laser photocoagulation  
- CRT > 250 μm on TD-OCT  
- no evidence of vitreo-retinal traction  
- good metabolic control (HbA1c < 7.0%) Exclusion criteria:  
- history of having received prior intraocular, peribulbar or systemic steroids or prior anti-VEGF therapy  
- uncontrolled diabetes mellitus  
- diabetic nephropathy  
- uncontrolled hypertension  
- history of myocardial infarction, stroke or other thromboembolic episode  
- monocular  
- not available for a follow-up duration of at least 6 months |
| Interventions | Intervention:  
- bevacizumab (1.25 mg) n = 20 (20 eyes)  
Comparator:  
- macular grid augmentation n = 20 (20 eyes)  
“IVB […] injected via pars plana route in the doses mentioned above by a single experienced investigator using full aseptic precautions. Postinjection, all patients were prescribed topical moxifloxacin 0.5% qid for 5 days. Macular grid laser augmentation was performed by a single experienced examiner according to the modified ETDRS protocol with a spot size of 100 μ, pulse duration of 100 ms, and a power of 50-100 mW titrated to produce mild intensity burns in areas showing diffuse leakage on the FFA in a ‘C’ shaped zone between 500 and 3000μ from the foveal center sparing the papilla-macular bundle.” Page 167  
Another intervention arm evaluated triamcinolone acetonide, but is not included in this review (n = 20 eyes) |
| Outcomes | Outcomes:  
- BCVA measured used Snellen chart (mean at follow-up, gain/loss of 3 lines)  
- CRT assessed using OCT |
Azad 2012  (Continued)

- adverse effects (increased IOP, cataract progression, others)
  Primary outcome: not specified
  Follow-up: 1, 3 and 6 months

Notes
Date study conducted: not reported
Funding: not reported
Conflict of interest: not reported
Trial registration: not reported

Risk of bias

<table>
<thead>
<tr>
<th>Bias</th>
<th>Authors' judgement</th>
<th>Support for judgement</th>
</tr>
</thead>
<tbody>
<tr>
<td>Random sequence generation (selection bias)</td>
<td>Unclear risk</td>
<td>Not reported</td>
</tr>
<tr>
<td>Allocation concealment (selection bias)</td>
<td>Unclear risk</td>
<td>Not reported</td>
</tr>
<tr>
<td>Blinding of participants and personnel (performance bias)</td>
<td>Unclear risk</td>
<td>Not reported</td>
</tr>
<tr>
<td>All outcomes</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Blinding of outcome assessment (detection bias)</td>
<td>Unclear risk</td>
<td>Not reported</td>
</tr>
<tr>
<td>All outcomes</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Incomplete outcome data (attrition bias)</td>
<td>Low risk</td>
<td>No loss to follow-up reported</td>
</tr>
<tr>
<td>All outcomes</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Selective reporting (reporting bias)</td>
<td>High risk</td>
<td>VA data and other outcomes incompletely reported</td>
</tr>
<tr>
<td>All outcomes</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Other bias</td>
<td>Low risk</td>
<td>No other bias identified</td>
</tr>
<tr>
<td>Overall risk of bias</td>
<td>Unclear risk</td>
<td>Unclear risk of bias for most items</td>
</tr>
</tbody>
</table>

BOLT 2010

Methods
Parallel group RCT
One eye per person; if both eyes were eligible eye with worse VA was selected

Participants
Country: UK
Number of people randomised: 80 (80 eyes)
Average age: 64 years (range 40 to 86)
Sex: 31% women
Inclusion criteria:
- 18 years or older
- diabetes mellitus
- BCVA in the study eye between 35 and 69 ETDRS letters at 4 m (Snellen
equivalent 6/60 or 6/12)  
- centre-involving CSMO with CRT on OCT of ≥ 270 µm  
- media clarity, pupillary dilation, and subject co-operation sufficient for adequate fundus imaging  
- at least 1 prior macular laser therapy  
- intraocular pressure < 30 mmHg  
- ability to return for regular study visits  
- fellow eye ≥ BCVA 3/60  
- fellow eye received no anti-VEGF treatment within the past 3 months and there was no expectation of such treatment during the study

Exclusion criteria: (for study eye)  
- macular ischaemia (FAZ ≥ 1000 µm GLD or severe perifoveal intercapillary loss on FFA)  
- macular oedema due to a cause other than DMO  
- pre-existing ocular condition that was likely to preclude VA improvement despite resolution of macular oedema  
- ocular condition that may affect macular oedema or alter VA during the course of the study, any treatment for DMO in the preceding 3 months  
- PRP within 3 months of enrolment or anticipated 6 months thereafter  
- PDR except for tufts of new vessels elsewhere < 1 disc in area with no vitreous haemorrhage  
- HbA1c > 11.0%  
- medical history of chronic renal failure requiring dialysis or kidney transplantation  
- BP > 170/100 mmHg  
- any thromboembolic event within 6 months  
- unstable angina, or evidence of active ischaemia on electrocardiogram at time of screening  
- major surgery within 28 days of randomisation or planned during the subsequent 12 months  
- participation in an investigational drug trial within 30 days of randomisation (or any time during the study)  
- systemic anti-VEGF or pro-VEGF treatment within 3 months of enrolment  
- pregnancy, breast feeding, or intention to become pregnant within the study period  
- intraocular surgery within 3 months of randomisation  
- aphakia  
- uncontrolled glaucoma  
- significant external ocular disease

Interventions

<table>
<thead>
<tr>
<th>Intervention:</th>
<th>Comparator:</th>
</tr>
</thead>
<tbody>
<tr>
<td>bevacizumab (1.25 mg) n = 42 (42 eyes)</td>
<td>macular laser therapy (MLT) n = 38 (38 eyes)</td>
</tr>
</tbody>
</table>

"Bevacizumab (1.25 mg in 0.05 ml) (Avastin; Roche Registration Limited, UK) was prepared by Moorfields Pharmaceuticals (London, UK) as a prefilled syringe containing 0.13 ml. In a designated intravitreal treatment room, under sterile conditions, using topical anesthesia and povidone-iodine 5% into the conjunctival sac and onto the lid margins, and following application of a drape and insertion of a lid speculum, injections were undertaken with a 30-gauge needle through the supra- or infratemporal quadrant, with a drop of ofloxacin placed
in the fornix at the end of the procedure. Patency of the central retinal artery was determined by indirect ophthalmoscopy and VA of hand movements or better. The IOP was checked 30 minutes after the injection, and if the pressure was increased (30 mmHg) appropriate treatment was commenced. After the injection, topical ofloxacin was instilled 4 times per day for 4 days”. Page 1080

"After baseline IVB, patients received 2 further IVB injections (6- and 12-week time points). Subsequent IVB injections were guided by an OCT-based retreatment protocol. In brief, if the thinnest recorded central retinal thickness was less than 270 m at 18 weeks, then treatment was continued only if macular thickness was not "stable." If central retinal thickness was greater than 270 m at 18 weeks and subsequent visits, then IVB injections were administered until a "stable" macular thickness was attained. "Stable macular thickness" was defined as 3 consecutive visits with the central retinal thickness within 20 m of the patient’s thinnest recorded central retinal thickness. Patients could thereby receive a minimum of 3 injections and a maximum of 9 injections in the first 12 months.” Page 1080

"Modified ETDRS MLT comprised 50 m argon laser spot size, laser applied only greater than 500 m from the edge of the FAZ, with focal treatment aiming to cause mild blanching of the retinal pigment epithelium and not darkening/whitening of microaneurysms. Areas of diffuse leakage or nonperfusion were similarly treated in a grid pattern.” Page 1080

<table>
<thead>
<tr>
<th>Outcomes</th>
<th>Primary outcome:</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>• mean change in BCVA (EDTRS letters measured at 4 m)</td>
</tr>
<tr>
<td></td>
<td>Secondary outcomes:</td>
</tr>
<tr>
<td></td>
<td>• mean CRT and mean change in CRT</td>
</tr>
<tr>
<td></td>
<td>• gain and loss of 15 and 10 letters of ETDRS</td>
</tr>
<tr>
<td></td>
<td>• loss of 30 ETDRS letters</td>
</tr>
<tr>
<td></td>
<td>• retinopathy severity (ETDRS grading)</td>
</tr>
<tr>
<td></td>
<td>• safety</td>
</tr>
<tr>
<td></td>
<td>○ GLD of the FAZ</td>
</tr>
<tr>
<td></td>
<td>○ area of the FAZ</td>
</tr>
<tr>
<td></td>
<td>○ Retinal Nerve Fibre Layer thickness</td>
</tr>
<tr>
<td></td>
<td>○ other ocular side effects</td>
</tr>
<tr>
<td></td>
<td>○ systemic side effects, including thromboembolic events, BP, and ECG findings</td>
</tr>
<tr>
<td>Follow-up:</td>
<td>12 and 24 months</td>
</tr>
</tbody>
</table>

| Notes | Date study conducted: May 2007 to August 2009 |
|       | Funding: “Supported by grants from Moorfields Special Trustees and the National Institute for Health Research UK to the Biomedical Research Center for Ophthalmology based at Moorfields Eye Hospital NHS Foundation Trust and UCL Institute of Ophthalmology.” |
|       | Conflict of interest: “The author(s) have no proprietary or commercial interest in any materials discussed in this article” |
|       | Trial registration: eudract.ema.europa.eu Identifier: 2007-000847-89 |

<p>| Risk of bias |</p>
<table>
<thead>
<tr>
<th>Bias</th>
<th>Authors’ judgement</th>
<th>Support for judgement</th>
</tr>
</thead>
</table>

Anti-vascular endothelial growth factor for diabetic macular oedema: a network meta-analysis (Review)
### Random sequence generation (selection bias)
- **Risk:** Low risk
- **Description:** Patients were randomised into 2 groups by means of an in-house computerized randomization program. The research investigator was not involved in the randomization process. Patients were stratified for BCVA, with the aim being that both groups would have comparable mean baseline BCVAs. Page 1080

### Allocation concealment (selection bias)
- **Risk:** Low risk
- **Description:** The doctor had to phone the Clinical Trial Unit in order to obtain a randomisation from the statistician [personal communication from investigators].

### Blinding of participants and personnel (performance bias)
- **Risk:** Unclear risk
- **Description:** Although the patient and the study physician were not masked to the therapeutic modality, the study optometrist, OCT technician, photographer, graders performing assessment of the FAZ and ETDRS retinopathy grading, and study statistician were all masked to the patient randomization. Page 1080

### Blinding of outcome assessment (detection bias)
- **Risk:** Low risk
- **Description:** See above

### Incomplete outcome data (attrition bias)
- **Risk:** Low risk
- **Description:** Two patients in the laser group did not complete 12 months of follow-up (1 patient moved away, and 1 patient could not be contacted). They were last reviewed at the 32-week time point, with these data being carried forward and an intention-to-treat analysis undertaken. All 42 patients in the IVB group completed the study. Page 1082

### Selective reporting (reporting bias)
- **Risk:** Low risk
- **Description:** We could not find a protocol but primary outcomes were stated in the methods and were those routinely used in the field.

### Other bias
- **Risk:** Low risk
- **Description:** No other bias identified

### Overall risk of bias
- **Risk:** Low risk
- **Description:** Low risk for most items; we considered masking of outcome assessors, though not of participants and physicians, sufficient to ensure unbiased outcome measurement.
### DA VINCI 2011

<table>
<thead>
<tr>
<th>Methods</th>
<th>Parallel group RCT</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>One eye per person, unclear how eye selected</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Participants</th>
<th>Country: USA, Canada and Austria</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number of people randomised:</td>
<td>221 (221 eyes)</td>
</tr>
<tr>
<td>Average age:</td>
<td>64 years (range 40 to 86)</td>
</tr>
<tr>
<td>Sex: 31% women</td>
<td></td>
</tr>
</tbody>
</table>

Inclusion criteria:
- 18 years or older
- diabetes mellitus
- DMO involving the central macula defined as CRT ≥ 250 µm in the central subfield based on Stratus OCT
- BCVA letter score at 4 m of 73-24 (Snellen equivalent: 20/40-20/320) measured by the ETDRS protocol
- women of childbearing potential were included only if they were willing to not become pregnant and to use a reliable form of birth control during the study period

Exclusion criteria:
- history of vitreoretinal surgery
- PRP or macular laser photocoagulation or use of intraocular or periocular corticosteroids or anti-angiogenic drugs within 3 months of screening
- vision decrease due to causes other than DMO
- PDR (unless regressed and currently inactive)
- ocular inflammation
- cataract or other intraocular surgery within 3 months of screening
- laser capsulotomy within 2 months of screening
- aphakia
- spherical equivalent of > −8 dioptres or any concurrent disease that would compromise VA or require medical or surgical intervention during the study period (in either eye)
- active iris neovascularisation
- vitreous haemorrhage
- traction retinal detachment
- preretinal fibrosis involving the macula
- visually significant vitreomacular traction or epiretinal membrane evident biomicroscopically or on OCT
- history of idiopathic or autoimmune uveitis
- structural damage to the center of the macula that is likely to preclude improvement in VA after the resolution of macular oedema
- uncontrolled glaucoma or previous filtration surgery
- infectious blepharitis, keratitis, scleritis, or conjunctivitis
- current treatment for serious systemic infection (systemic)
- uncontrolled diabetes mellitus
- uncontrolled hypertension
- history of cerebral vascular accident or myocardial infarction within 6 months
- renal failure requiring dialysis or renal transplant
- pregnancy or lactation
- history of allergy to fluorescein or povidone iodine
• only 1 functional eye
• ocular condition in the fellow eye with a poorer prognosis than the study eye

### Interventions

**Intervention:**
- VEGF Trap-Eye $n = 177$ (177 eyes)

**Comparator:**
- laser photocoagulation $n = 44$ (44 eyes)

*Patients were randomly assigned in a 1:1:1:1:1 ratio to 1 of 5 treatment regimens in 1 eye only: 0.5 mg VEGF Trap-Eye every 4 weeks (0.5q4); 2 mg VEGF Trap-Eye every 4 weeks (2q4); 2 mg VEGF Trap-Eye for 3 initial monthly doses and then every 8 weeks, (2q8); 2 mg VEGF Trap-Eye for 3 initial monthly doses and then on an as-needed (PRN) basis (2 PRN) ; or macular laser treatment by the modified ETDRS protocol* Page 1820

### Outcomes

**Primary outcome:**
- change in BCVA from baseline to week 24 (ETDRS chart at 4 m)

**Secondary outcomes:**
- retinal thickness assessed by OCT
- safety and tolerability
- change in BCVA from baseline at week 52
- proportion of eyes that gained at least 15 ETDRS letters in BCVA compared with baseline at weeks 24 and 52
- the change in CRT (central subfield on OCT) from baseline to weeks 24 and 52
- number of focal laser treatments given

**Follow-up:** 24 and 52 weeks

### Notes

Date study conducted: December 2008 to June 2009

Funding: "Sponsored by Regeneron Pharmaceuticals, Inc., Tarrytown, New York."


Trial registration:NCT00789477

### Risk of bias

<table>
<thead>
<tr>
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<th>Authors’ judgement</th>
<th>Support for judgement</th>
</tr>
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<tbody>
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<tr>
<td>Table 1: Methodological Quality Assessment Details</td>
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<tr>
<td>--------------------------------------------------</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Random sequence generation (selection bias)</strong></td>
<td>Low risk</td>
<td></td>
</tr>
<tr>
<td><em>&quot;The randomization was handled by an IVRS vendor. The study statistician at REGENERON provided the randomization plan and reviewed and approved the dummy rand table. Study Data Management at REGENERON tested the randomization function extensively along with the Clinical team.&quot;</em></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Allocation concealment (selection bias)</strong></td>
<td>Low risk</td>
<td></td>
</tr>
<tr>
<td><em>&quot;Sites called into IVRS to randomize patients and received the randomization number and drug kit assignment at the completion of the call. The site also received a confirmation email. Neither of these contained the actual randomization assignment. The randomization assignments were kept by the IVRS vendor in a secure, access-controlled database and were delivered to REGENERON by the IVRS vendor at the primary endpoint database lock.&quot;</em></td>
<td></td>
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</tr>
<tr>
<td><strong>Blinding of participants and personnel (performance bias)</strong></td>
<td>Low risk</td>
<td></td>
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<tr>
<td>All outcomes</td>
<td></td>
<td></td>
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<tr>
<td><em>&quot;To maintain participant masking, sham injections were performed on visits when an active dose was not given, and a sham laser was given to the VEGF Trap-Eye groups at week 1. Study drug and sham injections and laser and sham laser treatments were performed by an unmasked physician who had no other role in the study except to assess adverse events (AEs) immediately posttreatment. Sham injections followed the active treatment protocol with the exception that no needle was attached to the syringe, and the syringe hub was gently applied to the sclera to mimic an injection. Sham laser consisted of placing a contact lens on the study eye and positioning the patient in front of the laser machine for the approximate duration of a laser treatment.&quot;</em></td>
<td></td>
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</tr>
<tr>
<td><strong>Blinding of outcome assessment (detection bias)</strong></td>
<td>Unclear risk</td>
<td></td>
</tr>
<tr>
<td>All outcomes</td>
<td></td>
<td></td>
</tr>
<tr>
<td><em>A separate masked physician was assigned to assess adverse events (AEs) and retreatment and rescue criteria and to supervise the masked assessment of efficacy. Every effort was made to ensure that all other study site personnel remained masked to treatment assignment to facilitate an unbiased assessment of efficacy and safety.&quot;</em></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
### DA VINCI 2011 (Continued)

| Incomplete outcome data (attrition bias) | Low risk | “Two randomised patients did not receive treatment and 19 patients discontinued the study after receiving at least 1 treatment for the following reasons: lost to follow-up (6 patients), withdrew consent (6 patients), death (3 patients), treatment failures (2 patients), AE (1 patient), and protocol deviation (1 patient). Discontinuations were evenly distributed among the 5 treatment groups.” Page 1821 Comment: LOCF used |
| Selective reporting (reporting bias) | Low risk | Primary outcome declared and consistent with our review |
| Other bias | Low risk | No other bias identified |
| Overall risk of bias | Low risk | Low risk of bias for most items |

### DRCRnet 2010

#### Methods

Parallel group and within-person RCT
One or two study eyes per person. If both eyes eligible, right eye randomised first and then left eye assigned to “sham plus prompt laser group”. If right eye already assigned to this group then left eye assigned randomly to 1 of the other 3 groups

#### Participants

- **Country:** USA
- **Number of people randomised:** 691 (854 eyes)
- **Average age:** 63 years
- **Sex:** 44% women
- **Inclusion criteria:**
  - 18 years and older
  - diabetes (in study eye)
  - best-corrected Electronic-Early Treatment Diabetic Retinopathy Study (E-ETDRS Visual Acuity Test) VA letter score 78-24 (20/32-20/320)
  - definite retinal thickening due to DMO on clinical examination involving the centre of the macula assessed to be the main cause of visual loss
  - retinal thickness measured on TD-OCT ≥ 250 micron in the central subfield
- **Exclusion criteria:**
  - treatment for DMO within previous 4 months
  - PRP within the previous 4 months or anticipated need for PRP within the next 6 months
  - major ocular surgery within the previous 4 months
  - history of open-angle glaucoma or steroid-induced IOP elevation that required IOP-lowering treatment
  - IOP ≥ 25 mmHg (participant)
### Interventions

**Intervention:**
- ranibizumab (0.5 mg) and laser photocoagulation \(n = ? \) (375 eyes)

**Comparator:**
- sham injection and laser photocoagulation \(n = ? \) (293 eyes)

Ranibizumab group was also randomly allocated to prompt laser photocoagulation (187 eyes) which occurred within 3 to 10 days of the injection and deferred laser photocoagulation (188 eyes) which happened after 24 weeks. All eyes in comparator group were treated within 3 to 10 days of the sham injection.

Complex retreatment algorithm using web-based, real-time data-entry system (page 1066)

There was another intervention arm that combined triamcinolone with prompt laser photocoagulation, but this was not included in this review. \(n = ? \) (186 eyes)

### Outcomes

**Primary outcome:** BCVA and safety at 12 months

**Secondary outcomes:** CRT

Follow-up: every 4 weeks for 12 months. After 12 months, the trial was unmasked and follow-up continued to 3 years

### Notes

Dates participants enrolled: March 2007 to December 2008

Funding: “Supported through a cooperative agreement from the National Eye Institute and the National Institute of Diabetes and Digestive and Kidney Diseases, National Institutes of Health, Department of Health and Human Services EY14231, EY14229, and EY018817. The funding organization (National Institutes of Health) participated in oversight of the conduct of the study and review of the manuscript but not directly in the design or conduct of the study, the collection, management, analysis, or interpretation of the data; or the preparation of the manuscript. Genentech provided the ranibizumab for the study, and Allergan, Inc., provided the triamcinolone for the study. In addition, Genentech and Allergan, Inc., provided funds to the DRCR.net to defray the study's clinical site costs. As described in the DRCR.net Industry Collaboration Guidelines (available at www.drcr.net), the DRCR.net had complete control over the design of the protocol, the ownership of the data, and all editorial content of presentations and publications related to the protocol.”

Conflict of interest: “A complete list of all DRCR.net investigator financial disclosures can be found at www.drcr.net”

Trial registration: NCT00445003

### Risk of bias

<table>
<thead>
<tr>
<th>Bias</th>
<th>Authors’ judgement</th>
<th>Support for judgement</th>
</tr>
</thead>
<tbody>
<tr>
<td>Random sequence generation (selection bias)</td>
<td>Low risk</td>
<td>The randomisation sequence was computer-generated by the DRCR.net co-ordinating centre “...study participants with 1 study eye were assigned randomly on the DRCR.net study web-</td>
</tr>
</tbody>
</table>
### Allocation concealment (selection bias)

<table>
<thead>
<tr>
<th>Risk</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Low</td>
<td>Randomisation assignments were obtained through the DRCR.net study website, therefore no study personnel had access to the list or to the next assignment before it was assigned. “Study participants with 1 study eye were assigned randomly on the DRCR.net study website (using a permuted blocks design stratified by study eye visual acuity)” Page 1065</td>
</tr>
</tbody>
</table>

### Blinding of participants and personnel (performance bias)

<table>
<thead>
<tr>
<th>Risk</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Unclear</td>
<td>“Study participants in the 3 groups receiving laser were masked to treatment assignment through the primary outcome visit, whereas the ranibizumab deferred laser group was not masked.” Page 1065-6</td>
</tr>
</tbody>
</table>

### Blinding of outcome assessment (detection bias)

<table>
<thead>
<tr>
<th>Risk</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Low</td>
<td>“Visual acuity examiners and OCT technicians were masked to treatment group assignment before and at the 1-year primary outcome visit.” Page 1066</td>
</tr>
</tbody>
</table>

### Incomplete outcome data (attrition bias)

<table>
<thead>
<tr>
<th>Risk</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Low</td>
<td>Participants randomised in each group were: 293 laser, 187 ranibizumab + prompt laser, 188 ranibizumab + deferred laser and 186 IVTA + laser. At 1 year complete participants were 274, 171, 178, 176 respectively (91% to 95%) At 2 years complete participants were 211, 136, 139, 142 respectively (72% to 76%) Causes of missing data were balanced across groups</td>
</tr>
</tbody>
</table>

### Selective reporting (reporting bias)

<table>
<thead>
<tr>
<th>Risk</th>
<th>Description</th>
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<td>Low</td>
<td>We could not find a protocol but primary outcomes were stated in the methods and were those routinely used in the field</td>
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</table>

### Other bias

<table>
<thead>
<tr>
<th>Risk</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Low</td>
<td>No other source of bias identified</td>
</tr>
</tbody>
</table>

### Overall risk of bias

<table>
<thead>
<tr>
<th>Risk</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Low</td>
<td>Low risk of bias for most items</td>
</tr>
</tbody>
</table>
### Methods

Paragon group study

“One eye of each participant was randomly assigned in a 1:1:1 ratio to be injected with aflibercept (at a dose of 2.0 mg), bevacizumab (1.25 mg), or ranibizumab (0.3 mg). Randomization was performed at the DRCR.net study website, in permuted blocks and with stratification according to study site and visual acuity in the study eye.”

