

# Current treatments in diabetic macular oedema: systematic review and meta-analysis

Ford, J. A., Lois, N., Royle, P., Clar, C., Shyangdan, D., & Waugh, N. (2013). Current treatments in diabetic macular oedema: systematic review and meta-analysis. BMJ Open, 3(3), [e002269]. DOI: 10.1136/bmjopen-2012-002269

Published in: **BMJ Open** 

#### **Document Version:**

Publisher's PDF, also known as Version of record

# Queen's University Belfast - Research Portal:

Link to publication record in Queen's University Belfast Research Portal

**Publisher rights**Copyright 2013 the authors.

This is an open access Creative Commons Attribution-NonCommercial License (https://creativecommons.org/licenses/by-nc/2.0/), which permits use, distribution and reproduction for non-commercial purposes, provided the author and source are cited.

# General rights

Copyright for the publications made accessible via the Queen's University Belfast Research Portal is retained by the author(s) and / or other copyright owners and it is a condition of accessing these publications that users recognise and abide by the legal requirements associated with these rights.

# Take down policy

The Research Portal is Queen's institutional repository that provides access to Queen's research output. Every effort has been made to ensure that content in the Research Portal does not infringe any person's rights, or applicable UK laws. If you discover content in the Research Portal that you believe breaches copyright or violates any law, please contact openaccess@qub.ac.uk.

**Open Access** Research



# Current treatments in diabetic **DEN** macular oedema: systematic review and meta-analysis

John Alexander Ford, <sup>1</sup> Noemi Lois, <sup>2</sup> Pamela Royle, <sup>3</sup> Christine Clar, <sup>4</sup> Deepson Shyangdan, <sup>3</sup> Norman Waugh <sup>3</sup>

To cite: Ford JA, Lois N, Royle P. et al. Current treatments in diabetic macular oedema: systematic review and meta-analysis. BMJ Open 2013;3:e002269. doi:10.1136/bmjopen-2012-002269

Prepublication history for this paper are available online. To view these files please visit the journal online (http://dx.doi.org/10.1136/ bmjopen-2012-002269).

Received 26 October 2012 Accepted 1 February 2013

This final article is available for use under the terms of the Creative Commons Attribution Non-Commercial 2.0 Licence; see http://bmjopen.bmj.com

<sup>1</sup>Department of Population Health and Primary Care, Faculty of Medicine and Health Sciences, Norwich Medical School, University of East Anglia, Norwich, UK <sup>2</sup>Centre for Vascular and Visual Sciences, Queens University, Belfast, UK <sup>3</sup>Warwick Evidence, Division of Health Sciences, Warwick Medical School, Coventry, UK <sup>4</sup>Researcher in Systematic Reviews, Berlin, Germany

# Correspondence to

Dr John Alexander Ford; john.ford@uea.ac.uk

#### **ABSTRACT**

**Objectives:** The aim of this systematic review is to appraise the evidence for the use of anti-VEGF drugs and steroids in diabetic macular oedema (DMO) as assessed by change in best corrected visual acuity (BCVA), central macular thickness and adverse

Data source: MEDLINE, EMBASE, Web of Science with Conference Proceedings and the Cochrane Library (inception to July 2012). Certain conference abstracts and drug regulatory web sites were also searched.

Study eligibility criteria, participants and interventions: Randomised controlled trials were used to assess clinical effectiveness and observational trials were used for safety. Trials which assessed triamcinolone, dexamethasone, fluocinolone, bevacizumab, ranibizumab, pegaptanib or aflibercept in patients with DMO were included.

Study appraisal and synthesis methods: Risk of bias was assessed using the Cochrane risk of bias tool. Study results are narratively described and, where appropriate, data were pooled using random effects meta-analysis.

Results: Anti-VEGF drugs are effective compared to both laser and placebo and seem to be more effective than steroids in improving BCVA. They have been shown to be safe in the short term but require frequent injections. Studies assessing steroids (triamcinolone, dexamethasone and fluocinolone) have reported mixed results when compared with laser or placebo. Steroids have been associated with increased incidence of cataracts and intraocular pressure rise but require fewer injections, especially when steroid implants are

Limitations: The quality of included studies varied considerably. Five of 14 meta-analyses had moderate or high statistical heterogeneity.