### Participants

- **Country:** USA
- **Number of people (eyes) randomised:** 660
- **Average age:** 61 years
- **Sex:** not reported
- **Inclusion criteria:**
  - 18 years and older
  - diabetes (in study eye)
  - best-corrected Electronic-Early Treatment Diabetic Retinopathy Study (E-ETDRS Visual Acuity Test) VA letter score 78-24 (20/32-20/320)
  - definite retinal thickening due to DMO on clinical examination involving the centre of the macula assessed to be the main cause of visual loss
  - retinal thickness measured on TD-OCT ≥ 250 micron in the central subfield

- **Exclusion criteria:**
  - treatment for DMO within previous 4 months
  - PRP within the previous 4 months or anticipated need for PRP within the next 6 months
  - major ocular surgery within the previous 4 months
  - history of open-angle glaucoma or steroid-induced IOP elevation that required IOP-lowering treatment
  - IOP ≥ 25 mmHg (participant)
  - systolic BP was 180 mmHg or diastolic BP was 110 mmHg, or if a myocardial infarction, other cardiac event requiring hospitalisation, cerebrovascular accident, transient ischaemic attack, or treatment for acute congestive heart failure occurred within 4 months before randomisation

### Interventions

- **Interventions:**
  - aflibercept 2 mg: 224 eyes
  - bevacizumab 1.25 mg: 218 eyes
  - ranibizumab 0.3 mg: 218 eyes

Randomisation was stratified by site and visual acuity: ≥ 66 letter score/ ≤ 65 letter score

Retreatment algorithm:

“In general, an eye will continue to receive an injection if the eye is improving or worsening on OCT or visual acuity. The first time an eye has not improved or worsened, the eye will receive an injection. If the eye has not improved or worsened for at least 2 consecutive 4-week injections and OCT central subfield thickness is <250µ and visual acuity is 20/20 or better, the injection will be deferred.”

“In general, focal/grid laser will be initiated at or after the 24 week visit if 1) the OCT central subfield thickness is ≥250µ or there is edema that is threatening the fovea and 2) the eye has not improved on OCT or visual acuity from the last two consecutive injections.”
Outcomes
Primary outcome: BCVA and safety at 12 months
Secondary outcomes: CRT
Follow-up: after 12 months the trial was unmasked and follow-up continued to 3 years

Notes
Dates participants enrolled: March 2007 to December 2008
Funding: "Supported through a cooperative agreement from the National Eye Institute and
the National Institute of Diabetes and Digestive and Kidney Diseases, National Institutes of
Health, Department of Health and Human Services EY14231, EY14229, and EY018817.
The funding organization (National Institutes of Health) participated in oversight of the
conduct of the study and review of the manuscript but not directly in the design or conduct of the
study, the collection, management, analysis, or interpretation of the data; or the preparation
of the manuscript. Genentech provided the ranibizumab for the study, and Allergan, Inc.,
provided the triamcinolone for the study. In addition, Genentech and Allergan, Inc., provided
funds to the DRCR.net to defray the study's clinical site costs. As described in the DRCR.net
Industry Collaboration Guidelines (available at www.drcr.net), the DRCR.net had complete
control over the design of the protocol, the ownership of the data, and all editorial content of
presentations and publications related to the protocol."
Conflict of interest: “A complete list of all DRCR.net investigator financial disclosures can
be found at www.drcr.net”
Trial registration: NCT00445003 (Protocol T)

Risk of bias

<table>
<thead>
<tr>
<th>Bias</th>
<th>Authors' judgement</th>
<th>Support for judgement</th>
</tr>
</thead>
</table>
| Random sequence generation (selection bias) | Low risk           | The randomisation sequence was computer-generated by the DRCR.net co-ordinating centre
                                               |                     | “Randomization was performed at the DRCR.net study website, in permuted blocks and with stratification according to study site and visual acuity in the study eye.” Page 3 |
| Allocation concealment (selection bias) | Low risk           | Randomisation assignments were obtained through the DRCR.net study website, therefore no study personnel had access to the list or to the next assignment before it was assigned |
| Blinding of participants and personnel (performance bias) | Low risk           | “Study participants, reading-center graders, and the medical monitor who reviewed all adverse events were unaware of the treatment group assignments. Visual-acuity and OCT technicians were unaware of the treatment-group assignments at the 1-year visit. Investigators and study coordinators were aware of the treatment group assignments.” Page 3 |

<table>
<thead>
<tr>
<th>All outcomes</th>
<th>Low risk</th>
</tr>
</thead>
</table>

Anti-vascular endothelial growth factor for diabetic macular oedema: a network meta-analysis (Review)
Blinding of outcome assessment (detection bias)

| All outcomes | Low risk | “Visual-acuity and OCT technicians were unaware of the treatment-group assignments at the 1-year visit.” Page 3 |

Incomplete outcome data (attrition bias)

| All outcomes | Low risk | “The 2-year visit was completed by 90%, 85%, and 88% of the 660 randomised participants (91%, 90%, and 91% excluding deaths) in the aflibercept, bevacizumab, and ranibizumab groups, respectively (Fig S1, available at www.aaojournal.org). There were no substantial differences identified in the baseline characteristics of those who completed and those who did not complete the 2-year visit (Table S1, available at www.aaojournal.org).” |

Selective reporting (reporting bias)


Other bias

| Low risk | No other source of bias identified |

Overall risk of bias

| Low risk | Low risk of bias for most items |

**Ekinci 2014**

**Methods**

| Parallel group RCT |
| One eye per person, unclear how eye selected |

**Participants**

| Country: Turkey |
| Number of people randomised unclear: 100 (100 eyes) completed follow-up |
| Average age: 67 years (range 50 to 89) |
| Sex: 68% women |

Inclusion criteria:

- clinically significant DMO (CRT > 300 mm), as found through FFA and OCT evaluations and dilate fundus examination, after 1-year follow-up period

Exclusion criteria:

- participants who received intravitreal treatment at another centre
- additional diseases that might have an effect on sight (age related macular degeneration, uveitis, occlusion on the vein root or branch, hereditary macular diseases)
- PRP grid or focal laser photocoagulation application or intraocular surgery within 6 months
- participants with acute ocular infection, stroke, myocardial infarction, uncontrolled hypertension, pregnancy, renal failure and cataract formation during the follow-up period were excluded from the study
Interventions

<table>
<thead>
<tr>
<th>Interventions</th>
<th>Comparator</th>
</tr>
</thead>
<tbody>
<tr>
<td>Intervention: bevacizumab (1.25 mg) n = 50 (50 eyes)</td>
<td>Comparator: ranibizumab (0.05 mg) n = 50 (50 eyes)</td>
</tr>
</tbody>
</table>

"Topical anesthetic drops were instilled, and a drape application and blepharostat attachment were applied. Afterward, fornix lavage was applied using diluted povidone iodine. For Group 1, 1.25 mg (0.05 ml) of bevacizumab was injected into the eye that needed treatment, using a 30 gauge needle; for Group 2, 0.05 mg (0.05 cc) of ranibizumab was injected into the vitreous humor through the lower temporal quadrant, 3.5-4 mm behind the limbus. After the treatment, all patients were treated with topical antibiotics four-times a day for 1 week."

Outcomes

<table>
<thead>
<tr>
<th>Outcomes</th>
</tr>
</thead>
<tbody>
<tr>
<td>BCVA using the Snellen chart</td>
</tr>
<tr>
<td>CRT assessed with OCT</td>
</tr>
<tr>
<td>IOP assessed with applanation tonometry</td>
</tr>
</tbody>
</table>

Primary outcome not specified

Follow-up: monthly intervals after treatment to 12 months

Notes

Dates participants enrolled: 2011 to 2014

Funding: not reported

Conflict of interest: “The authors have no relevant affiliations or financial involvement with any organization or entity with a financial interest in or financial conflict with the subject matter or materials discussed in the manuscript. This includes employment, consultancies, honoraria, stock ownership or options, expert testimony, grants or patents received or pending, or royalties.”

Trial registration: not reported

Risk of bias

<table>
<thead>
<tr>
<th>Bias</th>
<th>Authors’ judgement</th>
<th>Support for judgement</th>
</tr>
</thead>
<tbody>
<tr>
<td>Random sequence generation (selection bias)</td>
<td>Unclear risk</td>
<td>Not reported</td>
</tr>
<tr>
<td>Allocation concealment (selection bias)</td>
<td>Unclear risk</td>
<td>Not reported</td>
</tr>
<tr>
<td>Blinding of participants and personnel (performance bias) All outcomes</td>
<td>Unclear risk</td>
<td>Unclear if participants, care providers or outcome assessors were masked to treatment method</td>
</tr>
</tbody>
</table>
### Ekinci 2014

(Continued)

<table>
<thead>
<tr>
<th>Bias Type</th>
<th>Risk Level</th>
<th>Description</th>
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<tbody>
<tr>
<td>Blinding of outcome assessment (detection bias)</td>
<td>Unclear risk</td>
<td>Unclear if participants, care providers or outcome assessors were masked to treatment method</td>
</tr>
<tr>
<td>Incomplete outcome data (attrition bias)</td>
<td>High risk</td>
<td>Exclusion after randomisation: 15 participants excluded. Patients with acute ocular infection (endophthalmitis after intravitreal injection, n = 3), stroke, myocardial infarction (n = 2), uncontrolled hypertension (n = 4), pregnancy (n = 1), renal failure (n = 1) and cataract formation during follow-up period (n = 4) were excluded from the study.</td>
</tr>
<tr>
<td>Selective reporting (reporting bias)</td>
<td>High risk</td>
<td>We could not find a protocol and our primary outcomes were not reported</td>
</tr>
<tr>
<td>Overall risk of bias</td>
<td>High risk</td>
<td>Most items at high or unclear risk of bias</td>
</tr>
</tbody>
</table>

### Ishibashi 2014

Methods

Allocation: randomised endpoint classification; safety/efficacy study intervention model; parallel assignment

Masking: double-masked

Participants

n = 243; country: Japan

43 recruiting hospitals

Interventions

Drug: pegaptanib sodium (n = 123)

Other: sham injection (n = 120)

Sex: female 113, male 130

Age: (SD) pegaptanib 65.9 (9.0), sham 66.0 (9.2) years

Inclusion criteria:

- Participants with macular oedema including central fovea diagnosed by fluorescein angiography
- Thickening of the retina (≥ 250 µm)
- Corrected VA is 35-68 letters by ETDRS charts

Exclusion criteria:

- Participants who underwent focal/grid laser within 4 month before study started
- Atrophy, scar and fibrosis including the centre of macula
- Underwent any eye surgery within 3 months before study started
- Participants with HbA1c 12.5% ≤ or with symptoms of uncontrolled diabetes

Outcomes

- Number of participants who experience a ≥ 10 letter improvement of VA in ETDRS chart from baseline to week 24
- Change from baseline in VA: double-masked phase (time frame: baseline, weeks 6, 12, 18, and 24); changes in VA were monitored through refraction and BCVA

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Anti-vascular endothelial growth factor for diabetic macular oedema: a network meta-analysis (Review)

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measurements using retro-illuminated, modified Ferris-Bailey ETDRS charts
- Number of participants underwent focal/grid laser, or vitrectomy; double-masked phase (time frame: up to 24 weeks; included focal laser photocoagulation, grid laser photocoagulation, and vitrectomy
- Number of participants who experience a ≥ 10 letter improvement of VA in ETDRS chart from baseline at week 54: open phase (time frame: baseline and week 54); BCVA measurements performed using retro-illuminated, modified Ferris-Bailey ETDRS charts
- Change from baseline in VA: open phase (time frame: baseline, weeks 30, 36, 42, 48 and 54); changes in VA were monitored through refraction and BCVA measurements using retro-illuminated, modified Ferris-Bailey ETDRS charts
- Number of participants who underwent focal/grid laser, or vitrectomy: open phase (time frame: weeks 24 to 54; included focal laser photocoagulation, grid laser photocoagulation, and vitrectomy

<table>
<thead>
<tr>
<th>Bias</th>
<th>Authors’ judgement</th>
<th>Support for judgement</th>
</tr>
</thead>
<tbody>
<tr>
<td>Random sequence generation (selection bias)</td>
<td>Unclear risk</td>
<td>No information</td>
</tr>
<tr>
<td>Allocation concealment (selection bias)</td>
<td>High risk</td>
<td>“Pegaptanib sodium or a syringe of sham was enclosed by aluminium bag, then the aluminium bag was put in a box. These boxes were supplied to each hospitals. These box could maintain masking. However, Pfizer Japan Inc, which requested this drug trial, made the mistake of allowing to open the box for some hospitals during transportation, so it was clear that it was possible not to ensure the masking sufficiently.” In 71 cases, there was the evidence of opening the box of study drugs In 172 cases, there was no evidence of opening the box. In 50 cases, there was evidence of not opening the box, so it was clear that masking is sufficient</td>
</tr>
<tr>
<td>Blinding of participants and personnel (performance bias) All outcomes</td>
<td>Low risk</td>
<td>Treating physician and his/her assistants were not masked and other staff (physician who did not directly give the drug, orthoptist, clinical research coordinator,</td>
</tr>
</tbody>
</table>
**Ishibashi 2014**  *(Continued)*

| Blinding of outcome assessment (detection bias) | Low risk | See above |
| All outcomes |          |          |

| Incomplete outcome data (attrition bias) | Low risk | Lost to follow-up: 6 pegaptanib, 4 sham |
| All outcomes |          |          |

| Selective reporting (reporting bias) | Unclear risk | No information |
| Other bias |          |          |

| Overall risk of bias | High risk | Unclear or high risk for most items |
|                      |          |          |

**Korobelnik 2014**

**Methods**

Parallel group RCT

Eyes: 862 eyes from 862 participants. One eye per participant. *"For patients who met eligibility criteria in both eyes, the eye with the worst BCVA was selected as the study eye. If a patient had DME with similar BCVA in both eyes, the eye with the clearest media was selected as the study eye. If the ocular media of the both eyes were similar in clarity, the patient's non-dominant eye (if identifiable) was selected as the study eye. If neither eye is dominant, the right eye was designated as the study eye."* (Appendix 2)

**Participants**

Country: 54 centres in USA (VISTA study, 446 participants) and 73 centres in Europe, Japan, and Australia (VIVID study, 406 participants)

Number of people randomised: 852 (852 eyes)

Average age: 63 years

Sex: 42% women

Inclusion criteria:

- adults ≥ 18 years with type 1 or 2 diabetes mellitus
- central DMO involvement (defined as retinal thickening involving the 1 mm central (OCT) subfield thickness)
- retinal thickness ≥ 300 µm (assessed by OCT)
- decrease in vision determined to be primarily the result of DME in the study eye
- BCVA ETDRS letter score of 73-24 (20/40-20/320) in the study eye

Exclusion criteria:

- laser photocoagulation (panretinal or macular) in the study eye within 90 days of day 1
- more than 2 previous macular laser treatments in the study eye
- previous use of intraocular or periocular corticosteroids in the study eye within 120 days of day 1
- previous treatment with antiangiogenic drugs in either eye (pegaptanib sodium, bevacizumab, ranibizumab etc.) within 90 days of day 1
Korobelnik 2014  (Continued)

<table>
<thead>
<tr>
<th>Interventions</th>
<th>Interventions:</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>• aflibercept 2q4 n = 290 (290 eyes): aflibercept 2 mg every 4 weeks</td>
</tr>
<tr>
<td>Comparator</td>
<td>• aflibercept 2q8 n = 286 (286 eyes): aflibercept 2 mg monthly for 5 months, then every 8 weeks</td>
</tr>
<tr>
<td></td>
<td>• laser photocoagulation and sham monthly injection = 286 (286 eyes)</td>
</tr>
</tbody>
</table>

"Eyes were randomised in a 1:1:1 ratio to receive either 2 mg IAI every 4 weeks (2q4), 2 mg IAI every 8 weeks after 5 initial monthly doses (from baseline to week 16) with sham injections on non-treatment visits (2q8), or macular laser photocoagulation at baseline and sham injections at every visit (laser control group)."

<table>
<thead>
<tr>
<th>Outcomes</th>
<th>Outcomes:</th>
</tr>
</thead>
<tbody>
<tr>
<td>Primary outcome:</td>
<td>• change in BCVA from baseline to week 52 (ETDRS chart at 4 m)</td>
</tr>
<tr>
<td>Secondary outcomes:</td>
<td>• proportion of eyes that gained at least 10 ETDRS letters in BCVA at week 52 compared with baseline</td>
</tr>
<tr>
<td></td>
<td>• proportion of eyes that gained at least 15 ETDRS letters in BCVA compared with baseline</td>
</tr>
<tr>
<td></td>
<td>• change in CRT (central subfield on OCT) from baseline to week 52</td>
</tr>
<tr>
<td></td>
<td>• proportion of eyes with a 2-step improvement in the ETDRS Diabetic Retinopathy Severity Scale (DRSS) score</td>
</tr>
<tr>
<td></td>
<td>• change from baseline in the National Eye Institute Visual Function Questionnaire-25 (NEI VFQ-25) near activities subscale score</td>
</tr>
<tr>
<td></td>
<td>• change from baseline in the NEI VFQ-25 distance activities subscale score</td>
</tr>
<tr>
<td>Follow-up:</td>
<td>52 weeks</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Notes</th>
<th>Notes:</th>
</tr>
</thead>
<tbody>
<tr>
<td>Date study conducted:</td>
<td>May 2011 to June 2013</td>
</tr>
<tr>
<td>Funding:</td>
<td>&quot;The VISTA and VIVID studies were funded by Regeneron Pharmaceuticals, Inc., Tarrytown, NY and Bayer HealthCare, Berlin, Germany. The sponsors participated in the design and conduct of the study, analysis of the data, and preparation of the manuscript.”</td>
</tr>
<tr>
<td>Conflict of interest:</td>
<td>Assistance with the study design and conduct and data analysis was provided by Karen Chu, MS, and Xiaoping Zhu, PhD, Regeneron Pharmaceuticals, Inc. (VISTA), and Jana Sachsinger, PhD, and Christiane Norenberg, MS, Bayer HealthCare (VIVID). Editorial and administrative assistance to the authors was provided by Hadi Moini, PhD, and S. Balachandra Dass, PhD, Regeneron Pharmaceuticals, Inc. “Other conflicts of interest reported in the paper.</td>
</tr>
<tr>
<td>Trial registration:</td>
<td>VISTA NCT01363440, VIVID NCT01331681</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Risk of bias</th>
<th>Bias</th>
<th>Authors’ judgement</th>
<th>Support for judgement</th>
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</thead>
<tbody>
<tr>
<td>Random sequence generation (selection bias)</td>
<td>Unclear risk</td>
<td>No details available</td>
<td></td>
</tr>
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</table>
### Korobelnik 2014  (Continued)

<table>
<thead>
<tr>
<th>Bias</th>
<th>Risk</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Allocation concealment (selection bias)</td>
<td>Unclear risk</td>
<td>No details available</td>
</tr>
<tr>
<td>Blinding of participants and personnel (performance bias) All outcomes</td>
<td>Low risk</td>
<td>“A masked investigator assessed safety and efficacy and decided on the need for laser re-treatment and additional treatment.”</td>
</tr>
<tr>
<td>Blinding of outcome assessment (detection bias) All outcomes</td>
<td>Unclear risk</td>
<td>“Masked graders at independent central reading centers evaluated OCT images for central retinal thickness (center subfield)”</td>
</tr>
<tr>
<td>Incomplete outcome data (attrition bias) All outcomes</td>
<td>Low risk</td>
<td>About 93% participants completed 52-week follow-up in each arm and causes of loss to follow-up were balanced across arms. Slightly higher loss to follow-up in laser group in VIVID - approx 15% compared to 8% and 11% in aflibercept groups</td>
</tr>
<tr>
<td>Selective reporting (reporting bias)</td>
<td>Unclear risk</td>
<td>Some differences between trial registration and final reports</td>
</tr>
<tr>
<td>Other bias</td>
<td>Low risk</td>
<td>No other bias identified</td>
</tr>
<tr>
<td>Overall risk of bias</td>
<td>Low risk</td>
<td>Though some items could not be fully assessed, we believe randomisation and allocation concealment should be adequate in this multicentre trial aiming at drug registration, as per regulatory requirement</td>
</tr>
</tbody>
</table>

### Lopez-Galvez 2014

<table>
<thead>
<tr>
<th>Method</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Methods</td>
<td>Multicentre, randomised, and open-label controlled trial</td>
</tr>
</tbody>
</table>
| Participants                                                         | Country: Spain  
Number of people randomised: 83 participants (40 ranibizumab, 43 grid laser)  
Average age: 63.5 (9.4) years. Sex: 59.8% M  
Inclusion criteria:  
- ≥18 years old  
- diabetes mellitus type 1/2  
- altered VA due to DMO. The study eye must have had a BCVA = 78-25 letters, and CRT = 250 μm |
| Interventions                                                        | Participants were randomised to intravitreal injection of ranibizumab (0.5 mg) with 3 loading doses and then PRN treatment or to LP (ratio 1:1) |
| Outcomes                                                             | Primary outcome:  
- Differences in mean change in best corrected visual acuity (BCVA) of treatment with ranibizumab 0.5 mg versus laser photocoagulation (LP) over 12 months in |
participants with DMO.
Secondary outcomes:
• % of participants with VA > 73 letters with ranibizumab (0.5 mg) versus laser
• Time and mean change in CRT by OCT with ranibizumab (0.5 mg) versus laser
• Monitoring and registry of all adverse events, serious adverse events, VA, concomitant medications, ophthalmologic exams (including count of fingers and movement of the hands), IOP, vital constants and analytical parameters

Notes
Sponsor: Novartis
Trial Registration: NCT00901186

Risk of bias

<table>
<thead>
<tr>
<th>Bias</th>
<th>Authors’ judgement</th>
<th>Support for judgement</th>
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<tbody>
<tr>
<td>Random sequence generation (selection bias)</td>
<td>Unclear risk</td>
<td>No information (abstract only; authors contacted but no response yet)</td>
</tr>
<tr>
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<td>Unclear risk</td>
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</tr>
<tr>
<td>Blinding of participants and personnel (performance bias)</td>
<td>Unclear risk</td>
<td>No information (abstract only; authors contacted but no response yet)</td>
</tr>
<tr>
<td>All outcomes</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Blinding of outcome assessment (detection bias)</td>
<td>Unclear risk</td>
<td>No information (abstract only; authors contacted but no response yet)</td>
</tr>
<tr>
<td>All outcomes</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Incomplete outcome data (attrition bias)</td>
<td>Unclear risk</td>
<td>9 and 11 participants lost to follow-up for ranibizumab and laser respectively</td>
</tr>
<tr>
<td>All outcomes</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Selective reporting (reporting bias)</td>
<td>Unclear risk</td>
<td>No information (abstract only; authors contacted but no response yet)</td>
</tr>
<tr>
<td>Other bias</td>
<td>Unclear risk</td>
<td>Insufficient information (abstract only; authors contacted but no response yet)</td>
</tr>
<tr>
<td>Overall risk of bias</td>
<td>Unclear risk</td>
<td>Most items at unclear risk</td>
</tr>
</tbody>
</table>
**LUCIDATE 2014**

| Methods | Parallel group RCT  
|---------|------------------  
|         | One eye per person, unclear how eye selected  
|         | “One eye per participant was included to avoid exposure of both eyes to the study drug. If both eyes were eligible, the eye with worse visual acuity became the study eye. Subjects were randomised with 2:1 probability to receive the intervention or standard care (ETDRS macular laser). The randomization list was created using permuted blocks of varying sizes, held by the trial statistician and concealed from the researcher who enrolled, assessed, and allocated treatment to participants.” (Page 961) |

| Participants | Country: UK  
|--------------|------------------  
| Number of people randomised: 37 (37 eyes)  
| Average age: 66 years  
| Sex: 36% women  
| Inclusion criteria:  
| • adult participants with type 1 or 2 diabetes  
| • BCVA of 55-79 ETDRS letters (Snellen equivalent, 20/30-20/80) resulting from centre-involving DMO, with Spectralis OCT (Heidelberg Engineering GmbH, Heidelberg, Germany) central subfield thickness of 300 mm or more in the study eye  
| Exclusion criteria:  
| • uncontrolled glaucoma  
| • aphakia  
| • cataract precluding fundus photography  
| • external ocular infections  
| • previous anti-VEGF or laser treatment in the preceding 3 months in both eyes  
| • angiographic evidence of macular ischaemia defined as FAZ GLD of > 1000 mm or severe perifoveal capillary loss  
| • other causes for macular oedema, for example after cataract surgery  
| • other causes of visual loss in the study eye; other diseases that may affect the course of macular oedema in the study eye  
| • PDR, either active or treated within the previous 3 months  
| • systemic conditions that precluded trial enrolment included HbA1c > 11.0%; past medical history of chronic renal failure requiring either dialysis or kidney transplantation; BP > 170/100 mmHg; an arteriothrombotic event within 6 months before randomisation, including myocardial infarction, acute congestive heart failure or other cardiac event, and stroke or transient ischaemic attack  
| • planned surgery  
| • pregnancy or breastfeeding |

| Interventions | Intervention:  
|---------------|------------------  
| ranibizumab (0.5 mg) n = 25  
| Comparator:  
| • laser photocoagulation n = 12  
| “Subjects were randomised with 2:1 probability to receive the intervention or standard care (ETDRS macular laser).” Page 961. "Intravitreal injections of ranibizumab (Lucentis, 0.5 mg in 0.05 mL solution for injection; Novartis Pharmaceuticals UK Ltd., Frimley, United Kingdom) at baseline, 4 weeks, and 8 weeks then every 4 weeks as required according to predefined retreatment criteria to a maximum of 12 injections. Retreatment occurred if BCVA was reduced by 5 letters or more from maximum acuity or if OCT central subfield thickness was more than 300 mm. Subjects in the laser arm received ETDRS macular laser at baseline guided by fluorescein angiography, OCT, and clinical examination. Laser retreatment occurred |
at 12, 24, and 36 weeks if clinically significant macular edema was still present, in accordance with standard clinical practice at the time; this was guided by the most recent fluorescein angiogram, OCT, and clinical examination results."

### Outcomes

Outcomes:
- change in ETDRS BCVA
- retinal sensitivity
- colour vision
- electrophysiologic parameters
- macular thickness and volume
- change in ETDRS severity grade of diabetic retinopathy from fundus photographs

Follow-up: 48 weeks

### Notes

Date study conducted: November 2010 to July 2011
Sponsor: Moorfields Eye Hospital NHS Foundation Trust
Conflict of interest: "Dr Comyn receives travel support from Novartis. Dr Sivaprasad is a consult for and receives payment for lectures or speaker bureaus and travel support from Novartis, Allergan, and Bayer, and receives payment for development of educational materials from Allergan. Dr Holder is a consultant to Servier. Dr Patel receives grant support from Allergan, Heidelberg United Kingdom, and Topcon United Kingdom and is a consultant to Bayer, Novartis, and Thrombogenics. Dr Hykin is a consultant to and receives grant support from Novartis, Allergan, and Bayer. Drs Comyn, Sivaprasad, Patel, Egang, Egan, Bainbridge, and Hykin have received a proportion of their funding from the Department of Health’s National Institute for Health Research Biomedical Research Centre for Ophthalmology at Moorfields Eye Hospital and University College London, Institute of Ophthalmology. Dr Bainbridge is supported by a National Institute for Health Research Professorship. Supported by an unrestricted research grant from Novartis and the National Institute for Health Research Biomedical Research Centre based at Moorfields Eye Hospital National Health Service Foundation Trust and University College London Institute of Ophthalmology."

Trial registration: NCT01223612

### Risk of bias

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<thead>
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</tr>
<tr>
<td>Allocation concealment (selection bias)</td>
<td>Low risk</td>
<td>See above</td>
</tr>
<tr>
<td>Blinding of participants and personnel (performance bias) All outcomes</td>
<td>High risk</td>
<td>No sham procedure</td>
</tr>
</tbody>
</table>
### Blinding of outcome assessment (detection bias)

<table>
<thead>
<tr>
<th>All outcomes</th>
<th>Unclear risk</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>“The microperimetry and electrophysiologic assessors were masked to the patient treatment arm. Evaluation of OCT scans, fundus photographs and fluorescein angiograms was performed by masked Reading Centre graders. The protocol states that the visual acuity assessors were also masked to the patient treatment arm but due to a protocol deviation they had access to the source notes and were potentially unmasked.”</td>
</tr>
</tbody>
</table>

### Incomplete outcome data (attrition bias)

<table>
<thead>
<tr>
<th>All outcomes</th>
<th>Low risk</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>22/25 (88%) of anti-VEGF group compared to 11/12 (92%) laser group followed up</td>
</tr>
</tbody>
</table>

### Selective reporting (reporting bias)

<table>
<thead>
<tr>
<th>Unclear risk</th>
</tr>
</thead>
</table>

### Other bias

| Low risk                                                                 |
| No other source of bias identified                                          |

### Overall risk of bias

| Unclear risk                                                                 |
| High or unclear risk of bias for nearly half the items                     |

---

### Macugen 2005

#### Methods

<table>
<thead>
<tr>
<th>Parallel group RCT</th>
</tr>
</thead>
<tbody>
<tr>
<td>One eye per person, chosen by participant and physician. In 81% of cases the eye with the worse VA was chosen</td>
</tr>
</tbody>
</table>

#### Participants

<table>
<thead>
<tr>
<th>Country: USA</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number of people randomised: 172 (172 eyes)</td>
</tr>
<tr>
<td>Average age: 62 years (range 27 to 89)</td>
</tr>
<tr>
<td>Sex: 49% women</td>
</tr>
<tr>
<td>Inclusion criteria:</td>
</tr>
<tr>
<td>- 18 years or older</td>
</tr>
<tr>
<td>- diabetes (study eyes)</td>
</tr>
<tr>
<td>- macular oedema involving the centre of the macula demonstrated on OCT with corresponding leakage from microaneurysms, retinal telangiectasis, or both on fluorescein angiography</td>
</tr>
<tr>
<td>- an area of retinal thickening of at least half a disc area involving the central macula as confirmed by graders at an independent fundus photograph and angiogram reading center (University of Wisconsin, Madison, Wisconsin)</td>
</tr>
<tr>
<td>- clear ocular media and adequate pupillary dilation to permit good stereoscopic fundus photographs (participants)</td>
</tr>
<tr>
<td>- BCVA letter scores between 68-25 inclusive (approximate Snellen equivalent, 20/50-20/320) in the study eye and at least 35 (20/100 or better) in the fellow eye</td>
</tr>
<tr>
<td>- IOP ≤ 23 mmHg</td>
</tr>
</tbody>
</table>
Macugen 2005  (Continued)

• assessment by the treating ophthalmologist that focal photocoagulation could be deferred safely for 16 weeks
• an electrocardiogram that demonstrated no abnormalities judged to be clinically relevant and serological test results that suggested no clinically meaningful haematological, liver, or renal abnormalities
• women enrolling in the study were required to be postmenopausal for 12 months before the study, surgically sterile, or not pregnant and on 2 forms of effective contraception

Exclusion criteria:
• history of PRP or focal photocoagulation
• neodymium:yttrium-aluminum-garnet laser or peripheral retinal cryoablation within the previous 6 months
• any abnormality thought likely to confound VA assessments or fundus photography, including cataract; vitreoretinal traction within 1 disc diameter of the fovea confirmed either clinically or on OCT
• vitreous incarceration in a previous wound or incision
• any retinal vein occlusion involving the macula; and atrophy/scarring/fibrosis or hard exudates involving the centre of the macula that would preclude improvement in VA
• a history of any intraocular surgery within the previous 12 months, myopia of ≥ 8 dioptres, axial length of ≥ 25 mm, and the likelihood of requiring either scatter (panretinal) photocoagulation within the ensuing 9 months or cataract surgery within 12 months
• active ocular or periocular infection
• previous therapeutic radiation to the eye, head, or neck
• any treatment with an investigational agent for any condition in the 60 days before enrolment. Known serious allergies to fluorescein dye
• glycosylated haemoglobin (GHb) levels of ≥ 13%
• 3 episodes of severe hypoglycaemia within 3 months of study entry
• 2 episodes of ketoacidosis within 1 year of baseline
• any episode of ketoacidosis within 3 months of baseline
• evidence of severe cardiac disease
• clinically significant peripheral vascular disease (previous surgery, amputation, or symptoms of claudication)
• uncontrolled hypertension (treated systolic BP 155 or diastolic BP 95), or stroke within the preceding 12 months

Interventions

Intervention:
• pegaptanib (0.3 mg, 1 mg, or 3 mg) n = 130 (130 eyes)

Comparator:
• sham injection n = 42 (42 eyes)

"Intravitreous pegaptanib or sham injections were administered at entry, week 6, and week 12, for a minimum of 3 injections. Thereafter, additional injections were administered every 6 weeks at the discretion of investigators if judged indicated, to a maximum of 6 injections up to week 30. [...] Pegaptanib was formulated for intravitreous injection at 0.3 mg/90 µl, 1 mg/90 µl, and 3 mg/90 µl concentrations in preservative-free phosphate-buffered saline (pH 5-7). Pegaptanib was packaged in sterile, single-use, United States Pharmacopeia type 1 graduated glass 1-ml syringes with preattached 27-gauge needles" Page 1748
### Outcomes

Outcomes:
- BCVA (measured using ETDRS chart)
- CRT on OCT
- change in retinal thickness derived by comparing measurements at baseline with those at week 36 or final examination if before week 36
  - focal photocoagulation applied at week 12 or later
  - size of the area of retinal thickness measured by photography
  - macular capillary leakage and cystoid spaces
  - adverse events
  - laboratory test abnormalities

Follow-up: 36 weeks

### Notes

Dates participants enrolled: not reported, study published 2005
Funding: "The study was sponsored by Eyetech Pharmaceuticals, Inc., New York, New York, and Pfizer Inc., New York, New York." Page 1747
Conflict of interest: not reported
Trial registration: NCT00040313

### Risk of bias

<table>
<thead>
<tr>
<th>Bias</th>
<th>Authors' judgement</th>
<th>Support for judgement</th>
</tr>
</thead>
<tbody>
<tr>
<td>Random sequence generation (selection bias)</td>
<td>Low risk</td>
<td>&quot;Patients were allocated [...] by a dynamic minimization procedure using a stochastic treatment allocation algorithm based on the variance method. Randomization was stratified by study site, size of the thickened retina area [...] and baseline VA [...]&quot; Page 1748</td>
</tr>
<tr>
<td>Allocation concealment (selection bias)</td>
<td>Low risk</td>
<td>&quot;An independent fundus photograph and angiogram reading center confirmed eligibility and appropriate retinal thickness classification both for study entry and for randomization and stratification using baseline fluorescein angiography and OCT.&quot; Page 1748</td>
</tr>
<tr>
<td>Blinding of participants and personnel (performance bias) All outcomes</td>
<td>Low risk</td>
<td>&quot;Study subjects receiving sham or study medication were treated identically in all regards, including ocular antisepsis procedures and subconjunctival anesthetic, except that subjects receiving active treatment had pegaptanib injected into the vitreus, whereas those receiving sham had a needleless syringe pressed against the conjunctiva and sclera. The injection procedure prevented subjects from seeing the syringe and needle, to minimize the risk of unmasking. In all but 3 subjects, injection was administered by a staff member and...&quot;</td>
</tr>
</tbody>
</table>

---

Macugen 2005 (Continued)
than the study ophthalmologist responsible for all other aspects of the protocol, to maintain investigator masking." Page 1748

<table>
<thead>
<tr>
<th>Blinding of outcome assessment (detection bias)</th>
<th>Low risk</th>
<th>“Study subjects receiving sham or study medication were treated identically in all regards, including ocular antisepsis procedures and subconjunctival anesthetic, except that subjects receiving active treatment had pegaptanib injected into the vitreous, whereas those receiving sham had a needleless syringe pressed against the conjunctiva and sclera. The injection procedure prevented subjects from seeing the syringe and needle, to minimize the risk of unmasking. In all but 3 subjects, injection was administered by a staff member other than the study ophthalmologist responsible for all other aspects of the protocol, to maintain investigator masking. Visual acuity was determined by a separate VA examiner masked to treatment.” Page 1748</th>
</tr>
</thead>
<tbody>
<tr>
<td>Incomplete outcome data (attrition bias)</td>
<td>Low risk</td>
<td>Nine participants were discontinued from the study before week 36. None in pegaptanib groups 0.3 mg and 1 mg, 3 in pegaptanib 3 mg group (3 mg subgroup: 2 participants by request at weeks 12 and 16 and 1 by other reason at week 1), 6 in sham group (5 participants by request at weeks 6, 11, 18, 30, and 33 and 1 due to death at week 8)</td>
</tr>
<tr>
<td>Selective reporting (reporting bias)</td>
<td>Low risk</td>
<td>The study protocol is available and all (primary and secondary) outcomes that are of interest in the study have been reported in the pre-specified way</td>
</tr>
<tr>
<td>Other bias</td>
<td>Low risk</td>
<td>No other source of bias identified</td>
</tr>
<tr>
<td>Overall risk of bias</td>
<td>Low risk</td>
<td>Low risk of bias for all items</td>
</tr>
</tbody>
</table>
### Methods

Parallel group RCT

One eye per person, unclear how eye selected

### Participants

Country: Australia, Europe, India, North America, and South America

Number of people randomised: 288 (288 eyes)

Average age: 62 years (20 to 83)

Sex: 43% women

Inclusion criteria:

- 18 years or older
- diabetes
- DMO involving the centre of the macula not associated with ischaemia (study eye)
- foveal thickness of ≥ 250 µm (centre point thickness measured on OCT)
- BCVA with a letter score of 65-35 (20/50-20/200 Snellen equivalents)
- IOP ≤ 21 mmHg
- clear ocular media and adequate pupillary dilation to allow good quality stereoscopic fundus photography
- focal or grid laser photocoagulation could be deferred for 18 weeks in the opinion of the treating ophthalmologist

Exclusion criteria:

- yttrium-aluminum-garnet laser, peripheral retinal cryoablation, laser retinopexy for retinal tears, or focal or grid photocoagulation within the prior 16 weeks or scatter PRP 6 months before baseline or likely to be needed within 9 months
- macular ischaemia if a nonperfusion area of > 1 disc area involving the foveal avascular zone (2 quadrants centred around the FAZ)

### Interventions

**Intervention:**

- pegaptanib sodium (0.3 mg) n = 145 (145 eyes)

**Comparator:**

- sham injection n = 143 (143 eyes)

Participants received pegaptanib 0.3 mg or sham injections every 6 weeks in year 1 (total 9 injections) and could receive focal/grid photocoagulation beginning at week 18. During year 2, participants received injections as often as every 6 weeks according to pre-specified criteria

### Outcomes

**Primary outcome:**

- 10-letter (2-line) improvement from baseline at 12 months (ETDRS chart)

**Secondary outcomes:**

- 10-letter improvement from baseline at 24 months
- changes from baseline in mean VA
- 15-letter (3-line) improvement in VA
- change in degree of retinopathy of 2 steps based on the 12-step scale of retinopathy
- decrease in retinal thickness at the centre point by 25% and 50%
- focal or grid laser
- change in NEI VFQ-25 and EQ-5D

Follow-up: 12 and 24 months

### Notes

Dates participants enrolled: September 2005 to July 2009

Funding: *Sponsored by Pfizer Inc, New York, New York. The sponsor participated in the*
Risk of bias

<table>
<thead>
<tr>
<th>Bias</th>
<th>Authors’ judgement</th>
<th>Support for judgement</th>
</tr>
</thead>
<tbody>
<tr>
<td>Random sequence generation (selection bias)</td>
<td>Low risk</td>
<td>“[...] subjects were centrally allocated to receive either pegaptanib 0.3 mg or sham injections (1:1) using a dynamic minimization procedure stratified by the site, hemoglobin A1c (&lt;7.6% vs &gt;=7.6%), systolic blood pressure (&lt;140 vs &gt;=140 mmHg), diastolic blood pressure (80 vs 80 mmHg), and baseline BCVA (&lt;54 vs &gt;=54 letters); the dynamic minimization used a stochastic treatment allocation algorithm based on the variance method.” Page 3</td>
</tr>
<tr>
<td>Allocation concealment (selection bias)</td>
<td>Low risk</td>
<td>“…subjects were centrally allocated...” Page 3</td>
</tr>
<tr>
<td>Blinding of participants and personnel (performance bias)</td>
<td>Low risk</td>
<td>“To maintain masking, the intravitreal procedure was identical between the sham and comparator arms, with the difference lying only in the application of an empty barrel of a needleless syringe in the sham procedure designed to mimic the intravitreal injection.” Page 3</td>
</tr>
<tr>
<td>Blinding of outcome assessment (detection bias)</td>
<td>Low risk</td>
<td>“Throughout the study, BCVA was measured at 4 m by the study refractionist/ophthalmologist, who was masked to the subject’s treatment and to the subject’s previous visual acuity (VA) assessments”. Page 3</td>
</tr>
<tr>
<td>Incomplete outcome data (attrition bias)</td>
<td>Unclear risk</td>
<td>At 1 year 116/144 (81%) pegaptanib-treated participants and 114/142 (80%) controls completed the 54-week visit. Adverse events led to discontinuation of 5 treated and 7 control participants. At 2 years 66 participants in each group completed the 102 week visit. ITT analysis with LOCF was used leading to the analysis of 133 treated and 127 control participants.</td>
</tr>
</tbody>
</table>
Selective reporting (reporting bias) | Low risk | All primary outcomes reported
---|---|---
Other bias | Low risk | No other biases identified
Overall risk of bias | Low risk | Low risk of bias for most items

### Nepomuceno 2013

**Methods**
- Parallel group RCT and within-person study
- People randomised to treatment but two eyes sometimes included. If two eyes included then fellow eye randomised to other treatment

**Participants**
- Country: Brazil
- Number of people randomised: 48 (63 eyes)
- Average age 64 years
- Sex: 55% women (based on eyes included in analyses)
- Inclusion criteria:
  - centre-involved DMO defined as a central subfield thickness > 300 mm on Spectral Domain-OCT, despite at least 1 session of macular laser photocoagulation performed at least 3 months previously
  - BCVA ETDRS measurement between 0.3 logMAR (Snellen equivalent: 20/40) and 1.6 logMAR (Snellen equivalent: 20/800)
- Exclusion criteria:
  - vitreomacular traction on SD-OCT
  - PDR needing PRP or anticipated to need PRP in the next 12 months
  - macular capillary dropout on fluorescein angiography
  - history of glaucoma or ocular hypertension (defined as an intraocular pressure > 22 mmHg)
  - an ocular condition (other than diabetes) that, in the opinion of the investigator, might affect macular oedema or alter VA during the course of the study (e.g. retinal vein occlusion, uveitis or other ocular inflammatory disease, neovascular glaucoma, etc)
  - systemic corticosteroid therapy
  - any condition that, in the opinion of the investigator, might preclude follow-up throughout the study period

**Interventions**
- Intervention:
  - bevacizumab (1.5 mg) n = 32 eyes
- Comparator:
  - ranibizumab (0.5 mg) n = 28 eyes

“Retreatment with the originally assigned treatment was performed monthly if central subfield thickness was greater than 275 mm.”

“If, after 3 consecutive injections, there was not a reduction in central subfield thickness of at least 10% or an increase in BCVA of at least 5 letters compared with baseline, the patient could, at the discretion of the treating ophthalmologist, receive focal/grid laser photocoagulation or continue to receive the same intravitreal medication for an additional 3 consecutive visits.”
Outcomes

Outcomes reported in publication (primary outcome not specified):
- BCVA (standardised ETDRS refraction protocol)
- Retinal thickness (using OCT)

On ClinicalTrials.gov following outcomes listed:
- Primary outcome measures: CSFT change (time frame: monthly from baseline to week 48; not designated as a safety issue); CSFT measured with SD-OCT
- Secondary outcome measures: BCVA change (time frame: monthly from baseline to week 48; not designated as a safety issue); BCVA using ETDRS charts

Notes

Dates participants enrolled: July 2010 to August 2011

Funding: "Fundacao de Amparo a Pesquisa do Estado de Sao Paulo (FAPESP), grant number 2010/013368; and Fundacao Apoioao Ensino, Pesquisa e Assistencia (FAEPA) do Hospital das Clinicas da Faculdade de Medicina de Ribeirao Preto da Universidade de Sao Paulo."