# Conclusions and implications of key findings:

The anti-VEGFs ranibizumab and bevacizumab have consistently shown good clinical effectiveness without major unwanted side effects. Steroid results have been mixed and are usually associated with cataract formation and intraocular pressure increase. Despite the current wider spectrum of treatments for DMO, only a small proportion of patients recover good vision (≥20/40), and thus the search for new therapies needs to continue.

# ARTICLE SUMMARY

# **Article focus**

■ To review the evidence for triamcinolone, dexamethasone, fluocinolone, bevacizumab, ranibizumab, pegaptanib and aflibercept in the treatment of diabetic macular oedema.

#### **Kev messages**

- The anti-VEGFs ranibizumab and bevacizumab have consistently shown good clinical effectiveness in the short term without major unwanted side effects.
- Steroid results have been mixed and are usually associated with cataract formation and IOP increase.

# Strengths and limitations of this study

- A robust, detailed review of the literature has been undertaken and, when appropriate, data have been combined in meta-analysis.
- The quality of studies included varied considerably.

# INTRODUCTION

Diabetic macular oedema (DMO) is a complication of diabetic retinopathy and a leading cause of blindness. The prevalence of DMO is likely to increase with more people suffering from diabetes.<sup>1</sup> Increasing DMO has significant implications for patients, healthcare providers and wider society. Laser has been the mainstay of treatment, but recently antivascular endothelial growth factor (anti-VEGF) drugs and steroids have been introduced as potential alternatives to laser photocoagulation.

# **Burden of disease**

Diabetic retinopathy is present at the time of diagnosis of diabetes mellitus in 0-30% of individuals.2 The incidence is estimated to be 2.3/100 person-years for the overall diabetic population and 4.5 for patients on insulin therapy.<sup>3</sup> There is good evidence that progression to DMO is associated with

duration of disease,<sup>4–7</sup> poor glycaemic control<sup>8</sup> and, in type 2 diabetes, the need for insulin,<sup>9</sup> though the need for insulin therapy is more a marker for duration and poor control.

The number of people with DMO is likely to increase as diabetes becomes more common. Some reports have suggested a decrease in progression to severe visual loss between 1975–1985 and 1986–2008 in a combined population of types 1 and 2.<sup>10</sup> Regular screening for retinopathy and better glycaemic control are thought to have reduced the progression to severe visual loss. Diabetic retinopathy is associated with a reduced quality of life. Compared with all diabetic complications, blindness was perceived to be the third worst health state after a major stroke and amputation.<sup>11</sup>

In the USA, the presence of DMO at diagnosis is associated with 29% additional costs within the first 3 years compared with individuals without retinopathy at diagnosis. <sup>12</sup> In 2010, the estimated healthcare costs for DMO in England were £92 million, with £65.6 million being spent on hospital treatment and related costs. <sup>13</sup>

Visual impairment results in increased welfare costs, early retirement and costs of home help and carers. <sup>14</sup> In England in 2010 (total population 52.23 million), the estimated population with diabetes was 2.34 million; the above social costs were estimated to be £11.6 million for DMO. <sup>13</sup>

#### Overview of pathophysiology

DMO is caused mainly by disruption of the blood-retinal barrier. The complex pathway that leads to this disruption has been previously described in this journal. Sustained hyperglycaemia causes a multifactorial cascade of physiological processes, involving increased permeability, cytokine activation, altered blood flow, hypoxia and inflammation. Vascular endothelial growth factor-A (VEGF-A) is a major contributor to the inflammatory process and, in particular, to angiogenesis and permeability. Hypoxia caused by microvascular disease stimulates the release of VEGF-A to aid perfusion. There are six major isoforms of VEGF-A: 121, 145, 165, 183, 189 and 206. In addition to causing widespread microvascular injury, there is now evidence that hyperglycaemia results in preceding neuronal dysfunction, which may contribute to visual loss.

# **Overview of current treatments**

Laser photocoagulation has been the mainstay of treatment for DMO. The landmark Diabetic Retinopathy Study<sup>18</sup> and the Early Treatment Diabetic Retinopathy Study (ETDRS)<sup>19</sup> <sup>20</sup> demonstrated its clinical effectiveness. However, although laser photocoagulation was clearly effective in preserving vision, it was less successful in restoring it, once lost. Furthermore, patients with perifoveal ischaemia are not amenable to this form of therapy. In EDTRS, although laser was shown to reduce the risk of moderate visual loss (a loss of three ETDRS lines) by 50%, visual acuity improved in only 3% of patients.<sup>20</sup> However, in some recent trials, laser has

improved the proportion of patients with more than or equal to 10 letters by 7–31%. <sup>21–24</sup> In addition, laser is not without side effects. Foveal burns, visual field defects, retinal fibrosis and laser scars have been reported. <sup>25</sup> Over the following decade it became apparent that certain patients suffered severe visual loss despite aggressive treatment. <sup>26</sup>

Steroids and anti-VEGF drugs are newer treatments in DMO. Intravitreal corticosteroids have potent anti-inflammatory effects. Triamcinolone (Kenalog) is not licensed for eye use but has been used to treat DMO for over 10 years. Triamcinolone (Trivaris), recently, was licensed for eye use. The development of intravitreal implants has allowed sustained release formulations. Fluocinolone acetonide (Iluvien, Alimera Sciences) and dexamethasone (Ozudex, Allergan) are implants that have been introduced recently.

Anti-VEGF agents have shown efficacy compared with laser. Bevacizumab (Avastin, Genenetch/Roche) is a monoclonal antibody that targets all VEGF isoforms. Although being developed for colorectal cancer, it is widely used off-label, as an intravitreal treatment for macular oedema of different aetiologies. Ranibizumab (Lucentis, Genentech/Roche) is a fragment of the bevacizumab antibody (molecular weight of ranibizumab 48.4 KDa compared with 149 KDa for bevacizumab). It was designed specifically for use in the eye. Ranibizumab is considerably more expensive than bevacizumab (the estimated cost of ranibizumab is \$2000/dose compared with \$50 for bevacizumab). 27 Pegaptanib (Macugen, Evetech Pharmaceuticals/Pfizer) is a PEGvlated aptamer, with a high affinity to the VEGF isoform 165, and was approved for the treatment of exudative AMD in 2004. Aflibercept (Regeneron/Bayer HealthCare) is a recent addition to the anti-VEGF class that targets all forms of VEGF-A and placental growth factor.

# Aim of the review

The aim of this review is to provide clinicians with an up-to-date overview of current intraocular drug treatments for DMO. It is hoped that the information contained herein will assist clinicians to present their patients with the best evidence supporting each treatment, including possible complications. In addition, this review may be helpful to policy makers. The review focuses on the current evidence for the use of anti-VEGF drugs and steroids to treat DMO, as assessed by change in best corrected visual acuity (BCVA) (mean and proportion with more than two lines improvement), central macular thickness (CMT), as determined by optical coherence tomography (OCT), and their adverse events.

# **EVIDENCE ACQUISITION**

A systematic literature search was performed. The databases searched included MEDLINE, EMBASE, Web of Science with Conference Proceedings and the Cochrane Library. The dates searched were from the inception of each database until July 2012.

The search terms combined the following key words: ranibizumab or lucentis or bevacizumab or avastin or pegaptanib or macugen or aflibercept or vegf trap-eye or steroid\* or corticosteroid\* or dexamethasone or fluocinolone or triamcinolone or anti-VEGF\* or anti-vascular endothelial growth factor\*

#### AND

DMO or diabetic macular edema or diabetic retinopathy or diabetic maculopathy

#### AND

(masked or sham or placebo OR control group or random\*) OR (systematic review or meta-analysis) OR (risk or safety or adverse or harm or pharmacovigilance or side-effect\* or precaution\* or warning\* or contraindication\* or contra-indication\* or tolerability or toxic)

The meeting abstracts of the Association for Research in Vision and Ophthalmology, the American Diabetes Association (2002–2012) and the European Association for the Study of Diabetes were searched from 2002 to 2012.