Conflict of interest: Rodrigo Jorge received travel support from Novartis to attend the 2012 American Society of Retina Specialists (ASRS) meeting

Trial registration: NCT01487629

Risk of bias

<table>
<thead>
<tr>
<th>Bias</th>
<th>Authors' judgement</th>
<th>Support for judgement</th>
</tr>
</thead>
<tbody>
<tr>
<td>Random sequence generation (selection bias)</td>
<td>Low risk</td>
<td>&quot;... received the randomised treatment according to a computer-generated sequence&quot; Page 503</td>
</tr>
<tr>
<td>Allocation concealment (selection bias)</td>
<td>Unclear risk</td>
<td>Not mentioned</td>
</tr>
<tr>
<td>Blinding of participants and personnel (performance bias)</td>
<td>Low risk</td>
<td>&quot;Examiners (E.T., F.P.P.A., R.P.) were masked regarding which treatment drug was used for each patient. Throughout the study, a single masked, certified examiner performed BCVA measurements prior to any other study procedure. Patients, OCT technicians, and fundus photographers were also masked to treatment group&quot;. Page 504</td>
</tr>
<tr>
<td>Blinding of outcome assessment (detection bias)</td>
<td>Low risk</td>
<td>&quot;Examiners (E.T., F.P.P.A., R.P.) were masked regarding which treatment drug was used for each patient. Throughout the study, a single masked, certified examiner performed BCVA measurements prior to any other study procedure. Patients, OCT technicians, and fundus photographers were also masked to treatment group&quot;. Page 504</td>
</tr>
<tr>
<td>Incomplete outcome data (attrition bias)</td>
<td>Low risk</td>
<td>&quot;The 3 patients excluded from the outcomes analyses consisted of 1 patient in the IV ranibizumab group who developed Staphylo-</td>
</tr>
</tbody>
</table>
Nepomuceno 2013  (Continued)

Coccus aureus endophthalmitis after the first injection (this patient chose to exit the study and he did not complete any further study visits); 1 patient in the IV bevacizumab group who developed advanced posterior subcapsular cataract, which precluded adequate SDOCT images, after the ninth follow-up visit; and 1 patient from the IV bevacizumab group who missed 3 consecutive follow-up visits.” Page 504

<table>
<thead>
<tr>
<th>Selective reporting (reporting bias)</th>
<th>Low risk</th>
<th>Both outcomes listed on trial registration reported</th>
</tr>
</thead>
<tbody>
<tr>
<td>Other bias</td>
<td>Unclear risk</td>
<td>Fifteen out of 48 participants with both eyes in analyses</td>
</tr>
<tr>
<td>Overall risk of bias</td>
<td>Low risk</td>
<td>Low risk of bias for most items</td>
</tr>
</tbody>
</table>

READ2 2009

Methods
Parallel group RCT
One eye per person; if both eyes were eligible, the eye with the greater centre subfield thickness was entered

Participants
Country: USA
Number of people randomised: 126 (126 eyes)
Average age: 62 years
Sex: 59% women
Inclusion criteria:
- 18 years and older
- diabetes
- DMO
- reduction in VA between 20/40-20/320
- centre subfield thickness measured by OCT ≥ 250 µm
- HbA1c ≥ 6% within 12 months before randomisation
- no potential contributing causes to reduced VA other than DMO
- reasonable expectation that scatter laser photocoagulation would not be required for the next 6 months
Exclusion criteria:
- received focal/grid laser treatment within 3 months
- intracocular injection of steroid within 3 months
- intracocular injection of a VEGF antagonist within 2 months

Interventions
Intervention:
- ranibizumab 0.5 mg n = 42 (42 eyes)
- ranibizumab 0.5 mg plus laser photocoagulation n = 42 (42 eyes)
Comparator:
- laser photocoagulation n = 42 (42 eyes)
Participants were randomised 1:1:1 to receive 0.5 mg ranibizumab at baseline and months 1, 3, and 5 (group 1), focal or grid laser photocoagulation at baseline and month 3 if needed (group 2), or a combination of 0.5 mg ranibizumab and focal or grid laser at baseline and month 3 (group 3). Starting at month 6, if retreatment criteria were met, all participants could be treated with ranibizumab.

Duration: primary outcome at 6 months, extension to 24 and 36 months

### Outcomes

As reported in publications:

**Primary outcome:**
- change in BCVA between baseline and follow-up

**Secondary outcomes:**
- change in BCVA between baseline and month 24
- 3 or more lines or 2 or more lines improvement at month 24
- change in foveal thickness between baseline and month 24
- elimination of 90% or 50% excess foveal thickness

On ClinicalTrials.gov

"Primary Outcome Measures: Improvement in vision of 15 or more letters, or achieve a final vision of 50 letters (20/25) or better if baseline VA was 40 letters (20/40) [Time Frame: 6 mos, 12 mos and 24 mos. Study Extended to 36 mos.] [Designated as safety issue: Yes]

Secondary Outcome Measures: Several outcomes related to OCT measurements and fluorescein angiography. [Time Frame: 6 mos, 12 mos and 24 mos, study extended to 36 mos.] [Designated as safety issue: Yes]"

Follow-up: 6 months and 24 months.

### Notes

Dates participants enrolled: not reported

Funding: "Sponsored by the Juvenile Diabetes Research Foundation and Genentech, Inc."

Conflict of interest: "QDN and PAC have served as members of Expert Panels for Genentech, Inc. without receiving an honorarium during the time of this study, but JHU has recently negotiated a contract through which JHU receives compensation. QDN is a consultant for Bausch and Lomb and has research support from Genentech, Inc., and Regeneron, Inc. PAC serves on the data and safety monitoring committee for a phase III trial sponsored by Regeneron, Inc., and has research support from Genentech, Alimera, and CoMentis for diabetic macular edema trials. Diana Do receives research support from Genentech. These activities are being managed by the Conflict of Interest Committee of the Johns Hopkins University School of Medicine. JSH is a consultant for Genentech, Alcon, Allergan, Bausch and Lomb, Eyemaginations, Fovea, Genzyme, Heidelberg, iScience, ISTA, Jerini, LPath, NeoVista, Nidal Vision, Novagadi, Novartis, Ophtherion, Oxigene, Paloma, Pfizer, Regeneron, Resolvyx, Schering Plough, Scyfix, and VisionCare and has received honoraria from Genentech, Heidelberg, Jerini, NeoVista, Optimedica, and Regeneron. JL has received honoraria from Genentech. DB is a consultant and has received honoraria from Genentech, Novartis, Alcon, Allergan, and Pfizer. PA is a consultant for Genentech" Page 2181

Trial registration: NCT00407381

### Risk of bias

<table>
<thead>
<tr>
<th>Bias</th>
<th>Authors’ judgement</th>
<th>Support for judgement</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
### READ2 2009 (Continued)

<table>
<thead>
<tr>
<th>Source of bias</th>
<th>Risk of Bias</th>
<th>Rationale</th>
</tr>
</thead>
<tbody>
<tr>
<td>Random sequence generation (selection bias)</td>
<td>Unclear risk</td>
<td>Unclear method of sequence generation and information could not be obtained from the authors</td>
</tr>
<tr>
<td>Allocation concealment (selection bias)</td>
<td>Unclear risk</td>
<td>Unclear method of allocation concealment and information could not be obtained from the authors</td>
</tr>
<tr>
<td>Blinding of participants and personnel (performance bias)</td>
<td>High risk</td>
<td>Unclear if masked and who was masked and information could not be obtained from the authors</td>
</tr>
<tr>
<td>Blinding of outcome assessment (detection bias)</td>
<td>High risk</td>
<td>Unclear if masked and who was masked and information could not be obtained from the authors</td>
</tr>
<tr>
<td>Incomplete outcome data (attrition bias)</td>
<td>Unclear risk</td>
<td>Participants randomised to each group: 33 ranibizumab, 34 ranibizumab + laser, 34 laser. Completed participants at 1 year: 29, 29, 30 (85% to 88%). Completed participants at 2 years: 24, 26, 24 (71% to 76%). Causes of missing data were balanced across groups</td>
</tr>
<tr>
<td>Selective reporting (reporting bias)</td>
<td>High risk</td>
<td>The primary outcome differed in the protocol and the final report</td>
</tr>
<tr>
<td>Other bias</td>
<td>Low risk</td>
<td>No other source of bias identified</td>
</tr>
<tr>
<td>Overall risk of bias</td>
<td>High risk</td>
<td>High risk of bias for nearly half the items and unclear risk for the others</td>
</tr>
</tbody>
</table>

### RELATION 2012

<table>
<thead>
<tr>
<th>Methods</th>
<th>Parallel group RCT One eye per person, eye with worse VA selected</th>
</tr>
</thead>
<tbody>
<tr>
<td>Participants</td>
<td>Country: Germany Number of people randomised: 128 (128 eyes) Average age: 64 years (range 31 to 79) Sex: 37% women Inclusion criteria: ● 18 years or older ● diabetes ● visual impairment (BCVA between 78-39 letters, testing distance 4 m) due to focal or diffuse DMO in at least one eye eligible for laser treatment in the opinion of</td>
</tr>
<tr>
<td>Exclusion criteria:</td>
<td></td>
</tr>
<tr>
<td>---------------------</td>
<td></td>
</tr>
<tr>
<td>• other eye diseases and conditions that might affect VA</td>
<td></td>
</tr>
<tr>
<td>• other eye and systemic treatments</td>
<td></td>
</tr>
<tr>
<td>• pregnancy or possibility of being pregnant</td>
<td></td>
</tr>
<tr>
<td>• Inability to comply with follow-up</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Interventions</th>
<th>Intervention:</th>
</tr>
</thead>
<tbody>
<tr>
<td>• ranibizumab (0.5 mg) plus laser n = 85 (85 eyes)</td>
<td></td>
</tr>
<tr>
<td>• laser plus sham injection n = 43 (43 eyes)</td>
<td></td>
</tr>
</tbody>
</table>

Ranibizumab was applied at baseline, 30, 60, 90 days and reapplied at intervals no shorter than 28 days and laser was applied at baseline and re-applied if needed at intervals no shorter than 3 months.

<table>
<thead>
<tr>
<th>Outcomes</th>
<th>Primary outcome:</th>
</tr>
</thead>
<tbody>
<tr>
<td>• mean change in BCVA from baseline to month 12 (ETDRS chart, 4 m)</td>
<td></td>
</tr>
</tbody>
</table>

Secondary outcomes:
• adverse events

<table>
<thead>
<tr>
<th>Notes</th>
<th>Dates participants enrolled: July 2010 to May 2011, terminated early</th>
</tr>
</thead>
<tbody>
<tr>
<td>Funding: Novartis</td>
<td></td>
</tr>
<tr>
<td>Conflict of interest: Novartis</td>
<td></td>
</tr>
<tr>
<td>Trial registration: NCT01131585</td>
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</table>

<table>
<thead>
<tr>
<th>Risk of bias</th>
<th>Authors' judgement</th>
<th>Support for judgement</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bias</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Random sequence generation (selection bias)</td>
<td>Unclear risk</td>
<td>Not reported</td>
</tr>
<tr>
<td>Allocation concealment (selection bias)</td>
<td>Unclear risk</td>
<td>Not reported</td>
</tr>
<tr>
<td>Blinding of participants and personnel (performance bias)</td>
<td>Unclear risk</td>
<td>Reported as double-masked, but no details given</td>
</tr>
<tr>
<td>All outcomes</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Blinding of outcome assessment (detection bias)</td>
<td>Unclear risk</td>
<td>Reported as double-masked, but no details given</td>
</tr>
<tr>
<td>All outcomes</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Incomplete outcome data (attrition bias)</td>
<td>High risk</td>
<td>Missing data: combined laser and ranibizumab: 7/85 (7%); laser 11/43 (26%)</td>
</tr>
<tr>
<td>All outcomes</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Selective reporting (reporting bias)</td>
<td>High risk</td>
<td>Only mean change of VA and harms reported</td>
</tr>
</tbody>
</table>
**RELATION 2012 (Continued)**

<table>
<thead>
<tr>
<th>Other bias</th>
<th>Low risk</th>
<th>Study terminated early due to European Medicine Agency approval of ranibizumab for DMO but this is independent of effect estimates</th>
</tr>
</thead>
<tbody>
<tr>
<td>Overall risk of bias</td>
<td>Unclear risk</td>
<td>Unclear risk of bias for most items</td>
</tr>
</tbody>
</table>

**RESOLVE 2010**

| Methods | Parallel group RCT  
One eye per person, eye with worse VA selected |
| --- | --- |
| Participants | Country: unclear exactly where conducted. Investigators from Australia, Denmark, Austria, France, Germany, Italy, Korea, Portugal, Spain, Switzerland, UK  
Number of people randomised: 151 (151 eyes)  
Average age: 64 years (range 32 to 85)  
Sex: 46% women  
Inclusion criteria:  
• 18 years or older  
• diabetes mellitus  
• stable HbA1c levels (≤ 12%)  
• DMO with centre involvement in at least one eye (study eye)  
• CRT ≥ 300 µm (Stratus Zeiss Meditec)  
• BCVA score between 73-39 letters inclusively, using ETDRS charts at a testing distance of 4 m (approximate Snellen equivalent of 20/40-20/160)  
• decreased vision attributed to foveal thickening from DMO, that was not explained by any other causes in the opinion of the investigator  
• laser photocoagulation, additional or first treatment, could be withheld for at least 3 months after randomisation  
Exclusion criteria:  
• PRP (focal peripheral laser photocoagulation) performed within 6 months prior to study entry. Grid/central laser photocoagulation was excluded except for participants with only mild laser burns at least 1000 µm from the centre of the fovea performed more than 6 months before the trial commenced  
• PDR in the study eye, with the exception of tufts of neovascularization < 1 disc area with no vitreous haemorrhage. As well as those with area of retinal ischaemia ≥ 500 µm and located ≤ 500 µm from the centre of the macula of the study eye as assessed by fluorescein angiography at visit 1 and confirmed by a central reading centre  
• participants with unstable medical conditions such as poor glycaemic or BP control  
• participants with hypertension for whom a change in antihypertensive treatment was initiated within 2 months preceding start of trial were not enrolled unless BP was maintained below 160/100 mmHg for at least 1 month prior to the first day of the trial by antihypertensive treatment  
• history of treatment with systemic corticosteroids within 4 months prior to randomisation or topical, rectal or inhaled corticosteroids in current use more than 2 times per week |
RESOLVE 2010  (Continued)

- previous participation in a study on antiangiogenic drugs
- ocular disorders and history of any condition that might confound the interpretation of study results or might render participant at high risk for treatment complications
- ocular inflammation in either eye or history of cataract surgery in the study eye within 6 months before study initiation
- pre-menopausal women not using adequate contraception and pregnant or nursing women

### Interventions

<table>
<thead>
<tr>
<th>Intervention</th>
<th>Comparator</th>
</tr>
</thead>
<tbody>
<tr>
<td>- ranibizumab (0.3 mg or 0.5 mg) n = 102 (102 eyes)</td>
<td>- sham injection n = 49 (49 eyes)</td>
</tr>
</tbody>
</table>

### Outcomes

<table>
<thead>
<tr>
<th>Primary outcome</th>
<th>Secondary outcomes</th>
</tr>
</thead>
<tbody>
<tr>
<td>mean change in BCVA from baseline at 1 month and 12 months</td>
<td>mean change in BCVA and CRT from baseline at 12 months</td>
</tr>
<tr>
<td>categorised BCVA outcome</td>
<td>safety</td>
</tr>
</tbody>
</table>

### Notes

- Dates participants enrolled: not reported
- Funding: Novartis
- Conflict of interest: authors served on advisory boards for Novartis and received honoraria and travel and accommodation payments; Novartis employees assisted with the analysis, interpretation and writing
- Trial registration: NCT00284050

### Risk of bias

<table>
<thead>
<tr>
<th>Bias</th>
<th>Authors’ judgement</th>
<th>Support for judgement</th>
</tr>
</thead>
<tbody>
<tr>
<td>Random sequence generation</td>
<td>Low risk</td>
<td>&quot;Eligible patients were randomised 1:1:1 to either ranibizumab (0.3 mg or 0.5 mg) or sham treatment according to a computer-generated randomised allocation schedule&quot; Online appendix page 1</td>
</tr>
<tr>
<td>Allocation concealment</td>
<td>Low risk</td>
<td>&quot;...allocation schedule (kept at a secure site and accessible only to the injecting physician&quot; Online appendix page 1</td>
</tr>
</tbody>
</table>

"Based on the patient strata the injecting physician would take the treatment allocation card and tear-off the cover and follow instructions to choose vial from the box as indicated (3 boxes, randomisation block size 3). The randomisation data were kept strictly confidential until database"
**RESOLVE 2010 (Continued)**

<table>
<thead>
<tr>
<th>Source of Bias</th>
<th>Type of Bias</th>
<th>Risk of Bias</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Blinding of participants and personnel (performance bias)</td>
<td>Low risk</td>
<td></td>
<td>Sham injection for masking participants</td>
</tr>
<tr>
<td>All outcomes</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Blinding of outcome assessment (detection bias)</td>
<td>Unclear risk</td>
<td></td>
<td>“Masking was maintained through appointment of a minimum of 2 investigators at each study site; unmasked injecting physician and a masked evaluating physician (roles could not be switched).” Online appendix page 1</td>
</tr>
<tr>
<td>All outcomes</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Incomplete outcome data (attrition bias)</td>
<td>Low risk</td>
<td></td>
<td>Participants who completed the trial at 1 year: 92/102 ranibizumab and 40/49 sham. Causes of missingness were balanced ITT analysis with LOCF was used</td>
</tr>
<tr>
<td>All outcomes</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Selective reporting (reporting bias)</td>
<td>Low risk</td>
<td></td>
<td>We could not find a protocol, but primary outcomes were stated in the methods and were those routinely used in the field</td>
</tr>
<tr>
<td>Other bias</td>
<td>Low risk</td>
<td></td>
<td>No other source of bias identified</td>
</tr>
<tr>
<td>Overall risk of bias</td>
<td>Low risk</td>
<td></td>
<td>Low risk of bias for most items</td>
</tr>
</tbody>
</table>

**RESPOND 2013**

<table>
<thead>
<tr>
<th>Method</th>
<th>Parallel group RCT</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>One eye per person, unclear how eye selected</td>
</tr>
</tbody>
</table>

**Participants**

Country: Canada  
Number of people randomised: 239 (239 eyes)  
Average age: 62 years (range 26 to 87)  
Sex: 40% women  
Inclusion criteria:  
- 18 years or older  
- stable type 1 or type 2 diabetes with HbA1c ≤ 10%  
- visual impairment due to focal or diffuse DMO in at least one eye eligible for laser treatment in the opinion of the investigator  
Exclusion criteria:  
- active conditions in study eye that could prevent improvement in VA  
- active eye infection or inflammation  
- history of stroke, renal failure or active hypertension
## Interventions

**Intervention:**
- ranibizumab (0.5 mg) n = 80 (80 eyes)
- ranibizumab (0.5 mg) plus laser n = 78 (78 eyes)

**Comparator:**
- laser n = 81 (81 eyes)

For combination and monotherapy, ranibizumab was administered as 3 monthly injections, then 10 months PRN injections given/withheld based on DME stability criteria. Laser was administered according to ETDRS guidelines at intervals of > 3 months.

## Outcomes

**On ClinicalTrials.gov**

**Primary Outcome Measures:** Measure: mean change from baseline in Best Correct Visual Acuity (BCVA) [Time Frame: 12 months] [Designated as safety issue: No]

**Secondary Outcome Measures:**
- number of patients with visual acuity above 73 letters [Time Frame: 3, 6, 9 and 12 months]
- number of patients with improvement in BCVA [Time Frame: 3, 6, 9 and 12 months]
- time course of BCVA changes [Time Frame: 3, 6, 9 and 12 months]
- change in central retinal thickness and other anatomical changes [Time Frame: 3, 6, 9 and 12 months]
- 15-letter (3-line) gain in BCVA [Time Frame: 3, 6, 9 and 12 months]

## Notes

Dates participants enrolled: July 2010 to March 2013

Funding: Novartis

Conflict of interest: authors not reported since the study was obtained as a Novartis public report

Trial registration: NCT01135914

## Risk of bias

<table>
<thead>
<tr>
<th>Bias</th>
<th>Authors’ judgement</th>
<th>Support for judgement</th>
</tr>
</thead>
<tbody>
<tr>
<td>Random sequence generation (selection bias)</td>
<td>Low risk</td>
<td>“Randomization was stratified by centre and followed a permuted block size of 6.”</td>
</tr>
<tr>
<td>Allocation concealment (selection bias)</td>
<td>Unclear risk</td>
<td>Not reported</td>
</tr>
<tr>
<td>Blinding of participants and personnel (performance bias) All outcomes</td>
<td>High risk</td>
<td>Unmasked study (described as open-label)</td>
</tr>
<tr>
<td>Blinding of outcome assessment (detection bias) All outcomes</td>
<td>High risk</td>
<td>Unmasked study (described as open-label)</td>
</tr>
<tr>
<td>Incomplete outcome data (attrition bias) All outcomes</td>
<td>High risk</td>
<td>More missing data in the laser arm (27%), mainly due to lack of efficacy, compared to the 2 ranibizumab arms (5% to 6%)</td>
</tr>
</tbody>
</table>
### RESPOND 2013 (Continued)

<table>
<thead>
<tr>
<th>Selective reporting (reporting bias)</th>
<th>Low risk</th>
<th>VA, OCT data and harms adequately reported (only loss of vision not reported)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Other bias</td>
<td>Low risk</td>
<td>No other bias identified</td>
</tr>
<tr>
<td>Overall risk of bias</td>
<td>High risk</td>
<td>High risk of bias for most items</td>
</tr>
</tbody>
</table>

### RESTORE 2011

<table>
<thead>
<tr>
<th>Methods</th>
<th>Parallel group RCT One eye per person, eye with worse VA selected unless other eye more suitable for treatment</th>
</tr>
</thead>
</table>
| Participants      | Country: 10 European countries, Australia, Canada, Turkey  
Number of people randomised: 345 (345 eyes)  
Average age: 63 years  
Sex: 42% women  
Inclusion criteria:  
- 18 years or older  
- diabetes mellitus (according to the American Diabetes Association or World Health Organization guidelines)  
- HbA1c ≤ 10%  
- visual impairment due to DMO  
- stable medication for the management of diabetes within 3 months before randomisation and expected to remain stable during the study  
- visual impairment due to focal or diffuse DMO in at least 1 eye that was eligible for laser treatment in the opinion of the investigator  
- BCVA letter score between 78-39, both inclusive, based on ETDRS-like VA testing charts administered at a starting distance of 4 m (approximate Snellen equivalent 20/32-20/160)  
- decreased vision due to DMO and not other causes, in the investigator's opinion (at visit 1)  
Exclusion criteria:  
- concomitant conditions in the study eye that could prevent the improvement in VA on the study treatment in the investigator's opinion  
- active intraocular inflammation or infection in either eye  
- uncontrolled glaucoma in either eye (e.g. IOP > 24 mmHg on medication, or from the investigator's judgement)  
- laser PRP (within 6 months) or focal/grid laser photocoagulation (within 3 months) before study entry  
- treatment with antiangiogenic drugs in the study eye within 3 months before randomisation  
- history of stroke  
- systolic BP > 160 mmHg or diastolic BP > 100 mmHg  
- untreated hypertension  
- change in antihypertensive treatment within 3 months preceding baseline |
<table>
<thead>
<tr>
<th>Interventions</th>
<th>Intervention:</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>• ranibizumab (0.5 mg) plus sham laser n = 116 (116 eyes)</td>
</tr>
<tr>
<td></td>
<td>• ranibizumab (0.5 mg) plus laser n = 118 (118 eyes)</td>
</tr>
<tr>
<td>Comparator</td>
<td>• laser treatment plus sham injections n = 111 (111 eyes)</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Outcomes</th>
<th>Primary outcome:</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>• mean average change in BCVA from baseline over 12 months</td>
</tr>
<tr>
<td></td>
<td>Secondary outcomes:</td>
</tr>
<tr>
<td></td>
<td>• VA improvement</td>
</tr>
<tr>
<td></td>
<td>• BCVA letter score 73 (20/40 Snellen equivalent) at month 12</td>
</tr>
<tr>
<td></td>
<td>• mean change in BCVA letter score</td>
</tr>
<tr>
<td></td>
<td>• mean change in central retinal (subfield) thickness</td>
</tr>
<tr>
<td></td>
<td>• patient-reported outcomes</td>
</tr>
<tr>
<td></td>
<td>• safety</td>
</tr>
</tbody>
</table>

Follow-up: 12 months

<table>
<thead>
<tr>
<th>Notes</th>
<th>Dates participants enrolled: not reported</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Funding: Novartis</td>
</tr>
<tr>
<td></td>
<td>Conflict of interest: authors reported financial support of Novartis or were Novartis employees</td>
</tr>
<tr>
<td></td>
<td>Trial registration: NCT00906464</td>
</tr>
</tbody>
</table>

**Risk of bias**

<table>
<thead>
<tr>
<th>Bias</th>
<th>Authors’ judgement</th>
<th>Support for judgement</th>
</tr>
</thead>
<tbody>
<tr>
<td>Random sequence generation (selection bias)</td>
<td>Low risk</td>
<td>“A randomization list was produced by, or under the responsibility of, Novartis Drug Supply Management using a validated system that automated the random assignment of treatment arms to randomization numbers in the specified ratio.” Appendix 1</td>
</tr>
<tr>
<td>Allocation concealment (selection bias)</td>
<td>Low risk</td>
<td>Central randomisation using an electronic Case Report Form after each participant was included by study investigators</td>
</tr>
</tbody>
</table>
| Blinding of participants and personnel (performance bias) All outcomes | Low risk          | “The masked BCVA assessor evaluated the visual acuity of the patient and provided the results to the evaluating investigator who also was masked to the treatment assignment. The evaluating investigator was responsible for all other aspects of the study, excluding the injection procedures. Based on all the performed clinical assessments and the visual acuity (VA) results received from the BCVA assessor, the evaluating investigator had to decide on the
treatment requirements for the patient each month and communicated this decision to the treating investigator. The treating investigator was unmasked to the treatment assignment and performed all injections or laser treatment as well as the corresponding sham treatments. He/she was required not to be involved in any other aspect of the study and not to divulge the patient’s treatment assignment to anyone. Once the designated roles were determined, the roles could not be switched at any time during the conduct of the study. Every effort was made to limit the number of unmasked study personnel to ensure the integrity of this masked study. An independent review and standardized grading of fundus photography, fluorescein angiography, and optical coherence tomography (OCT) images for the patients screened and enrolled in the study was performed at a central reading center that did not have access to any other data of the patients.” Appendix 1

<table>
<thead>
<tr>
<th><strong>Blinding of outcome assessment (detection bias)</strong></th>
<th><strong>Low risk</strong></th>
<th><strong>See above</strong></th>
</tr>
</thead>
<tbody>
<tr>
<td>All outcomes</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

| **Incomplete outcome data (attrition bias)**      | **Low risk** | **Participants randomised in each group were: 116 ranibizumab, 118 ranibizumab + laser, 111 laser** |
| All outcomes                                      |              | **At 1 year complete participants were 87.9%, 87.3% and 88.3% respectively** |
|                                                   |              | **There were 2 deaths in each of the 3 treatment arms** |
|                                                   |              | **Used ITT analysis with LOCF** |

| **Selective reporting (reporting bias)**          | **Low risk** | **We could not find a protocol, but primary outcomes were stated in the methods and were those routinely used in the field** |
|                                                  |              |               |

| **Other bias**                                    | **Low risk** | **No other source of bias identified** |
|                                                  |              |               |

| **Overall risk of bias**                          | **Low risk** | **Low risk of bias for most items** |
|                                                  |              |               |
## REVEAL 2015

| Methods | Parallel group RCT  
<table>
<thead>
<tr>
<th></th>
<th>One eye per person, eye with worse VA selected unless other eye more suitable for treatment</th>
</tr>
</thead>
</table>
| Participants | Country: Asian population from 52 centres across 6 countries, or regions: China, Hong Kong, Japan, South Korea, Singapore, and Taiwan  
|         | Number of people randomised: 396 (396 eyes)  
|         | Average age: 61 years  
|         | Sex: 44% women  
|         | Inclusion criteria:  
|         | • 18 years or older  
|         | • diabetes mellitus (according to the American Diabetes Association or World Health Organization guidelines)  
|         | • HbA1c ≤ 10%  
|         | • visual impairment due to DMO  
|         | • stable medication for the management of diabetes within 3 months before randomisation and expected to remain stable during the study  
|         | • visual impairment due to focal or diffuse DMO in at least 1 eye that was eligible for laser treatment in the opinion of the investigator  
|         | • BCVA letter score between 78-39, both inclusive, based on ETDRS-like VA testing charts administered at a starting distance of 4 m (approximate Snellen equivalent 20/32-20/160)  
|         | • decreased vision due to DMO and not other causes, in the investigator’s opinion (at visit 1)  
|         | Exclusion criteria:  
|         | • concomitant conditions in the study eye that could prevent the improvement in VA on the study treatment in the investigator’s opinion  
|         | • active intraocular inflammation or infection in either eye  
|         | • uncontrolled glaucoma in either eye (e.g. IOP > 24 mmHg on medication, or from the investigator’s judgement)  
|         | • laser PRP (within 6 months) or focal/grid laser photocoagulation (within 3 months) before study entry  
|         | • treatment with antiangiogenic drugs in the study eye within 3 months before randomisation  
|         | • history of stroke  
|         | • systolic BP > 160 mmHg or diastolic BP > 100 mmHg  
|         | • untreated hypertension  
|         | • change in antihypertensive treatment within 3 months preceding baseline |
| Interventions | “Patients were randomised in a 1:1:1 ratio to 1 of 3 treatment arms: intravitreal ranibizumab 0.5 mg injection + sham laser, intravitreal ranibizumab 0.5 mg injection + active laser, or active laser + sham injections for 12 months” Page 1404  
|         | Number in each group: ranibizumab + sham laser (n = 133), ranibizumab + active laser (n = 132), or sham injection + active laser (n = 131) |
| Outcomes | Primary outcome:  
|         | • mean average change in BCVA from baseline over 12 months  
|         | Secondary outcomes:  
|         | • several BCVA expressions  
|         | • mean change in central retinal (subfield) thickness |

Anti-vascular endothelial growth factor for diabetic macular oedema: a network meta-analysis (Review)  
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<table>
<thead>
<tr>
<th>Risk of bias</th>
<th>Authors’ judgement</th>
<th>Support for judgement</th>
</tr>
</thead>
<tbody>
<tr>
<td>Random sequence generation (selection bias)</td>
<td>Low risk</td>
<td>“At Visit 2, all patients who fulfilled all the inclusion/exclusion criteria were given the lowest available number on the randomization list. This number assigned them to one of the treatment arms. The investigator entered the randomization number on the electronic case report form. A randomization list was produced by, or under the responsibility of Novartis Drug Supply Management using a validated system that automated the random assignment of treatment arms to randomization numbers in the specified ratio”</td>
</tr>
<tr>
<td>Allocation concealment (selection bias)</td>
<td>Low risk</td>
<td>See above</td>
</tr>
<tr>
<td>Blinding of participants and personnel (performance bias)</td>
<td>Low risk</td>
<td>“To ensure successful masking in this double-masked study, at the start of the study and at each study site, the following site personnel were required to demonstrate their role: BCVA assessor and evaluating investigator (masked to the treatment assignment) and treating investigator (unmasked to the treatment assignment)”</td>
</tr>
<tr>
<td>Blinding of outcome assessment (detection bias)</td>
<td>Low risk</td>
<td>See above</td>
</tr>
<tr>
<td>Incomplete outcome data (attrition bias)</td>
<td>High risk</td>
<td>Higher proportion of loss to follow-up in the laser group; this can decrease the benefit with anti-VEGF (see below)</td>
</tr>
</tbody>
</table>

Overall, 345 (87.1%) patients completed the study. The proportion of patients who discontinued the study was 7.5% in the ranibizumab arm, 13.6% in the ranibizumab active laser treatment arm, and
### REVEAL 2015  (Continued)

<table>
<thead>
<tr>
<th>Selective reporting (reporting bias)</th>
<th>Unclear risk</th>
<th>No protocol available</th>
</tr>
</thead>
<tbody>
<tr>
<td>Other bias</td>
<td>Low risk</td>
<td>No other bias identified</td>
</tr>
<tr>
<td>Overall risk of bias</td>
<td>Low risk</td>
<td>Low risk for most items</td>
</tr>
</tbody>
</table>

### RISE-RIDE

#### Methods
- Parallel group RCT
- One eye per person, unclear how eye selected

#### Participants
- Country: USA and South America
- Number of people randomised: 759 (759 eyes)
- Average age: 62 years
- Sex: 43% women
- Inclusion criteria:
  - 18 years or older
  - diabetes mellitus
  - decreased vision from DMO (study eye BCVA, 20/40-20/320 Snellen equivalent using ETDRS testing)
  - macular oedema (TD-OCT) central subfield thickness ≥ 275 µm
- Exclusion criteria:
  - prior vitreoretinal surgery
  - recent history (within 3 months of screening) of panretinal or macular laser in the study eye
  - intraocular corticosteroids antiangiogenic drugs
  - uncontrolled hypertension
  - uncontrolled diabetes (HbA1c > 12%)
  - recent (within 3 months) cerebrovascular accident, or myocardial infarction

#### Interventions
- Intervention:  
  - ranibizumab (0.3 mg or 0.5 mg) n = 244 (244 eyes)
- Comparator:  
  - sham injection n = 122 (122 eyes)

"The median number of ranibizumab injections was 24. The mean number of macular laser treatments over 24 months was 1.8 and 1.6 in the sham groups and 0.3 to 0.8 in the ranibizumab groups. Substantially more sham-treated patients received macular laser under the protocol-specified criteria or underwent panretinal photocoagulation for proliferative diabetic retinopathy." Page 5
### Outcomes

<table>
<thead>
<tr>
<th>Primary outcome:</th>
</tr>
</thead>
<tbody>
<tr>
<td>• gain of 15 or more ETDRS letters in BCVA score from baseline at 24 months (corresponding to 3 lines on the eye chart)</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Secondary outcomes: (at 24 months)</th>
</tr>
</thead>
<tbody>
<tr>
<td>• mean change from baseline BCVA score over time</td>
</tr>
<tr>
<td>• proportion of participants with BCVA Snellen equivalent of 20/40</td>
</tr>
<tr>
<td>• mean change from baseline BCVA score over time in participants with focal oedema as assessed on fluorescein angiography</td>
</tr>
<tr>
<td>• proportion of participants losing 15 letters in BCVA score from baseline</td>
</tr>
<tr>
<td>• mean change from baseline in OCT CFT over time</td>
</tr>
<tr>
<td>• proportion of participants with a 3-step progression from baseline in ETDRS retinopathy severity on fundus photography</td>
</tr>
<tr>
<td>• proportion of participants with resolution of leakage on FA</td>
</tr>
<tr>
<td>• mean number of macular laser treatments</td>
</tr>
</tbody>
</table>

Follow-up: 24 months

### Notes

Dates participants enrolled: June 2007 to January 2009

Funding: "This study was supported by Genentech Inc. Support for third-party writing assistance by Ivo Stoilov, MD, CMPP, of Envision Scientific Solutions was provided by Genentech Inc." "The sponsor participated in the design and conduct of the study; collection, management, analysis, and interpretation of the data; and preparation and review of the manuscript." Page 1121

Conflict of interest: "Dr Ip is a consultant/advisor for Eye Technology Ltd, Genentech Inc, NicOx, Notal Vision, QLT Phototherapeutics Inc, Regeneron, and Sirion and has received grant support from Allergan Inc. Drs Hopkins and Ehrlich and Ms Wong are employees of Genentech Inc, a member of the Roche Group. Drs Hopkins and Ehrlich hold equity and/or options in Roche." Page 1121

Trial registration: RIDE NCT00473382 RISE NCT00473330

### Risk of bias

<table>
<thead>
<tr>
<th>Bias</th>
<th>Authors' judgement</th>
<th>Support for judgement</th>
</tr>
</thead>
<tbody>
<tr>
<td>Random sequence generation (selection bias)</td>
<td>Low risk</td>
<td>“Randomization was stratified by study eye BCVA (55 vs 55 ETDRS letters), baseline HbA1c (&lt;=8% vs &gt;8%), prior DME therapy in the study eye (yes vs no), and study site. Dynamic randomization was used to obtain approximately a 1:1:1 ratio among groups (Fig 1). Randomization was done via interactive phone system. The sponsor developed the specifications for the randomization, and a third party programmed and held the randomization algorithm.” Page 3, Nguyen et al</td>
</tr>
<tr>
<td>Allocation concealment (selection bias)</td>
<td>Low risk</td>
<td>“Randomization was stratified by study eye BCVA (55 vs 55 ETDRS letters), baseline HbA1c (&lt;=8% vs &gt;8%), prior DME therapy in the study eye (yes vs no), and study site. Dynamic randomization was used to obtain approximately a 1:1:1 ratio among groups (Fig 1). Randomization was done via interactive phone system. The sponsor developed the specifications for the randomization, and a third party programmed and held the randomization algorithm.” Page 3, Nguyen et al</td>
</tr>
</tbody>
</table>
Blinding of participants and personnel (performance bias)  
All outcomes | Low risk | “Ocular assessments, including the need for macular laser, were made by evaluating ophthalmologists masked to patients’ treatment assignments. Study treatments were administered by treating ophthalmologists unmasked to treatment assignments but masked to ranibizumab dose. To improve patient masking, all patients received subconjunctival anesthesia before sham or active injections (performed as previously described).22 Study site personnel (except treating physicians and assistants), central reading center personnel, and the sponsor and its agents (except drug accountability monitors) were masked to treatment assignment. Treating physicians were masked to the assigned dose of ranibizumab.” Page 3, Nguyen et al

Blinding of outcome assessment (detection bias)  
All outcomes | Low risk | See above

Incomplete outcome data (attrition bias)  
All outcomes | Unclear risk | The 2-year study period was completed by 83.3% of participants in RISE and by 84.6% in RIDE; causes of missingness not reported

Selective reporting (reporting bias)  
All outcomes | Low risk | All VA cut-offs and secondary outcomes available at 2 years, although not at 1 year, as pre-planned

Other bias | Low risk | No other bias identified

Overall risk of bias | Low risk | Low risk of bias for most items
### Methods

Parallel group RCT

One or two eyes per person, in bilateral cases unclear how the second eye allocated

### Participants

Country: Iran  
Number of people randomised: 129 (150 eyes)  
Average age: 61 years  
Sex: 49% women  
Inclusion criteria:  
- clinically significant DMO based on ETDRS criteria  
Exclusion criteria:  
- previous PRP or focal laser photocoagulation  
- prior intraocular surgery or injection  
- history of glaucoma or ocular hypertension  
- VA of 20/40 or better, or worse than 20/300  
- presence of iris neovascularisation  
- high-risk PDR  
- significant media opacity  
- monocularity  
- pregnancy  
- serum creatinine ≥ 3 mg/dL  
- uncontrolled diabetes mellitus

### Interventions

**Intervention:**  
- bevacizumab (1.25 mg) n = 50 eyes  

**Comparator:**  
- laser photocoagulation n = 50 eyes  

Re-treatment was performed at 12-week intervals whenever indicated  
There was another intervention arm which combined bevacizumab with triamcinolone, but this is not included in this review (n = 50 eyes)

### Outcomes

**Primary outcome:**  
- change in BCVA (logMAR) at week 24 (data available at 36 weeks)  

**Secondary outcomes:**  
- VA change  
- CRT change assessed by OCT  
- injection-related complications

### Notes

Dates participants enrolled: September 2005 to May 2007  
Funding: “Supported by the Ophthalmic Research Center of Shahid Beheshti University (MC) Tehran, Iran.” Page 1150  
Conflict of interest: “The author(s) have no proprietary or commercial interest in any materials discussed in this article” Page 1150  
Trial registration: NCT00370669

### Risk of bias

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<thead>
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<td></td>
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<tr>
<td><strong>Soheilian 2007</strong></td>
<td>(Continued)</td>
<td></td>
</tr>
<tr>
<td>------------------</td>
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<tr>
<td><strong>Random sequence generation (selection bias)</strong></td>
<td>Low risk</td>
<td>“Randomization was performed using random block permutation method according to a computer-generated randomization list. The block length varied randomly (6, 12). Random allocation sequence was performed by a biostatistician. The detail of series was unknown by the study investigators.” Page 2 Soheilian 2009</td>
</tr>
<tr>
<td><strong>Allocation concealment (selection bias)</strong></td>
<td>Low risk</td>
<td>“Randomization was performed using random block permutation method according to a computer-generated randomization list. The block length varied randomly (6, 12). Random allocation sequence was performed by a biostatistician. The detail of series was unknown by the study investigators.” Page 2 Soheilian 2009</td>
</tr>
<tr>
<td><strong>Blinding of participants and personnel (performance bias)</strong></td>
<td>Low risk</td>
<td>“A sham laser procedure (20 seconds) was performed by aiming the laser beam on the macula for the eyes in the IVB and IVB/IVT groups. In the MPC group, a sham injection was done by a needleless syringe pressed against the conjunctiva. To keep the masking process, patients were prevented from seeing the syringes. All procedures were run by staff members other than the study investigators to preserve investigator masking. Best-corrected VA measurement and OCT were performed by certified examiners masked both to the randomization and to the findings of previous measurements.” Page 2-3 Soheilian 2009</td>
</tr>
<tr>
<td><strong>Blinding of outcome assessment (detection bias)</strong></td>
<td>Low risk</td>
<td>See above</td>
</tr>
<tr>
<td><strong>Incomplete outcome data (attrition bias)</strong></td>
<td>Unclear risk</td>
<td>There were 6 missing eyes out of 50 at 36 weeks in the IVB group and 12 out of 50 in the photocoagulation group and causes were not clearly unrelated to VA outcome, except for 2 deaths. In a subsequent publication in 2012 the authors reported 39 (78%) and 38 (76%) eyes in the two arms; 8 participants (12 eyes) missing were dead for causes unrelated to treatment, but other causes of death were not reported</td>
</tr>
</tbody>
</table>
**Soheilian 2007**  *(Continued)*

<table>
<thead>
<tr>
<th>Bias Type</th>
<th>Risk</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Selective reporting (reporting bias)</td>
<td>Low risk</td>
<td>The primary outcomes are continuous measures and no arbitrary cut-points were used</td>
</tr>
<tr>
<td>Other bias</td>
<td>High risk</td>
<td>There was an imbalance of baseline VA in the IVB and photocoagulation groups: 0.71 logMAR versus 0.55 logMAR. Although there was a potential unit of analysis issue (150 eyes of 129 patients, 16% of participants had both eyes included), comparisons were made in a marginal regression model (based on generalised estimating equation methods) adjusted for the baseline values and to eliminate any possible correlation effects between the 2 eyes of participants in bilateral enrolled cases. However, we could not take correlation into account when analysing dichotomous VA definitions.</td>
</tr>
<tr>
<td>Overall risk of bias</td>
<td>Unclear risk</td>
<td>High or unclear risk of bias for two items to a degree which we believe may influence effect estimates</td>
</tr>
</tbody>
</table>

**Turkoglu 2015**

<table>
<thead>
<tr>
<th>Methods</th>
<th>Prospective study, treatment in the better-seeing eye</th>
</tr>
</thead>
</table>
| Participants | Country: Turkey  
Number of people randomised: 70 participants (35 ranibizumab, 35 grid laser)  
Average age: 64.6 ± 8.2 years ranibizumab; 63.8 ± 7.4 years laser  
Sex: 21% male ranibizumab, 18% male laser  
Inclusion criteria:  
• evidence of CSME by means of FFA  
• at least 6 months of follow-up  
• no other systemic or ocular disease that might affect vision  
Exclusion criteria:  
• participants with a history of intravitreal injection and laser photocoagulation for proliferative diabetic retinopathy or CSME  
• participants with vitreous haemorrhage present at the time of recruitment or vitreous haemorrhage which developed after enrolment |
| Interventions | Focal or grid laser photocoagulation treatment was performed in 35 participants and laser settings, including power, spot size, duration and number of burns, were recorded  
35 participants received initial injection of ranibizumab 0.5 mg/0.05 mL. All participants of both groups received treatment in their better-seeing eye. After the induction phase, the intravitreal injections were administered if any of the following changes were observed: presence of visual acuity loss; persistent or recurrent subretinal or intraretinal fluid |
Continued

### Turkoglu 2015

**Outcomes**

Group comparisons of absolute scores and mean changes from baseline scores at 6-month visit were performed using analysis of the Turkish version of VFQ-25; it has modifications to adjust for Turkish culture and lifestyle.

### Notes

**Risk of bias**

<table>
<thead>
<tr>
<th>Bias</th>
<th>Authors' judgement</th>
<th>Support for judgement</th>
</tr>
</thead>
<tbody>
<tr>
<td>Random sequence generation (selection bias)</td>
<td>Unclear risk</td>
<td>No data available</td>
</tr>
<tr>
<td>Allocation concealment (selection bias)</td>
<td>Unclear risk</td>
<td>No data available</td>
</tr>
<tr>
<td>Blinding of participants and personnel (performance bias)</td>
<td>Unclear risk</td>
<td>No data available</td>
</tr>
<tr>
<td>Blinding of outcome assessment (detection bias)</td>
<td>Unclear risk</td>
<td>No data available</td>
</tr>
<tr>
<td>Incomplete outcome data (attrition bias)</td>
<td>Low risk</td>
<td>No loss to follow-up reported</td>
</tr>
<tr>
<td>Selective reporting (reporting bias)</td>
<td>Unclear risk</td>
<td>No specific statement nor protocol available</td>
</tr>
<tr>
<td>Other bias</td>
<td>Low risk</td>
<td>No other bias identified</td>
</tr>
<tr>
<td>Overall risk of bias</td>
<td>Unclear risk</td>
<td>Most items not reported</td>
</tr>
</tbody>
</table>

**Wiley 2016**

**Methods**

Randomised, double-masked, 3-period, 2-treatment crossover design with 4 treatment sequence patterns. Each of 3 12-week periods consisted of 3 intravitreous injections of ranibizumab (0.3 mg) or bevacizumab (1.25 mg), given every 4 weeks, with evaluation of the treatment period 4 weeks after the third dose (i.e. weeks 12, 24, and 36).