In addition, the web sites of the European Medicines Agency and the US Food and Drug Association were searched for data on registration status and safety. Clinicaltrials.gov and the EU Clinical Trials Register were searched in July 2012 for data on ongoing research.

Full details of the searches are shown in appendix 1.

Randomised controlled trials (RCT) were used to evaluate clinical effectiveness. Safety was assessed through both RCTs and observational studies.

RCTs were included provided that they (1) addressed the use of triamcinolone, dexamethasone, fluocinolone, bevacizumab, ranibizumab, pegaptanib or aflibercept in patients with DMO, (2) had a minimum follow-up of 6 months and (3) had a minimum of 25 eyes per study arm. Studies were excluded if they (1) evaluated laser only, (2) assessed the effect of the aforementioned treatments in macular oedema due to other retinal diseases (instead of DMO), (3) used only a single dose, (4) were combined with a surgical intervention or (5) published studies in languages other than English. There were no exclusions based on drug dose. Trials were excluded if they evaluated combined drug treatment with surgery or systemic treatment.

Search results were screened by two independent authors (JF and PR/DS). Data were extracted by one author (CC) and checked by a second (JF). Data extracted included inclusion/exclusion criteria, baseline demographics, BCVA expressed as a change in logMAR/ETDRS letters or proportion of participants with more than two or three lines BCVA improvement, CMT and adverse events. Risk of bias was assessed using the Cochrane risk of bias tool.

Studies were assessed for similarity in study population, interventions (dose and frequency), outcomes and

time to follow-up, with a view to including similar studies in a meta-analysis. Conference abstracts were excluded from the meta-analysis because their quality and detailed methodology were not clear. A difference of 6 months was allowed between study follow-ups because of the potential heterogeneity from disease progression and differences in the number of doses prescribed. If salient data were not reported, such as SDs, data were sought by personal communication with authors. Data were analysed using Review Manager software. If data from multiple time-points were available, the primary end-point data were used. Data were entered by one author (JF) and double-checked by a second (DS). Mean differences were calculated for change in BCVA and CMT and ORs were calculated for proportion of participants with more than two lines improvement. The 95% CIs were calculated for all outcomes. Statistical heterogeneity was measured through  $I^2$  scores. A score of less than 30% was considered as low heterogeneity, a score of more than 70% was considered as high heterogeneity and scores between 30% and 70% were considered as moderate. A random effects model was used throughout. The random effects model assumes variability between studies and therefore models uncertainty into the meta-analysis. Fixed assumes no variability. Generally speaking, the random effects model results in wider CIs.

#### **RESULTS**

The literature search identified 430 unique articles for possible inclusion, as shown in figure 1. In total, 328 articles were excluded on the basis of title and abstract, leaving 102 full papers to be read. Fifty-one of these articles were excluded; the reasons for their exclusion are summarised in table 1. Fifty-one articles from 29 studies met the inclusion criteria and were included in the review; these are described in tables 3–16. Seven studies were suitable for meta-analysis.

# Study quality

The quality of the included studies was, in general, good as is shown in table 2. (Note that the meeting abstracts were not quality assessed, owing to the lack of details reported on the methods.) Most studies adequately described sequence generation, except in three studies where it was unclear. 28-30 However, allocation concealment was poorly described throughout, with only eight appropriately.31-38 addressing this issue reports Reporting of masking also varied. A number of studies masked patients using sham injection or sham laser.  $^{21}$   $^{24}$   $^{29}$   $^{31}$   $^{33}$   $^{36}$   $^{38}$   $^{39}$   $^{40}$  Various studies reported that masking of patients was impossible. Assessors, where reported, were masked. In two studies, incomplete outcomes were not addressed. 31 41 Baseline characteristics consistent within study treatment Administration of laser followed the ETDRS protocol, or a modified version, in all studies that described laser administration. 21–24 28 30 33 34 42 43 Two studies, both

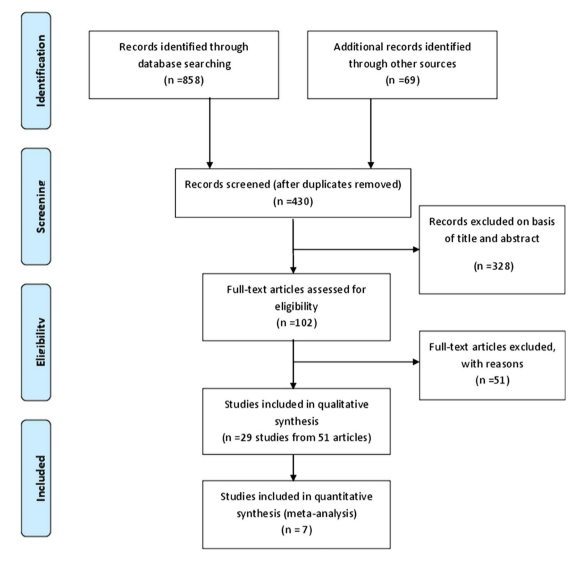


Figure 1 PRISMA flow diagram.

available only as meeting abstracts, did not report the laser administration details. 44 45

# **Intravitreal anti-VEGFs**

The characteristics of all published studies including design, inclusion/exclusion criteria, intervention, outcomes and their timing are shown in tables 3–8. Safety data for each drug are shown in tables 9–16.

# Ranibizumab

Nine RCTs have evaluated ranibizumab as a potential new treatment for patients with DMO (tables 3 and 8); seven were sponsored by industry, and two were led by independent investigators) (table 7). 21 46 READ-2 was the first large RCT (n=126). 28 47 It compared ranibizumab (0.5 mg) alone, and ranibizumab in combination with laser and laser alone. At 6 months, BCVA had improved significantly in the ranibizumab alone group compared with laser alone or ranibizumab plus laser. Addition of laser to ranibizumab did not provide additional BCVA

gain. REVEAL (n=396) compared ranibizumab (0.5 mg) with ranibizumab plus laser and laser alone. <sup>48</sup> At 12 months, both ranibizumab arms resulted in a statistically significantly better improvement in BCVA compared to laser alone. The addition of laser did not confer further benefit.

Within the past 2 years, the results of RESOLVE, <sup>36</sup> RESTORE <sup>24</sup> and RISE and RIDE <sup>38</sup> have been published in peer-reviewed journals. RESTORE (n=345) randomised similar groups as the READ-2 study (ranibizumab (0.5 mg) alone, laser alone and ranibizumab plus laser); outcomes were evaluated at 12 months. Ranibizumab improved mean BCVA, with laser providing no additional benefit. Two-year extended follow-up suggested that these results continued. <sup>49</sup> RESOLVE (n=151) compared two doses of ranibizumab (0.3 and 0.5 mg) with sham injection. The greatest improvement in BCVA at 12 months was in the 0.3 mg group (11.8 letter gain) compared to the 0.5 mg group (8.8 letter gain) or sham injection (1.4 letter loss). In this study, rescue laser was