**Participants**

Country: USA and UK

Number of people randomised: 56 participants (62 eyes, 6 piz both eyes)

Average age: 62 years

Sex: 38.7% women

Inclusion criteria:

Eligible participants had type 1 or type 2 diabetes mellitus, were at least 18 years of age, and could enter one or both eligible eyes in the study. Principal eligibility criteria for a
study eye included: (1) presence of DME involving the centre of the macula, (2) ETDRS BCVA letter score of 78 to 24 (Snellen equivalent, 20/32-20/400), and (3) CSMT of 330 mm or more on Cirrus (Carl Zeiss Meditec, Inc, Dublin, CA) OCT
Exclusion criteria:
Eye included presence of factors or other conditions judged to impact the course of oedema or to preclude possible improvement in vision with treatment; PRP, focal or grid laser photocoagulation, or depot corticosteroid injection within the previous 3 months; ocular injection with an anti-VEGF agent within the previous 2 months; more than 4 injections with an anti-VEGF agent within the previous year; or prior vitrectomy. Potential participants were excluded for history of renal failure (requiring haemodialysis or renal transplantation) and for a measured systolic BP of more than 180 mmHg or a diastolic BP of more than 110 mmHg

Interventions
Each of 3 12-week periods consisted of 3 intravitreous injections of ranibizumab (0.3 mg) or bevacizumab (1.25 mg), given every 4 weeks, with evaluation of the treatment period 4 weeks after the third dose (i.e. weeks 12, 24, and 36)
Each study eye received 9 monthly injections over the course of the trial, according to a pattern of treatments determined by 1 of 4 randomly assigned sequences: R-R-B (n = 17), R-B-B (n = 15), B-B-R (n = 16), or B-R-R (n = 14), where R indicates a series of 3 consecutive ranibizumab injections, and B represents a series of 3 consecutive bevacizumab injections. Study eyes meeting predefined criteria for significant worsening of DME at week 12 or later could receive focal or grid laser photocoagulation. Fellow eyes in participants only enrolling 1 eye could receive any necessary ocular treatment

Outcomes
The primary outcome: mean change in BCVA from baseline, estimated for a 3-month dosing period in a linear mixed-effects model
The main prespecified secondary outcome was the change in CRT, measured as OCT CSMT, estimated for a 3-month dosing period using the linear mixed-effects model

Notes
June 2012 and January 2014

Risk of bias

<table>
<thead>
<tr>
<th>Bias</th>
<th>Authors’ judgement</th>
<th>Support for judgement</th>
</tr>
</thead>
</table>
| Random sequence generation (selection bias) | Low risk | “Participants were assigned to 1 of the 4 treatment sequences using a randomization list generated by the Data and Statistical Coordinating Center before study initiation, with balance after every 12 enrollments. The list was provided to unmasked pharmacists at each site, who confirmed a valid participant identification code before dispensing study treatment. Both clinical sites used the same randomised list, but selected treatment assignments from opposite ends. For participants entering both eyes in the trial, the right eye was assigned randomly as above to 1 of the
### Allocation concealment (selection bias)

**Low risk**

Participants were assigned to 1 of the 4 treatment sequences using a randomization list generated by the Data and Statistical Coordinating Center before study initiation, with balance after every 12 enrollments. The list was provided to unmasked pharmacists at each site, who confirmed a valid participant identification code before dispensing study treatment. Both clinical sites used the same randomised list, but selected treatment assignments from opposite ends. For participants entering both eyes in the trial, the right eye was assigned randomly as above to 1 of the 4 treatment sequences, and the left eye was assigned automatically to the sequence with the inverse schedule (for example, B-R-R in the right eye and R-B-B in the left eye). Page 2

### Blinding of participants and personnel (performance bias)

**Low risk**

"Participants and investigators were masked to treatment. Site staff collecting study data, including research coordinators, technicians, and photographers, were also masked."

### Blinding of outcome assessment (detection bias)

**Low risk**

"Site staff collecting study data, including research coordinators, technicians, and photographers, were also masked."

### Incomplete outcome data (attrition bias)

**Unclear risk**

"One participant with a single eye assigned to the B-B-B group withdrew after the week 4 visit after a hemorrhagic stroke. All remaining participants completed the study, including the week 12, 24, and 36 visits, and were included in this analysis."

### Selective reporting (reporting bias)

**Low risk**

VA (no SD), OCT (no SD) data and harms adequately reported

### Other bias

**Low risk**

Six out of 56 participants had both eyes included in analyses

### Overall risk of bias

**Low risk**

Low risk of bias for most items
Abbreviations
BCVA: best-corrected visual acuity
BP: blood pressure
CRT: central retinal thickness
CSFT: central subfield thickness
CSMO: clinically significant macular oedema
CSMT: central subfield mean thickness
DMO: diabetic macular oedema (DME: US spelling edema)
ECG: electrocardiogram
EQ-5D: EuroQol 5D
ETDRS: Early Treatment Diabetic Retinopathy Study
FAZ: foveal avascular zone
FFA: fundus fluorescein angiography
GLD: greatest linear dimension
HbA1c: glycated haemoglobin
IOP: intraocular pressure
ITT: intention-to-treat
iv: intravenous
IV: intravitreal injection
IVB: intravitreal bevacizumab
IVT: intravitreal triamcinolone
LOCF: last observation carried forward
logMAR: log of the Minimum Angle of Resolution
NEI VFQ-25: National Eye Institute Visual Function Questionnaire-25
OCT: optical coherence tomography
PDR: proliferative diabetic retinopathy
PFCL: perifoveal capillary loss
PRP: panretinal photocoagulation
RCT: randomised controlled trial
SD: standard deviation
SD-OCT: spectral-domain optical coherence tomography
TD-OCT: time-domain optical coherence tomography
VA: visual acuity
VEGF: vascular endothelial growth factor

Characteristics of excluded studies  [ordered by study ID]

<table>
<thead>
<tr>
<th>Study</th>
<th>Reason for exclusion</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ahmadieh 2013</td>
<td>Not an RCT</td>
</tr>
<tr>
<td>CRFB002DFR08 (LUDIC)</td>
<td>Single-arm study</td>
</tr>
<tr>
<td>CRFB002DGB14 (RELIGHT)</td>
<td>Single-arm study</td>
</tr>
<tr>
<td>CRFB002DNO02 (PTIMAL)</td>
<td>Single-arm study</td>
</tr>
<tr>
<td>DRCRnet 2007</td>
<td>Follow-up at 12 weeks only</td>
</tr>
</tbody>
</table>
DRCRnet 2011  
Follow-up at 14 weeks only. RCT comparing ranibizumab (2 injections), triamcinolone (1 injection) to sham in participants with DMO undergoing grid and panretinal laser photocoagulation.

DRCRnet 2012  
Follow-up of DRCRnet 2010 comparing prompt to deferred laser in participants treated for ranibizumab for DMO: does not report on comparison of ranibizumab with laser.

Faghihi 2008  
Follow-up at 16 weeks only.

NCT02985619 (BEVATAAC)  
Triamcinolone as comparator, unpublished study.

Paccola 2008  
Single injection of intravitreal triamcinolone acetonide (4 mg/0.1 mL) compared to single injection of intravitreal bevacizumab (1.5 mg/0.06 mL). Duration: 24 weeks.

Solaiman 2010  
Single intravitreal injection of bevacizumab (inadequate dose); follow-up 6 months.

Zehetner 2013  
Physiological study of anti-VEGF levels only.

**Abbreviations**
DMO: diabetic macular oedema  
RCT: randomised controlled trial  
VEGF: vascular endothelial growth factor

**Characteristics of studies awaiting assessment**  
*[ordered by study ID]*

**Chen 2016**

| Methods | Allocation: randomised  
|         | Endpoint classification: efficacy study  
|         | Intervention model: parallel assignment  
|         | Masking: unclear  
|         | Primary purpose: treatment |

| Participants | 42 (72 eyes); country: China  
|             | Male 55%, average age 60 years |

| Interventions | Experimental: ranibizumab 0.5 mg; one week after ranibizumab, the participants received grid photocoagulation intervention: macular area C-shaped manner, spot size 100 µm to 200 µm, 1 spot width apart, level I to II power and exposure time 0.1 second  
|               | Active comparator: laser only |

| Outcomes | Primary outcome (time frame: 3 and 6 months):  
|          | • Improvement of VA (2 or more lines on a letter chart, but not ETDRS)  
|          | • Ranibizumab plus laser: 28/36 (78%)  
|          | • Laser only: 18/36 (50%)  
|          | Secondary outcome: |
### Chen 2016

- Reduction in retinal thickening based on OCT
- Baseline: ranibizumab plus laser: 487 (SD: 85) micron; laser 480 (SD: 83) micron
- 6 months: ranibizumab plus laser: 246 (SD: 26) micron; laser: 370 (SD: 36) micron

**Notes**
- Articles in Chinese
- Authors contacted to obtain additional data

### Fouda 2017

**Methods**
- Allocation: randomised study but no other information reported
- Endpoint classification: efficacy study
- Intervention model: parallel assignment
- Masking: unclear
- Primary purpose: treatment

**Participants**
- 70 eyes randomised (42 participants); country: Egypt
- Male % not reported, average age 55 years

**Interventions**
- Experimental: ranibizumab 0.5 mg; aflibercept 2 mg

**Outcomes**
- Primary outcome (time frame: 12 months): VA and CRT (cut-off and instrument not specified)
- Decimal VA used: no significant difference between ranibizumab and aflibercept
- Reduction in retinal thickening based on OCT: no significant difference between ranibizumab and aflibercept
- Number of re-injections after the loading dose: 2.62 (SD 0.68) aflibercept, 3.03 (SD 0.95) ranibizumab (P = 0.02)

**Notes**
- Authors contacted to obtain additional data

### Huang 2016

**Methods**
- Allocation: quasi-randomised, participants with even visit numbers were allocated to intervention group, whereas participants with odd visit numbers were allocated to control group
- Endpoint classification: efficacy study
- Intervention model: parallel assignment
- Masking: unclear
- Primary purpose: treatment

**Participants**
- 78; country: China
- Male 55%, average age 51 to 54 years

**Interventions**
- Experimental: ranibizumab 0.5 mg
- Active comparator: argon laser 532µm, green, 50 µm to 100 µm spots, intensity 100 mW to 200 mW and exposure time 0.1 second

**Outcomes**
- Primary outcome (time frame: 3 and 6 months):
  - Improvement of VA (cut-off and instrument not specified)
  - Ranibizumab plus laser: 24/41 (59%)
  - Laser only: 11/27 (30%)

Secondary outcome:
### Huang 2016

(Continued)

- Reduction in retinal thickening based on OCT
- Baseline: ranibizumab: 401 (SD: 39) micron; laser: 387 (SD: 31) micron
- 6 months: ranibizumab: 289 (SD: 34) micron; laser: 320 (SD: 37) micron

**Notes**
- Article in Chinese
- Authors contacted to obtain additional data

### Jovanovic 2015

**Methods**
- Allocation: randomised
- Endpoint classification: efficacy study
- Intervention model: parallel assignment
- Masking: unclear
- Primary purpose: treatment

**Participants**
- 72 (120 eyes); country: Poland
- Average age: 60 years
- Inclusion criteria: severe DMO that affects the fovea, reduction in VA and/or metamorphopsia, diffuse oedema with or without cystic oedema (confirmed by fluorescein angiography and by OCT), CRT \( \geq 300 \) µm, the absence of hard lipid exudates in the form of plaque in the subfoveal region, no prior laser treatment, no prior VEGF inhibitor treatment, and no previous intravitreal or subtenonian corticosteroid administration. The exclusion criteria were: high risk and advanced proliferative DR (PDR), the presence of other eye diseases that could affect VA, prior eye surgeries

**Interventions**
- Experimental: bevacizumab 1.25 mg, one or more injections, with or without laser photocoagulation after 4 to 6 weeks depending on clinical response
- Active comparator: laser only

**Outcomes**
- Primary outcome: not specified
- Outcomes reported: VA (logMAR), CRT based on OCT
- Quantitative results are available, but not at specified follow-up times and by number of injection and combination rather than by assigned subgroup

**Notes**
- Sponsor: none reported
- Authors contacted to obtain additional information

### NCT00387582

**Methods**
- Allocation: randomised
- Endpoint classification: efficacy study
- Intervention model: parallel assignment
- Masking: open-label
- Primary purpose: treatment

**Participants**
- 49, country: USA

---

Anti-vascular endothelial growth factor for diabetic macular oedema: a network meta-analysis (Review)

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### NCT00387582 (Continued)

<table>
<thead>
<tr>
<th>Interventions</th>
<th>Experimental: I</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>• Lucentis injections for the first 3 months of the study and then according to the protocol for the duration of the trial</td>
</tr>
<tr>
<td></td>
<td>Active comparator: II</td>
</tr>
<tr>
<td></td>
<td>• Argon laser treatment at enrolment and then according to the protocol for the duration of the study</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Outcomes</th>
<th>Primary outcome (time frame: 6 and 12 months):</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>• Prevention of vision loss at 1 year as evidenced by ETDRS VA</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Secondary outcome:</th>
</tr>
</thead>
<tbody>
<tr>
<td>• Reduction in retinal thickening based on OCT</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Notes</th>
<th>Completion Date: February 2009 (No Results)</th>
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</thead>
<tbody>
<tr>
<td></td>
<td>Author contact not found</td>
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<tr>
<td></td>
<td>Sponsor: Rocky Mountain Retina Consultants</td>
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<td></td>
<td>Collaborator: Genentech, Inc.</td>
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### NCT00997191 (IBeTA)

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<tbody>
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<td>Endpoint classification: safety/efficacy study</td>
</tr>
<tr>
<td></td>
<td>Intervention model: parallel assignment</td>
</tr>
<tr>
<td></td>
<td>Masking: open-label</td>
</tr>
<tr>
<td></td>
<td>Primary purpose: treatment</td>
</tr>
</tbody>
</table>

| Participants | 12; country: Brazil  |

<table>
<thead>
<tr>
<th>Interventions</th>
<th>Procedure: laser photocoagulation</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Drug: intravitreal triamcinolone</td>
</tr>
<tr>
<td></td>
<td>Drug: intravitreal bevacizumab</td>
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<table>
<thead>
<tr>
<th>Outcomes</th>
<th>Primary outcome (time frame: 1 year):</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>• BCVA</td>
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</table>

<table>
<thead>
<tr>
<th>Secondary outcomes:</th>
</tr>
</thead>
<tbody>
<tr>
<td>• macular mapping test</td>
</tr>
<tr>
<td>• multifocal electroretinogram</td>
</tr>
<tr>
<td>• CRT</td>
</tr>
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<table>
<thead>
<tr>
<th>Notes</th>
<th>Completion date: November 2011 (no results)</th>
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<tbody>
<tr>
<td></td>
<td>Author could not be contacted</td>
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<tr>
<td></td>
<td>Sponsor: University of Sao Paulo</td>
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<tr>
<td></td>
<td>Collaborator: Fundacao de Amparo a Pesquisa do Estado de Sao Paolo</td>
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</table>
**NCT01445899 (MATISSE)**

**Methods**
- Allocation: randomised
- Endpoint classification: safety/efficacy study
- Intervention model: parallel assignment
- Masking: double-masked (participant, caregiver, investigator)

**Participants**
- 264; countries: USA, Israel

**Interventions**
- Drug: PF-04523655 (Stratum I)
- Drug: PF-04523655 and ranibizumab
- Drug: ranibizumab
- Drug: PF-04523655 (Stratum II)

**Outcomes**
- Primary outcomes:
  - Safety and dose-limiting toxicities (Stratum I): to determine the safety and dose-limiting toxicities of a single intravitreal (IVT) injection of PF-04523655 in people with low vision
  - Pharmacokinetics (Stratum I): to determine the pharmacokinetics (PK) of a single IVT injection of PF-04523655 in people with low vision
  - Safety and tolerability (Stratum II): to evaluate the safety and tolerability of PF-04523655 alone and in combination with ranibizumab in participants with DMO
  - Efficacy (Stratum II): to evaluate the ability of PF-04523655 alone and in combination with ranibizumab to improve VA compared to ranibizumab alone in people with DMO

**Notes**
- Completion date: November 2013 (no results)
- Author contact not found
- Sponsor: Quark Pharmaceuticals
- Consider putting in excluded studies

---

**NCT01565148 (IDEAL)**

**Methods**
- Allocation: randomised
- Endpoint classification: safety/efficacy study
- Intervention model: factorial assignment
- Masking: open-label

**Participants**
- 208; country: USA

**Interventions**
- Experimental Group 1: drug: iCo-007 350 µg as an intravitreal injection at baseline followed by another iCo-007 dose (350 µg) at month 4
- Experimental Group 2: drug: iCo-007 700 µg as an intravitreal injection at baseline followed by another iCo-007 dose (700 µg) at month 4
- Experimental Group 3: drug: iCo-007 350 µg as an intravitreal injection at baseline followed 7 days later by laser photocoagulation. At month 4, intravitreal injection of iCo-007 (350 µg) will be given as mandatory treatment. If the eye also meets retreatment criteria, it will also receive the second laser photocoagulation
- Experimental Group 4: drug: ranibizumab 0.5 mg intravitreal injection at baseline followed by iCo-007 350 µg intravitreal injection 2 weeks later; re-treatment with ranibizumab 0.5 mg mandatory at month 4 followed by iCo-007 350 µg 2 weeks later
NCT01565148 (IDEAL)

Outcomes

Primary outcome:
- Change in VA from baseline to month 8

Secondary outcomes:
- Number of participants in a given study arm experiencing the same drug-related serious adverse event as a measure of safety and tolerability
- Safety of repeated iCo-007 intravitreal injections in treatment of people with DMO as monotherapy and in combination with ranibizumab or laser photocoagulation. Serious consideration will be given if 2 or more participants in a particular treatment arm experience the same drug-related serious adverse event
- Change in VA from baseline to month 12
- Change in retinal thickness measured by OCT from baseline to month 8
- Change in retinal thickness measured by OCT from baseline to month 12
- Duration of iCo-007 treatment effect during the 12-month follow-up period as measured by VA and OCT thickness
- Peak plasma concentration (Cmax) of iCo-007 after multiple injections

Notes

Study has passed its completion date and status has not been verified in more than 2 years on ClinicalTrials.gov
Author contacted
Sponsor: Quan Dong Nguyen
Collaborators: Juvenile Diabetes Research Foundation, iCo Therapeutics Inc

Abbreviations

BCVA: best-corrected visual acuity
CFST: central subfield macular thickness
CRT: central retinal thickness
DMO: diabetic macular oedema
EQ-5D: EuroQol 5D
ETDRS: Early Treatment Diabetic Retinopathy Study
OCT: optical coherence tomography
PRN: pro re nata
SD: standard deviation
TE: treat and extend
VA: visual acuity
VEGF: vascular endothelial growth factor
VFQ-25: Visual Function Questionnaire 25-item

Characteristics of ongoing studies [ordered by study ID]

ChiCTR-TRC-12002417

Trial name or title
A randomised controlled trial to compare the efficacy and safety of 1) macular laser vs 2) repeated intravitreal bevacizumab vs 3) combined repeated intravitreal bevacizumab with macular laser for diabetic macular edema

Methods
Parallel group RCT

Participants
People with type 2 diabetes and DMO
Interventions

Group 1 (Control): macular laser photocoagulation performed every 4 months unless the deferral criteria are met. Group 2: intravitreal bevacizumab injections (1.25 mg each) given at 0, 1, 2 months and repeated en bloc every 4 months unless the deferral criteria are met. Group 3: Intravitreal bevacizumab injections (1.25 mg each) given at 0, 1, 2 months, followed by macular laser photocoagulation at month 3; and repeated en bloc every 4 months unless the deferral criteria are met.

Outcomes

BCVA at 2 years

Starting date

Unknown; trial registered 13 August 2013

Contact information

Joyce Kung (joycekung@cuhk.edu.hk); Carmen Chan (kmcc2001@hotmail.com)

Notes

Status checked on Chictr.Org.Cn on 1 December 2016. Author contacted

NCT01635790 (BRDME)

Trial name or title

Comparing the effectiveness and costs of bevacizumab to ranibizumab in participants with diabetic macular edema (BRDME)

Methods

Parallel group RCT

Participants

246 people with DMO

Interventions

Ranibizumab compared to bevacizumab

Outcomes

From clinical trials record:
Primary outcome:
- change in BCVA in the study eye from baseline to month 6 (designated as safety issue: no)
Secondary outcome measures:
- proportion of participants with a gain or loss of 15 letters or more at 6 months compared to baseline BCVA (designated as safety issue: no)
- change in leakage on fluorescein angiography, baseline compared to 6 month exit visit (designated as safety issue: no)
- change in foveal thickness (central retinal area) by OCT, 6 month exit visit compared to baseline (designated as safety issue: no)
- total number of adverse events that occurred during the 6 month study, with secondary a classification of the types of adverse events (designated as safety issue: yes)
- costs per quality adjusted life-year of the 2 treatments (time frame: 6 months; designated as safety issue: no), results will be based on the use of standardised health questionnaires (EQ-5D or Health Utility Index Mark 3)
- proportion of participants with a BCVA of 20/40 or more at 6 months compared to baseline BCVA (designated as safety issue: no)

http://clinicaltrials.gov/show/NCT01635790

Starting date

Study start date: June 2012
Estimated primary completion date: June 2016 (Final data collection date for primary outcome measure)
### NCT01635790 (BRDME)  (Continued)

<table>
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<tr>
<th>Contact information</th>
<th>Reinier O Schlingemann (<a href="mailto:r.schlingemann@amc.uva.nl">r.schlingemann@amc.uva.nl</a>); Monique Wezel (<a href="mailto:m.wazel@amc.uva.nl">m.wazel@amc.uva.nl</a>)</th>
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### NCT02194634

<table>
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<tr>
<th>Trial name or title</th>
<th>Safety and Efficacy Study of Conbercept in Diabetic Macular Edema (DME) (Sailing)</th>
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<tbody>
<tr>
<td>Methods</td>
<td>Parallel group RCT</td>
</tr>
<tr>
<td>Participants</td>
<td>type 1 or type 2 diabetes mellitus with DMO</td>
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</tbody>
</table>
| Interventions       | Experimental: Conbercept treatment group  
Conbercept injection and sham laser treatment at day 0 for 1st time, the investigators will decide whether the participants need to get repeated treatment according to monthly assessment  
Active Comparator: Laser treatment group  
Laser treatment and sham injection at day 0 for 1st time, the investigators will decide whether the repeated laser treatment is needed according to monthly results during the visit after 3 months |
| Outcomes            | Primary outcome measures:  
- Mean change from baseline in BCVA at month 12 (time frame: baseline and month 12) (designated as safety issue: no)  
- To compare mean change from baseline BCVA between treatment group and controlled group at month 12  
Secondary outcome measures:  
- Mean change from baseline in CRT between two groups (time frame: baseline and month 12 (designated as safety issue: no)  
- To compare mean change from baseline CRT between two groups at month 12  
- Safety (e.g. incidence of adverse events) of Conbercept ophthalmic injection (time frame: 12 months) (designated as safety issue: yes)  
- To assess safety parameters during the study, such as incidence of adverse events, incidence of adverse drug reactions etc |
| Starting date       | July 2014  
Estimated Study Completion Date: September 2017  
Estimated Primary Completion Date: December 2016 (Final data collection date for primary outcome measure) |
| Contact information | Chengdu Kanghong Biotech Co., Ltd. |
| Notes               | This study is recruiting participants. (Status checked on ClinicalTrials.Gov on 5 December 2016) |
### NCT02259088