Study	Reason
Active comparator trials	
Cho et al <sup>87</sup>	Single dose
DRCRN 2010	<6 months f/u
(Googe et al) <sup>88</sup>	Comonus I/a
Faghihi <i>et al</i> <sup>89</sup>	Single dose
Figueroa <i>et al</i> <sup>90</sup>	Single dose
Isaac <i>et al</i> <sup>91</sup>	Single dose
Paccola <i>et al</i> <sup>92</sup>	Single dose
Prager <i>et al</i> <sup>93</sup>	<25 pts per arm
Ozturk <i>et al</i> <sup>94</sup>	Non-RCT
Marey and Ellakwa <sup>95</sup>	<6 months
Shahin and El-Lakkany <sup>96</sup>	Single dose
Pegaptanib	Origic dosc
Loftus <i>et al</i> <sup>97</sup>	Quality of life data
Ranibizumab	Quality of life data
Ferrone and Jonisch <sup>98</sup>	<25 nte ner arm
Bevacizumab	<25 pts per arm
Solaiman <i>et al</i> <sup>99</sup>	Single door
DRCRN—Scott et al <sup>100</sup>	Single dose
Lee <sup>101</sup>	<25 pts per arm Non-RCT
Isaac <i>et al<sup>91</sup></i>	Single dose
Trimacinolone	Cinale deservation at the
Audren et al <sup>102</sup>	Single dose (dosing study)
Audren <i>et al</i> <sup>103</sup> Avitabile <sup>104</sup>	Single dose
	Mixed RVO and DMO
Bandello <i>et al</i> <sup>105</sup>	Case report+PDR
Bonini <i>et al</i> <sup>106</sup>	Single dose injection technique
Cellini <i>et al</i> <sup>107</sup>	Single injection PSTI
Cardillo <i>et al</i> <sup>108</sup>	Single injection PSTI
Chung et al <sup>109</sup>	Single injection PSTI
Dehghan et al <sup>110</sup>	Single dose
DRCRN—Chew et al 111	<25 pts per arm
Gil et al <sup>112</sup>	<25 pts per arm
Entezari <i>et al</i> <sup>113</sup>	<6 months
Hauser et al <sup>114</sup>	Single dose
Jonas et al <sup>115</sup>	Single dose
Joussen et al <sup>116</sup>	Study protocol
Avci and Kaderli <sup>117</sup>	Anaesthetic technique
Kang et al <sup>118</sup>	Single dose
Kim et al <sup>119</sup>	Single injection and CME
Lam et al <sup>120</sup>	Single injection
Lee <sup>121</sup>	Single injection
Maia <i>et al</i> <sup>122</sup>	Single dose
Massin et al <sup>123</sup>	Single dose
Mohamed et al <sup>124</sup>	Post hoc analysis
Nakamura et al <sup>125</sup>	Single dose
Spandau <i>et al</i> <sup>126</sup>	Single dose
Tunc <sup>127</sup>	<6 months
Verma <i>et al</i> <sup>128</sup>	Single dose
Wickremasinghe et al <sup>129</sup>	Single dose
Yalcinbayir et al <sup>130</sup>	Single dose
Dexamethasone	
Haller <i>et al</i> <sup>131</sup>	<6 months
Haller <i>et al</i> <sup>132</sup>	<25 pts per arm
Kuppermann <i>et al</i> 133	Mixture of macular oedema
	causes
Boyer et al <sup>134</sup>	Non-randomised
Fluocinolone	
Campochiaro et al <sup>135</sup>	<25 pts per arm
Diclofenac	
Elbendary <sup>71</sup>	<35 pts per arm
,	p p

allowed after 3 months of treatment, if BCVA had decreased by 10 letters or more, or if the investigator considered the macula not to be flat as assessed by OCT. Only 4.9% of the ranibizumab group required rescue laser, compared with 34.7% in the sham injection group.

READ-2 and RESTORE were suitable for pooling through meta-analysis and, when doing so, it was found that ranibizumab statistically significantly improved mean BCVA compared with laser (figure 2). In regard to the proportion of patients gaining more than or equal to 15 letters, individual trials showed a statistically significant difference between laser and ranibizumab but when these two trials were pooled using a random effects model, the result was no longer statistically significant. When a fixed effects model was used, the result was statistically significant (figure not shown). Adding laser to ranibizumab did not add any significant benefit (figure 3). In fact, the mean change in BCVA and the proportion of patients with more than 15 letter gain favoured, although not statistically significantly so, ranibizumab alone compared with ranibizumab plus laser. This was probably a chance effect.