<table>
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<tr>
<th>Trial name or title</th>
<th>A 12-month, Randomized, Efficacy and Safety Study of 0.5 mg Ranibizumab vs Laser in Chinese DME Patients</th>
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<tr>
<td>Methods</td>
<td>Parallel group RCT</td>
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<tr>
<td>Participants</td>
<td>Participants with DMO with BCVA score between 78 and 39</td>
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<tr>
<td>Interventions</td>
<td>Ranibizumab PRN versus laser</td>
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<tr>
<td>Outcomes</td>
<td>Primary outcome measures:&lt;br&gt;- Mean average BCVA change (time frame: 12 months)&lt;br&gt;- Mean average BCVA change from Month 1 through Month 12 compared to baseline.&lt;br&gt;Secondary Outcome Measures:&lt;br&gt;- Mean BCVA change by visit (time frame: 12 months)&lt;br&gt;- Mean BCVA change from baseline at each visit&lt;br&gt;- Mean change in CSFT (time frame: 12 months)&lt;br&gt;- Mean change in CSFT from baseline at each visit&lt;br&gt;- BCVA improvement of ( \geq 10 ) and ( \geq 15 ) letters (time frame: 12 months)&lt;br&gt;- Proportion of participants achieving BCVA improvement of ( \geq 10 ) and ( \geq 15 ) letters from baseline to Month 12&lt;br&gt;- BCVA loss of (&lt; 10 ) and (&lt; 15 ) letters (time frame: 12 months)&lt;br&gt;- Proportion of participants with BCVA loss of (&lt; 10 ) and (&lt; 15 ) letters from baseline to Month 12&lt;br&gt;- VA ( \geq 73 ) letters (time frame: 12 months)&lt;br&gt;- Proportion of participants with VA ( \geq 73 ) letters (approximate 20/40 Snellen chart equivalent) at Month 12&lt;br&gt;- Mean average BCVA change after month 3 (time frame: 12 months)&lt;br&gt;- Mean average BCVA change from Month 4 to Month 12 compared to Month 3&lt;br&gt;- Mean change in patient-reported visual functioning scale (time frame: 6 and 12 months)&lt;br&gt;Mean change in the patient-reported visual functioning through VFQ-25 composite and subscale scores at Month 6 and Month 12 compared to baseline.&lt;br&gt;- Adverse events (time frame: 12 months)&lt;br&gt;- To evaluate the safety of 0.5 mg ranibizumab intravitreal injections relative to laser photocoagulation, as assessed by the type, frequency and severity of ocular and non-ocular adverse events over 12 months.&lt;br&gt;- Evaluation of treatment patterns (time frame: 12 months)&lt;br&gt;- To evaluate the number of re-treatments and retreatment patterns (assessed by time of stabilization/time to re-initiation of treatment) in participants treated with 0.5 mg ranibizumab</td>
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<td>Estimated study completion date: January 2017&lt;br&gt;Estimated primary completion date: January 2017 (final data collection date for primary outcome measure)</td>
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<td>Notes</td>
<td>This study is ongoing, but not recruiting participants. (Status checked on ClinicalTrials.gov on 5 December 2016)</td>
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## NCT02348918

### Trial name or title
A Phase 2 Randomized, Controlled, Double-Masked, Multicenter Clinical Trial Designed to Evaluate the Safety and Exploratory Efficacy of Luminate® (ALG-1001) as Compared to Avastin® and Focal Laser Photocoagulation in the Treatment of Diabetic Macular Edema

### Methods
Parallel group RCT

### Participants
DMO participants

### Interventions
**Experimental: Luminate 1.0 mg group**
- Luminate 1.0 mg intravitreal injection administered at baseline (Day 0), 4 weeks and 8 weeks with PRN Luminate injection at week 20 for a total of at least 3 and no more than 4 Luminate injections. Sham injections will be performed at weeks 12 and 16 and may also be performed at week 20 if PRN Luminate is not required; sham laser treatment will be administered at baseline and at 16 weeks

**Experimental: Luminate 2.0 mg group**
- Luminate 2.0 mg intravitreal injection administered at baseline (Day 0), 4 weeks and 8 weeks with PRN Luminate injection at week 20 for a total of at least 3 and no more than 4 Luminate injections. Sham injections will be performed at weeks 12 and 16 and may also be performed at week 20 if PRN Luminate is not required; sham laser treatment will be administered at baseline and at 16 weeks

**Experimental: Luminate 3.0 mg group**
- Luminate 3.0 mg intravitreal injection administered at baseline (Day 0), 4 weeks and 8 weeks with PRN Luminate injection at week 20 for a total of at least 3 and no more than 4 Luminate injections. Sham injections will be performed at weeks 12 and 16 and may also be performed at week 20 if PRN Luminate is not required; sham laser treatment will be administered at baseline and at 16 weeks

**Active Comparator: Avastin® group**
- Avastin 1.25 mg intravitreal injection administered at baseline (Day 0), 4 weeks and 8 weeks with PRN Avastin injection at weeks 12, 16, or 20 for a total of at least 3 and up to 6 Avastin injections. Sham injections may be performed at weeks 12, 16, and 20 if PRN Avastin is not required; sham laser treatment will be administered at baseline and at 16 weeks

**Active Comparator: focal laser photocoagulation group**
- Focal laser photocoagulation performed at baseline (Day 0) with possible PRN laser retreatment at week 16. Sham intravitreal injections will be performed at baseline (Day 0), 4 weeks, 8 weeks, 12 weeks, 16 weeks and 20 weeks

### Outcomes
**Primary outcome measures:**
- Change in OCT CSFT at Week 24 (time frame: 24 weeks)
- The primary efficacy outcome is OCT CSFT at Week 24 as compared to baseline

**Secondary outcome measures:**
- Change in BCVA at Week 24 (time frame: 24 weeks)
- Secondary efficacy outcome is BCVA changes at Week 24 as compared to baseline

### Starting date
October 2014

Estimated study completion date: March 2016

Estimated primary completion date: December 2015 (final data collection date for primary outcome measure)

### Contact information
Allegro Ophthalmics, LLC

### Notes
This study is recruiting participants; (status checked on ClinicalTrials.gov on 5 Dec. 2016)
### NCT02645734

**Trial name or title**  The Effect of Bevacizumab and Ziv-aflibercept in Diabetic Macular Edema  

**Methods**  Parallel group RCT  

**Participants**  DMO participants  

**Interventions**  
- Active comparator: injection intravitreous bevacizumab  
- Active comparator: injection ziv-aflibercept at dose of 1.25 mg  
- Active comparator: injection ziv-aflibercept at dose of 2.5 mg  

**Outcomes**  
- Primary outcome measures:  
  - VA (time frame: until 6 months)  
- Secondary outcome measures:  
  - CSFT (time frame: until 6 months)  

**Starting date**  
- Study first received: 2 January 2016  
- Last updated: 4 January 2016  
- Estimated primary completion date: February 2016 (final data collection date for primary outcome measure)  

**Contact information**  
- Zahra Rabbani Khah, Shahid Beheshti Medical University  

**Notes**  
- This study is recruiting participants; (status checked on ClinicalTrials.gov on 5 December 2016)  

### NCT02699450

**Trial name or title**  A Phase 2 Study of RO6867461 in Participants With Center-Involving Diabetic Macular Edema (CI-DME) (BOULEVARD)  

**Methods**  Parallel group RCT  

**Participants**  DMO  

**Interventions**  
- Experimental: RO6867461 1.5 mg  
  - Participants will receive 1.5 milligrams (mg) RO6867461 intravitreal (IVT) every 4 weeks up to Week 20, followed by 1 sham administration at Week 24.  
- Experimental: RO6867461 6 mg  
  - Participants will receive 6 mg RO6867461 IVT every 4 weeks up to Week 20, followed by 1 sham administration at Week 24.  
- Active Comparator: Ranibizumab 0.3 mg  
  - Participants will receive 0.3 mg ranibizumab IVT every 4 weeks up to Week 24  

**Outcomes**  
- Primary outcome measures:  
  - Mean change from baseline in BCVA at week 24 using ETDRS modified charts (time frame: baseline, Week 24)  
  - Apparent plasma clearance of RO6867461 (time frame: pre-dose on Days 1, 28, 84, 140, and 168; post-dose on Days 7, 182, and 196 or early termination)  
  - Apparent plasma volume of RO6867461 (time frame: pre-dose on Days 1, 28, 84, 140, and 168; post-dose on Days 7, 182, and 196 or early termination)  

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Secondary Outcome Measures:
- Percentage of participants gaining ≥ 15 letters from baseline BCVA at Week 24 (time frame: baseline, and Week 24)
- Percentage of participants with BCVA ≥ 69 letters (20/40 or better) at Week 24 (time frame: Week 24)
- Percentage of participants with BCVA ≥ 84 letters (20/20 or better) at Week 24 (time frame: Week 24)
- Mean change from baseline in BCVA at Week 28 (time frame: baseline, and Week 28)
- Mean change from baseline in foveal centre point thickness at Week 24 and 28, as measures by spectral domain OCT (SD-OCT) (time frame: baseline, Weeks 24 and 28)
- Mean change from baseline in mean CSFT at Week 24 and 28, as measures by SD-OCT (time frame: baseline, Weeks 24 and 28)
- Percentage of participants with resolution of subretinal and intraretinal fluid at Week 24 and 28, as measures by SD-OCT (time frame: Weeks 24 and 28)
- Percentage of participants with resolution of leakage at the macula at Week 24, as measures by FFA (time frame: Week 24)
- Change from baseline in the size of the foveal avascular zone at Week 24, as measures by FFA (time frame: baseline and Week 24)
- Change from baseline in plasma levels of vascular endothelial growth factor (VEGF) (time frame: baseline, Weeks 1, 4, 12, 24, 26, and 28 or early termination)
- Change from baseline in plasma levels of angiopoietin-2 (Ang-2) (time frame: baseline, Weeks 1, 4, 12, 24, 26, and 28 or early termination) (designated as safety issue: no)
- Maximum observed plasma concentration (Cmax) of RO6867461 (time frame: pre-dose on Days 1, 28, 84, 140, and 168; post-dose on Days 7, 182, and 196 or early termination)
- Area under the plasma concentration-time curve from time zero to extrapolated infinite time [AUC (0-inf)] of RO6867461 (time frame: pre-dose on Days 1, 28, 84, 140, and 168; post-dose on Days 7, 182, and 196 or early termination)
- Area under the plasma concentration-time curve from time zero to end of dosing interval [AUC (0-tau)] of RO6867461 (time frame: pre-dose on Days 1, 28, 84, 140, and 168; post-dose on Days 7, 182, and 196 or early termination)
- Time to reach maximum observed plasma concentration (Tmax) of RO6867461 (time frame: pre-dose on Days 1, 28, 84, 140, and 168; post-dose on Days 7, 182, and 196 or early termination)
- Plasma decay half-life (t1/2) of RO6867461 (time frame: pre-dose on Days 1, 28, 84, 140, and 168; post-dose on Days 7, 182, and 196 or early termination)
- Number of participants with adverse events (time frame: baseline up to Week 28 or early termination)
- Number of participants with anti-RO6867461 antibodies (time frame: baseline, Weeks 1, 4, 12, 20, 24, 26, and 28 or early termination)

Starting date | March 2016
---|---
Estimated Study Completion Date | October 2017
Estimated Primary Completion Date | October 2017 (final data collection date for primary outcome measure)

Contact information | Hoffmann-La Roche

Notes | This study is recruiting participants; (status checked on ClinicalTrials.gov on 5 December 2016)
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<td><strong>Methods</strong></td>
</tr>
<tr>
<td><strong>Participants</strong></td>
</tr>
<tr>
<td><strong>Interventions</strong></td>
</tr>
</tbody>
</table>
| **Outcomes** | Primary outcome measures:  
• Change from baseline in BCVA measured by the ETDRS letter score at week 12 (time frame: baseline to week 12)  
Secondary outcome measures:  
• Change from baseline in CSFT at week 12 (time frame: baseline to week 12)  
• Change from baseline in CSFT at week 36 (time frame: baseline to week 36)  
• Proportion of participants with a ≥2-step improvement in Diabetic Retinopathy Severity Scale from baseline at week 12 (time frame: baseline to week 12)  
• Proportion of participants with a ≥2-step improvement in Diabetic Retinopathy Severity Scale from baseline at week 36 (time frame: baseline to week 36)  
• Change from baseline in BCVA measured by the ETDRS letter score at week 36 (time frame: baseline to week 36)  
• Proportion of participants with no retinal and/or subretinal fluid at week 12 (time frame: baseline to week 12)  
• Proportion of participants with no retinal and/or subretinal fluid at week 36 (time frame: baseline to week 36)  
• Time to no retinal and/or subretinal fluid at week 12 (time frame: baseline to week 12)  
• Time to no retinal and/or subretinal fluid at week 36 (time frame: baseline to week 36) |
| **Starting date** | March 2016 |
| **Estimated study completion date** | October 2017 |
| **Estimated primary completion date** | April 2017 (final data collection date for primary outcome measure) |
| **Contact information** | Regeneron Pharmaceuticals |
| **Notes** | This study is on ongoing, but not recruiting participants; (status checked on ClinicalTrials.gov on 5 December 2016) |

**Abbreviations**

BCVA: best-corrected visual acuity  
CSFT: central subfield thickness  
CRT: central retinal thickness  
CSMO: clinically significant macular oedema  
DMO: diabetic macular oedema (DME: US spelling edema)
ETDRS: Early Treatment Diabetic Retinopathy Study
FFA: fundus fluorescein angiography
IOP: intraocular pressure
NA: not available
OCT: optical coherence tomography
PDR: proliferative diabetic retinopathy
PRN: pro re nata (as required in the circumstances)
PRP: panretinal photocoagulation
RCT: randomised controlled trial
VA: visual acuity
DATA AND ANALYSES

Comparison 1. Ranibizumab versus laser photocoagulation at 6 to 12 months

<table>
<thead>
<tr>
<th>Outcome or subgroup title</th>
<th>No. of studies</th>
<th>No. of participants</th>
<th>Statistical method</th>
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<tbody>
<tr>
<td>1 Quality of life: NEI-VFQ composite score at 6 to 12 months</td>
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<td>412</td>
<td>Mean Difference (Fixed, 95% CI)</td>
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ADDITIONAL TABLES

Table 1. Reporting of all outcomes across studies

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<td>151</td>
<td>151</td>
<td>151</td>
<td></td>
</tr>
<tr>
<td>RESPONSE</td>
<td>203</td>
<td>203</td>
<td>202</td>
<td>237</td>
<td>237</td>
<td>237</td>
<td></td>
</tr>
<tr>
<td>RESTORE</td>
<td>343</td>
<td>343</td>
<td>343</td>
<td>299</td>
<td>345</td>
<td>345</td>
<td>345</td>
</tr>
</tbody>
</table>
Table 1. Reporting of all outcomes across studies  
(Continued)

<table>
<thead>
<tr>
<th>Study</th>
<th>Gain lines</th>
<th>Mean VA change</th>
</tr>
</thead>
<tbody>
<tr>
<td>RE-VEAL 2015</td>
<td>12</td>
<td>1074</td>
</tr>
<tr>
<td>RISE-RIDE</td>
<td>13</td>
<td>1131</td>
</tr>
<tr>
<td>Soheilian 2007</td>
<td>87</td>
<td>87</td>
</tr>
<tr>
<td>Turkoglu 2015</td>
<td>70</td>
<td>70</td>
</tr>
<tr>
<td>Wiley 2016</td>
<td>124</td>
<td>124</td>
</tr>
<tr>
<td>Total studies</td>
<td>17</td>
<td>5</td>
</tr>
<tr>
<td>Total participants</td>
<td>4031</td>
<td>1844</td>
</tr>
</tbody>
</table>

Numbers in the table are the total number of eyes for each study, as available by follow-up year (1 or 2) and outcome measure.

Table 2. Network structure: efficacy at 12 months

<table>
<thead>
<tr>
<th>Laser</th>
<th>Aflibercept</th>
<th>Bevacizumab</th>
<th>Pegaptanib</th>
<th>Ranibizumal deferred laser</th>
<th>Ranibizumal prompt laser</th>
<th>Sham</th>
<th>Overall</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gain 3+</td>
<td>12</td>
<td>4</td>
<td>5</td>
<td>3</td>
<td>8</td>
<td>1</td>
<td>3</td>
</tr>
<tr>
<td></td>
<td>1074</td>
<td>539</td>
<td>344</td>
<td>410</td>
<td>713</td>
<td>188</td>
<td>545</td>
</tr>
<tr>
<td>Mean VA change</td>
<td>13</td>
<td>4</td>
<td>7</td>
<td>3</td>
<td>11</td>
<td>1</td>
<td>6</td>
</tr>
<tr>
<td></td>
<td>1131</td>
<td>539</td>
<td>476</td>
<td>410</td>
<td>861</td>
<td>188</td>
<td>629</td>
</tr>
</tbody>
</table>

Anti-vascular endothelial growth factor for diabetic macular oedema: a network meta-analysis (Review)  
Copyright © 2017 The Cochrane Collaboration. Published by John Wiley & Sons, Ltd.
Table 2. Network structure: efficacy at 12 months (Continued)

<table>
<thead>
<tr>
<th>Mean CRT change</th>
<th>11</th>
<th>4</th>
<th>7</th>
<th>10</th>
<th>1</th>
<th>4</th>
<th>2</th>
<th>16</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>986</td>
<td>538</td>
<td>476</td>
<td>779</td>
<td>175</td>
<td>451</td>
<td>86</td>
<td>3491</td>
</tr>
</tbody>
</table>

QOL

<table>
<thead>
<tr>
<th>4</th>
<th>4</th>
</tr>
</thead>
<tbody>
<tr>
<td>838</td>
<td>838</td>
</tr>
</tbody>
</table>

For each efficacy outcome, numbers in the table are the total number of studies (upper line for each outcome) and the total number of eyes (lower line for each outcome), as available by treatment and measured at one year.

Table 3. Gain of 3 or more lines of visual acuity at 12 months: direct (upper-right triangle) and mixed (lower-left triangle) estimates

<table>
<thead>
<tr>
<th>Procedure</th>
<th>LASER</th>
<th>AFLI</th>
<th>BEVA</th>
<th>PEGA</th>
<th>RANI</th>
<th>RANI-DL</th>
<th>RANI-PL</th>
</tr>
</thead>
<tbody>
<tr>
<td>3.66 (2.79 to 4.79)</td>
<td>3.82 (2.61 to 5.58)</td>
<td>2.74 (1.34 to 5.61)</td>
<td>2.82 (1.82 to 4.38)</td>
<td>1.88 (1.31 to 2.70)</td>
<td>2.30 (1.74 to 3.03)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>2.47 (1.81 to 3.37)</td>
<td>2.47 (1.81 to 3.37)</td>
<td>0.68 (0.53 to 0.86)</td>
<td>0.69 (0.24 to 1.89)</td>
<td>1.14 (0.88 to 1.48)</td>
<td>0.51 (0.30 to 0.89)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>1.70 (0.58 to 4.94)</td>
<td>1.70 (0.58 to 4.94)</td>
<td>0.46 (0.16 to 1.34)</td>
<td>0.69 (0.24 to 1.89)</td>
<td>0.69 (0.24 to 1.89)</td>
<td>0.51 (0.30 to 0.89)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>2.76 (2.12 to 3.59)</td>
<td>2.76 (2.12 to 3.59)</td>
<td>0.75 (0.60 to 0.94)</td>
<td>1.11 (0.87 to 1.43)</td>
<td>1.62 (0.58 to 4.57)</td>
<td>1.88 (1.31 to 2.70)</td>
<td>2.30 (1.74 to 3.03)</td>
<td></td>
</tr>
<tr>
<td>2.02 (1.46 to 2.81)</td>
<td>2.02 (1.46 to 2.81)</td>
<td>0.55 (0.37 to 0.82)</td>
<td>0.82 (0.54 to 1.24)</td>
<td>1.19 (0.40 to 3.58)</td>
<td>1.88 (1.31 to 2.70)</td>
<td>2.30 (1.74 to 3.03)</td>
<td></td>
</tr>
<tr>
<td>2.33 (1.81 to 3.00)</td>
<td>2.33 (1.81 to 3.00)</td>
<td>0.64 (0.47 to 0.86)</td>
<td>0.94 (0.68 to 1.31)</td>
<td>1.37 (0.47 to 3.99)</td>
<td>1.88 (1.31 to 2.70)</td>
<td>2.30 (1.74 to 3.03)</td>
<td></td>
</tr>
<tr>
<td>0.87 (0.35 to 2.17)</td>
<td>0.87 (0.35 to 2.17)</td>
<td>0.24 (0.10 to 0.59)</td>
<td>0.35 (0.14 to 0.87)</td>
<td>0.51 (0.30 to 0.89)</td>
<td>0.32 (0.13 to 0.76)</td>
<td>0.37 (0.15 to 0.93)</td>
<td></td>
</tr>
</tbody>
</table>

P value for overall inconsistency = 0.883 in the network meta-analysis.

Values in the table are risk ratios and 95% confidence intervals. Values in bold are ones where the 95% confidence intervals does not include 1 (null effect).
### Table 4. Mean visual acuity change at 12 months: direct (upper-right triangle) and mixed (lower-left triangle) estimates

<table>
<thead>
<tr>
<th>LASER</th>
<th>direct</th>
<th>indirect</th>
<th>direct</th>
<th>indirect</th>
</tr>
</thead>
<tbody>
<tr>
<td>AF LI</td>
<td>0.07 (0.03 to 0.11)</td>
<td>0.04 (0.00 to 0.08)</td>
<td>0.02 (−0.05 to 0.01)</td>
<td>0.01 (−0.02 to 0.03)</td>
</tr>
<tr>
<td>BEV A</td>
<td>0.08 (0.05 to 0.11)</td>
<td>0.01 (−0.02 to 0.03)</td>
<td>0.00 (−0.04 to 0.05)</td>
<td>0.01 (−0.04 to 0.05)</td>
</tr>
<tr>
<td>PEGA</td>
<td>0.01 (−0.09 to 0.07)</td>
<td>0.09 (0.04 to 0.19)</td>
<td>0.00 (−0.02 to 0.06)</td>
<td>0.01 (−0.02 to 0.06)</td>
</tr>
<tr>
<td>RANI</td>
<td>0.19 (0.11 to 0.27)</td>
<td>0.01 (−0.04 to 0.05)</td>
<td>0.00 (−0.01 to 0.03)</td>
<td>0.01 (−0.04 to 0.05)</td>
</tr>
<tr>
<td>RANI-DL</td>
<td>0.28 (0.21 to 0.35)</td>
<td>0.20 (0.13 to 0.27)</td>
<td>0.20 (0.13 to 0.27)</td>
<td>0.20 (0.11 to 0.26)</td>
</tr>
</tbody>
</table>

| SHAM | 0.08 (0.03 to 0.13) | 0.08 (0.03 to 0.13) | 0.08 (0.03 to 0.13) | 0.08 (0.03 to 0.13) |

*P value for differences between direct and indirect estimates = 0.031 in the network meta-analysis.
P value for overall inconsistency = 0.665.

### Table 5. Mean central retinal thickness change at 12 months: direct (upper-right triangle) and mixed (lower-left triangle) estimates

<table>
<thead>
<tr>
<th>LASER</th>
<th>direct</th>
<th>indirect</th>
<th>direct</th>
<th>indirect</th>
</tr>
</thead>
<tbody>
<tr>
<td>AF LI</td>
<td>68 (43 to 94)</td>
<td>22 (−4 to 48)</td>
<td>68 (43 to 94)</td>
<td>22 (−4 to 48)</td>
</tr>
<tr>
<td>BEV A</td>
<td>68 (29 to 108)</td>
<td>132 (72 to 187)</td>
<td>68 (29 to 108)</td>
<td>132 (72 to 187)</td>
</tr>
<tr>
<td>RANI</td>
<td>39 (2 to 76)</td>
<td>1470 (95 to 196)</td>
<td>39 (2 to 76)</td>
<td>1470 (95 to 196)</td>
</tr>
<tr>
<td>RANI-DL</td>
<td>57 (−6 to 120)</td>
<td>18 (−40 to 76)</td>
<td>57 (−6 to 120)</td>
<td>18 (−40 to 76)</td>
</tr>
</tbody>
</table>

*P value for differences between direct and indirect estimates = 0.031 in the network meta-analysis.
Table 5. Mean central retinal thickness change at 12 months: direct (upper-right triangle) and mixed (lower-left triangle) estimates (Continued)

<table>
<thead>
<tr>
<th></th>
<th>Direct Estimates</th>
<th>Mixed Estimates</th>
</tr>
</thead>
<tbody>
<tr>
<td>RANI-PL</td>
<td>72 (−103 to −42)</td>
<td>−16 (−71 to 38)</td>
</tr>
<tr>
<td>−72.90 (−103 to −42)b</td>
<td>41 (−2 to 84)</td>
<td>123 (−67 to 179)</td>
</tr>
<tr>
<td>77 (18 to 137)</td>
<td>191 (127 to 256)</td>
<td>153 (97 to 208)</td>
</tr>
<tr>
<td></td>
<td>SHAM</td>
<td></td>
</tr>
<tr>
<td></td>
<td>123 (−68 to 13)</td>
<td>2 (−31 to 35)a</td>
</tr>
<tr>
<td></td>
<td>121 (−65 to 17)</td>
<td>2 (−31 to 35)</td>
</tr>
</tbody>
</table>

a P value for differences between direct and indirect estimates = 0.003.
b P value for differences between direct and indirect estimates = 0.044.
* P value for heterogeneity = 0.002; I² = 80% in the direct meta-analysis.
* P value for heterogeneity = 0.000; I² = 91% in the direct meta-analysis.
P value for overall inconsistency = 0.209 in the network meta-analysis.

Table 6. Network structure: safety at the longest available follow-up

<table>
<thead>
<tr>
<th></th>
<th>Laser</th>
<th>Aflibercept</th>
<th>Bevacizumab</th>
<th>Pegaptanib</th>
<th>Ranibizumab</th>
<th>Sham</th>
<th>Overall</th>
</tr>
</thead>
<tbody>
<tr>
<td>SSAE</td>
<td>9</td>
<td>3</td>
<td>6</td>
<td>2</td>
<td>10</td>
<td>5</td>
<td>18</td>
</tr>
<tr>
<td></td>
<td>1013</td>
<td>556</td>
<td>410</td>
<td>186</td>
<td>1303</td>
<td>528</td>
<td>4229</td>
</tr>
<tr>
<td>ATC*</td>
<td>10</td>
<td>3</td>
<td>4</td>
<td>2</td>
<td>8</td>
<td>2</td>
<td>15</td>
</tr>
<tr>
<td></td>
<td>824</td>
<td>846</td>
<td>330</td>
<td>188</td>
<td>1113</td>
<td>184</td>
<td>3718</td>
</tr>
<tr>
<td>Death</td>
<td>11</td>
<td>3</td>
<td>4</td>
<td>2</td>
<td>10</td>
<td>3</td>
<td>17</td>
</tr>
<tr>
<td></td>
<td>903</td>
<td>846</td>
<td>333</td>
<td>188</td>
<td>1521</td>
<td>434</td>
<td>4455</td>
</tr>
</tbody>
</table>

For each safety outcome, numbers in the table are the total number of studies (upper line for each outcome) and the total number of eyes (lower line for each outcome), as available by treatment and measured at the longest available follow-up.

(*) combined incidence of (1) cardiovascular, hemorrhagic, and unknown death; (2) nonfatal MI; and (3) nonfatal stroke.

Table 7. All serious systemic adverse events (longest available follow-up)

<table>
<thead>
<tr>
<th></th>
<th>CONTROL</th>
<th>AFLI</th>
<th>BEVA</th>
<th>PEGA</th>
<th>RANI</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>0.95 (0.75 to 1.20)</td>
<td>1.29 (0.43 to 3.84)</td>
<td>1.02 (0.67 to 1.53)</td>
<td>0.98 (0.76 to 1.25)</td>
<td></td>
</tr>
<tr>
<td>0.98 (0.83 to 1.16)</td>
<td>0.95 (0.75 to 1.20)</td>
<td>0.95 (0.75 to 1.20)</td>
<td>0.95 (0.75 to 1.20)</td>
<td>1.04 (0.83 to 1.32)</td>
<td></td>
</tr>
<tr>
<td>0.93 (0.73 to 1.19)</td>
<td>0.95 (0.76 to 1.18)</td>
<td>0.95 (0.75 to 1.20)</td>
<td>0.96 (0.77 to 1.20)</td>
<td>1.04 (0.64 to 1.64)</td>
<td></td>
</tr>
<tr>
<td>1.02 (0.64 to 1.64)</td>
<td>1.04 (0.63 to 1.72)</td>
<td>1.09 (0.64 to 1.86)</td>
<td>0.95 (0.57 to 1.58)</td>
<td>0.97 (0.80 to 1.17)</td>
<td></td>
</tr>
</tbody>
</table>

P value for overall inconsistency = 0.859.
Table 8. Antiplatelet Trialists Collaboration arterial thromboembolic events at the longest available follow-up

<table>
<thead>
<tr>
<th></th>
<th>CONTROL</th>
<th>AFLI</th>
<th>BEVA</th>
<th>PEGA</th>
<th>RANI</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>1.50 (0.81 to 2.79)</td>
<td>0.92 (0.17 to 5.12)</td>
<td>0.78 (0.31 to 1.97)</td>
<td>0.64 (0.38 to 1.07)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>0.88 (0.37 to 2.13)</td>
<td>1.46 (0.71 to 2.98)</td>
<td></td>
<td>2.26 (1.15 to 4.23)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>0.94 (0.33 to 2.66)</td>
<td>1.06 (0.36 to 3.11)</td>
<td>BEVA</td>
<td>1.51 (0.85 to 2.69)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>0.79 (0.20 to 3.02)</td>
<td>0.89 (0.18 to 4.43)</td>
<td>PEGA</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>1.09 (0.52 to 2.29)</td>
<td>1.24 (0.48 to 3.19)</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

- P value for differences between direct and indirect estimates = 0.002.
- P value for overall inconsistency = 0.274 in the network meta-analysis.

Table 9. All-cause mortality at the longest available follow-up

<table>
<thead>
<tr>
<th></th>
<th>CONTROL</th>
<th>AFLI</th>
<th>BEVA</th>
<th>PEGA</th>
<th>RANI</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>1.69 (0.30 to 9.42)</td>
<td>0.95 (0.06 to 14.85)</td>
<td>0.82 (0.25 to 2.65)</td>
<td>0.64 (0.32 to 1.25)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>1.01 (0.34 to 3.03)</td>
<td>AFLI</td>
<td></td>
<td>2.26 (0.80 to 6.40)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>1.61 (0.45 to 5.69)</td>
<td>1.59 (0.43 to 5.94)</td>
<td>BEVA</td>
<td>0.85 (0.40 to 1.83)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>0.81 (0.16 to 4.03)</td>
<td>0.81 (0.12 to 5.62)</td>
<td>PEGA</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>0.90 (0.40 to 2.01)</td>
<td>1.16 (0.38 to 3.58)</td>
<td></td>
<td>RANI</td>
<td></td>
</tr>
</tbody>
</table>

- P value for differences between direct and indirect estimates = 0.011.
- P value for differences between direct and indirect estimates = 0.030.
- P value for differences between direct and indirect estimates = 0.015.
- P value for differences between direct and indirect estimates = 0.022.
- P value for overall inconsistency = 0.087 in the network meta-analysis.

Table 10. Similarity among studies. baseline values and number of injections

<table>
<thead>
<tr>
<th>Study</th>
<th>Participants</th>
<th>Interventions</th>
<th>Mean n. injections</th>
<th>Visual acuity (logMAR)</th>
<th>Retinal thickness (µm)</th>
<th>Study sponsor</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ahmadieh 2008</td>
<td>78</td>
<td>Bevacizumab Sham</td>
<td></td>
<td></td>
<td></td>
<td>None reported</td>
</tr>
<tr>
<td>BOLT 2010</td>
<td>80</td>
<td>Bevacizumab Laser</td>
<td>9* 3*</td>
<td>0.59 0.61</td>
<td>507 482</td>
<td>Public</td>
</tr>
<tr>
<td>DA VINCI 2011</td>
<td>89</td>
<td>Aflibercept Laser</td>
<td>3.6 to 5.5 1.7</td>
<td>0.55</td>
<td>441</td>
<td>Industry</td>
</tr>
<tr>
<td>Study</td>
<td>N</td>
<td>Treatment 1</td>
<td>Treatment 2</td>
<td>Treatment 3</td>
<td>Baseline 1</td>
<td>Baseline 2</td>
</tr>
<tr>
<td>------------------</td>
<td>-----</td>
<td>-------------</td>
<td>-------------</td>
<td>-------------</td>
<td>------------</td>
<td>------------</td>
</tr>
<tr>
<td>DRCRnet 2010</td>
<td>668</td>
<td>Laser</td>
<td>Ranibizumab-PL</td>
<td>Ranibizumab-DL</td>
<td>3*</td>
<td>9*</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>0.38</td>
<td>0.39</td>
</tr>
<tr>
<td>DRCRnet 2015</td>
<td>620</td>
<td>Aflibercept</td>
<td>Bevacizumab</td>
<td>Ranibizumab</td>
<td>9.2</td>
<td>9.4</td>
</tr>
<tr>
<td></td>
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<td></td>
<td></td>
<td></td>
<td>0.40</td>
<td>0.40</td>
</tr>
<tr>
<td>Ekinci 2014</td>
<td>100</td>
<td>Bevacizumab</td>
<td>Ranibizumab</td>
<td></td>
<td>5.1</td>
<td>6.5</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>0.24</td>
<td>0.24</td>
</tr>
<tr>
<td>Ishibashi 2014</td>
<td>233</td>
<td>Pegaptanib</td>
<td>Sham</td>
<td></td>
<td>4</td>
<td>0.56</td>
</tr>
<tr>
<td>Korobelnik 2014</td>
<td>268</td>
<td>Aflibercept</td>
<td>Laser</td>
<td></td>
<td>8.5</td>
<td>2.4</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>0.50</td>
<td>0.50</td>
</tr>
<tr>
<td>Lopez-Galvez 2014</td>
<td>83</td>
<td>Ranibizumab</td>
<td>Laser</td>
<td></td>
<td>5.3</td>
<td>2.1</td>
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<td>Pegaptanib</td>
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Table 10. Similarity among studies, baseline values and number of injections  (Continued)

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<tr>
<th>Study</th>
<th>N</th>
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<th>N</th>
<th>Baseline</th>
<th>Number of injections</th>
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<td>Ranibizumab-PL</td>
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<td>0.52</td>
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<td>Soheilian 2007</td>
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<td>Bevacizumab Laser</td>
<td>352</td>
<td>0.71</td>
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<tr>
<td>Turkoglu 2015</td>
<td>70</td>
<td>Laser Ranibizumab</td>
<td>460</td>
<td>0.84</td>
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<tr>
<td>Wiley 2016</td>
<td>124</td>
<td>Bevacizumab Ranibizum</td>
<td>477</td>
<td>3</td>
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</tbody>
</table>

DL: plus deferred laser  
PL: plus prompt laser  
(*): median, not mean, available and reported

FEEDBACK
Feedback, 25 June 2013

Summary
Comments: 1. In the electronic searches, did you not find the article: Lim JW, Lee HK, Shin MC. Comparison of intravitreal bevacizumab alone or combined with triamcinolone versus triamcinolone in diabetic macular edema: A randomized clinical trial. Ophthalmologica. 2012;227(2):100-6. The article was published online: October 12, 2011, so it should have been found in the last electronic search, June 2012. I understand this article would have been excluded because of the triamcinolone comparison (it compares bevacizumab 1.25 mg versus bevacizumab 1.25 mg plus triamcinolone 2 mg versus triamcinolone 2 mg) but maybe it should appear in the ‘Characteristics of excluded studies’ section?

2. About the outcome results for ‘Quality of life’: Quality of life results should be included from the RESTORE 2011 trial. In the RESTORE 2011 trial (RESTORE 2011) data on quality of life have been reported using EQ-5D and NEI VFQ-25. It reported 12 months results, so it could also have been included. Mitchell P, Bandello F, Schmidt-Erfurth U, Lang G, Massin P, Schlingemann R, et al. The RESTORE 2011 Study ranibizumab monotherapy or combined with laser versus laser monotherapy for diabetic macular edema. Ophthalmology. 2011;118(4):615-25.

3. In the section Effects of interventions/Anti-VEGF versus sham treatment/ Quality of the evidence: “READ2 2009 provided visual gain, but not visual loss data”. This section evaluates anti-VEGF versus sham treatment and the READ trial is about ranibizumab versus laser.

4. For the included study: DCRRNet 2010 [published data only] Diabetic Retinopathy Clinical Research Network, Elman MJ, Aiello LP, Beck RW, Bressler NM, Bressler SB, et al. Randomized trial evaluating ranibizumab plus prompt or deferred laser or triamcinolone plus prompt laser for diabetic macular edema. Ophthalmology 2010;117(6):1064-77. It seems that you have also considered results from this trial, from the 2011 publication for 2 years results (Analysis 3.7-3.11): Elman MJ, Bressler NM, Qin H, Beck RW, Ferris FL 3rd, Friedman SM, et al. Expanded 2-year follow-up of ranibizumab plus prompt laser or deferred laser or triamcinolone plus prompt laser for diabetic macular edema. Ophthalmology. 2011;118(4):609-614. The values of “N”, total population evaluated belong to 2011 publication; the numbers are higher than those belonging to the 2010 publication. So this reference should also be cited.
5. For the included study: READ2 2009 [published data only] Nguyen QD, Shah SM, Khwaja AA, Channa R, Hatef E, Do DV, et al. Two-year outcomes of the ranibizumab for edema of the mAcula in diabetes (READ-2) study. Ophthalmology 2010;117(11):2146-51. The results that are considered in the review belong to the article by Nguyen 2009 (results and follow up at 6 months). Nguyen QD, Shah SM, Heier JS, Do DV, Lim J, Boyer D, et al. Primary end point (six months) results of the Ranibizumab for Edema of the mAcula in diabetes. Ophthalmology. 2009;116 (11):2175-81. All the analyses have been done with the 6 months follow up. Because after six months all patients could be treated with ranibizumab, data were not collected beyond six months. So this reference should also be cited.

6. In the 'Characteristics of included studies' table for RISE-RIDE, the 'outcomes' section should be completed.

7. In Tables 2, 5, 7, 8 and 9 'bevacacizumab' should be corrected to 'bevacizumab'.

Reply

We thank Ruth Ubago Pérez for her comments submitted through the Feedback system in The Cochrane Library.

1. In the 'Characteristics of excluded studies' table, we have added that not only Paccola 2008, but also Lim 2012 were excluded because another Cochrane review focuses on the use of intravitreal steroids in people with diabetic macular oedema.

2. We will include quality of life data in the next review update.

3. We have removed this sentence.

4 and 5. We have added these references.

6. We have completed the 'Outcomes' section.

7. We have corrected these typos.

Contributors

Comment from Ruth Ubago Pérez, Pharmacist Technician, Andalusian Agency for Health Technology Assessment, Spain

Reply from Gianni Virgili (lead author of review)

WHAT'S NEW

Last assessed as up-to-date: 26 April 2017.

<table>
<thead>
<tr>
<th>Date</th>
<th>Event</th>
<th>Description</th>
</tr>
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<tr>
<td>2 May 2017</td>
<td>New search has been performed</td>
<td>Issue 6, 2017: Updated protocol: objectives revised as comparing different antiangiogenic drugs using network meta-analysis technique</td>
</tr>
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</table>
HISTORY
Review first published: Issue 4, 2009

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<th>Date</th>
<th>Event</th>
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<tr>
<td>4 November 2014</td>
<td>Amended</td>
<td>Plain language summary title has been amended</td>
</tr>
<tr>
<td>17 October 2014</td>
<td>New citation required but conclusions have not changed</td>
<td>Issue 10, 2014: Five new studies (Azad 2012; Ekinci 2014; Nepomuceno 2013; RELATION 2012; RESPOND 2013) have been included in the update.</td>
</tr>
<tr>
<td>17 October 2014</td>
<td>New search has been performed</td>
<td>Issue 10, 2014: Electronic searches updated.</td>
</tr>
<tr>
<td>4 November 2013</td>
<td>Feedback has been incorporated</td>
<td>The authors have made some edits to the review in response to feedback received. See ‘Feedback 1’ for further details.</td>
</tr>
<tr>
<td>14 March 2013</td>
<td>Amended</td>
<td>The abstract has been amended to focus on the comparison with laser and presenting absolute effects</td>
</tr>
<tr>
<td>11 November 2012</td>
<td>New search has been performed</td>
<td>Updated searches yielded seven new trials for inclusion. One trial that had previously been included was excluded. An economic section has been added. One new author Massimo Brunetti has been added to the review team</td>
</tr>
<tr>
<td>11 November 2012</td>
<td>New citation required and conclusions have changed</td>
<td>Inclusion of seven new studies has changed the conclusions to this review from the previous version</td>
</tr>
</tbody>
</table>

CONTRIBUTIONS OF AUTHORS
Conceiving the review: GV, MP
Designing the review: GV, MP, EL
Co-ordinating the review: GV
Data collection for the review.

- Designing search strategies: IG
- Undertaking searches: IG
- Screening search results: GV, MP
- Organising retrieval of papers: IG
- Screening retrieved papers against inclusion criteria: GV, Cochrane Eyes and Vision
- Appraising quality of papers: GV, MP
- Extracting data from papers: GV, MP, EL
• Writing to authors of papers for additional information: GV, MP
• Obtaining and screening data on unpublished studies: GV, Cochrane Eyes and Vision

Data management for the review.
• Entering data into Review Manager 5: GV, MP, EL

Analysis of data: GV, EL

Interpretation of data.
• Providing a methodological perspective: GV, EL
• Providing a clinical perspective: GV, MP
• Providing a policy perspective: GV, MP, EL
• Providing a consumer perspective: none

Writing the review: GV, MP, EL

DECLARATIONS OF INTEREST

Gianni Virgili: none known
Mariacristina Parravano received payment for participating on the Advisory Board for Allergan, Bayer and Novartis.
Jennifer Evans: none known
Iris Gordon: none known
Ersilia Lucenteforte: none known

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• The Cochrane Review Incentive Scheme provided funding for Jennifer Evans to assist with the 2014 update of this review.

The views and opinions expressed therein are those of the authors and do not necessarily reflect those of the Systematic Reviews Programme, NIHR, NHS or the Department of Health.
D I F F E R E N C E S  B E T W E E N  P R O T O C O L  A N D  R E V I E W

Differences between protocol and review in the first published version of this review

We have added LILACS to the list of databases which have been searched for this review. We have used a sensitivity analysis for the robustness of results in comparisons including only one trial according to a statistical technique derived from a recent publication (Borm 2009).

Changes in update, 2012 compared to the protocol of the previous version

1. We have specified that studies comparing different anti-VEGF drugs will also be included in this review, but intravitreal steroids will be excluded as they are the subject of another Cochrane Review. Moreover, we decided not to consider the comparison of bevacizumab with bevacizumab plus triamcinolone, which included two studies; in fact this comparison investigates the additional effect of triamcinolone rather than the benefit of anti-VEGF drugs.
2. We have computed indirect comparison odds ratios (OR) of a gain of 3+ and 2+ lines for bevacizumab and pegaptanib versus ranibizumab as the reference drug using random-effects model logistic regression.

Changes in update, 2014 compared to the protocol of the previous version

1. We have included five more studies but the conclusions did not change.
2. We no longer consider economic evidence since antiangiogenic therapy is widely approved and reimbursed.
3. We eliminated the table on retinal detachment as an ocular adverse event since it proved to be extremely rare in all studies.
4. Units of analysis issue: in the update of this review we no longer performed a sensitivity analysis regarding the primary outcome to determine the impact of excluding studies with eyes, rather than participants, as the unit of analysis. In fact, a significant amount of evidence from studies with individuals as unit of analysis was achieved for the main comparisons.
5. Single trial issue: in the 2012 and 2014 updates of the review we did not use the sensitivity analysis on the robustness of single trial results recommended by Borm 2009, as was originally planned. Instead, we calculated the ‘Optimal Information Size’ to rate the quality of evidence regarding imprecision as recommended by the GRADE study group in Guyatt 2011.

Changes in update, 2016 compared to the protocol of the previous version

1. The objective was now to compare different anti-VEGF drugs and a new protocol was developed.
2. We used network meta-analysis technique to augment direct evidence with indirect evidence.
3. We restricted the number of outcomes to three efficacy outcomes, three safety outcomes and quality of life.
4. We have included six more studies and conclusions are changed.
5. The sensitivity analysis restricted to low risk of bias studies was added to the protocol.
6. We included a cross-over study and treated it as a parallel arm study in efficacy analyses.

I N D E X  T E R M S
Medical Subject Headings (MeSH)
Angiogenesis Inhibitors [*therapeutic use]; Antibodies, Monoclonal [therapeutic use]; Antibodies, Monoclonal, Humanized [therapeutic use]; Aptamers, Nucleotide [therapeutic use]; Bevacizumab; Diabetic Retinopathy [*complications]; Laser Coagulation [methods]; Macular Edema [*drug therapy; surgery]; Randomized Controlled Trials as Topic; Ranibizumab; Receptors, Vascular Endothelial Growth Factor [therapeutic use]; Recombinant Fusion Proteins [therapeutic use]; Triamcinolone [therapeutic use]; Vascular Endothelial Growth Factor A [*antagonists & inhibitors]

MeSH check words
Humans