RISE (n=377) and RIDE (n=382) were identical in design. The study arms are similar to those in the RESOLVE study, 0.3 or 0.5 mg ranibizumab compared with sham. In the RISE study, the proportion of patients with 15 or more letter gain was greatest in the 0.3 mg group at 24 months, whereas in the RIDE study this was greatest in the 0.5 mg group. In the DRCRN trial (n=854), Elman and colleagues compared ranibizumab (0.5 mg) plus prompt (within 3-10 days post ranibizumab) or deferred (≥24 weeks) laser with sham injection plus prompt laser, or triamcinolone (4 mg, Trivaris) plus prompt laser (table 8). At 1 year, both ranibizumab groups reported greater gains in mean BCVA change than triamcinolone or laser alone. Interestingly, at 2 years (n=628), the proportion of patients with 10 or more letter gain was not statistically significantly different between ranibizumab plus prompt laser and laser alone groups, but was statistically significant in the ranibizumab plus deferred laser compared with laser alone comparison. The reason for this is not clear.

READ-3 (n=152) has been published in abstract form and compared monthly injections of intravitreal ranibizumab high dose (2.0 mg) and low dose (0.5 mg).  $^{50}$  At 6 months, there was no statistically significant difference in BCVA between groups.

One study (n=63), published in abstract form, was identified which directly compared monthly injections of ranibizumab (0.5 mg) with bevacizumab (1.5 mg).<sup>51</sup> At 48 weeks, the authors found no statistically significant difference between bevacizumab and ranibizumab.

RESTORE, READ-2 and DRCRN (12 month data used) were suitable for pooling through meta-analysis to compare ranibizumab plus laser and laser alone (figure 4). Ranibizumab plus laser resulted in a statistically significantly greater change in mean BCVA, proportion of patients with more than 15 letter gain and CMT reduction versus laser alone.

injection; RVO, retinal vein occlusion.

Adequate sequence generation  Unclear  Yes	Allocation concealment  Unclear  Yes	Masking Unclear  Yes (patients and outcome assessors)	Incomplete outcome data addressed  Yes (91.3% completion)  Yes (82% completion in	Free of selective reporting  Yes	Free of other bias (eg, similarity at baseline, power assessment)  Comparison groups similar at baseline; power analysis not mentioned	Funder  Juvenile Diabetes Research Foundation, Genentech Inc
Yes		Yes (patients and	completion) Yes (82%		similar at baseline; power analysis not	Research Foundation,
Yes		Yes (patients and	completion) Yes (82%		similar at baseline; power analysis not	Research Foundation,
	Yes		•	Vaa		
Yes			sham arm, 90.2% with ranibizumab)	Yes	Comparison groups similar at baseline; power analysis unclear	Novartis Pharma, Switzerland
	Unclear	Yes (patients, outcome assessors)	Yes (87.3–88.3% completion)	Yes	Comparison groups similar at baseline; power analysis carried out (power adequate for VA changes)	Novartis Pharma, Switzerland
Yes	Yes	Yes (patients, treating physician masked to assigned dose of ranibizumab)	Yes (2 year study completed by 83.3% of patients in RISE and by 84.6% in RIDE)	Yes	Comparison groups similar at baseline; ITT analysis; power analysis carried out (power adequate for primary endpoint)	Genentech Inc
Yes	Unclear	Partial (outcome assessors, not patients)	Yes (97.5% completion)	Yes	Comparison groups similar at baseline (except laser group had longer duration of clinically significant DMO); power analysis carried out (power adequate for VA changes)	Moorfields Special Trustees, National Institute for Health Research
Yes	Unclear	Yes (patient	Yes (100% completion)	Yes	Comparable groups at baseline	Not specified
Yes	Yes	Yes (patients and technicians assessing BCVA, OCT and IOP)	Yes (92.3% follow-up at 6 months)	Yes	Comparison groups similar at baseline; power analysis carried out (power adequate for CMT changes)	Supported in part by the Action for Vision Eye Foundation Hong Kong (charity)
	Yes	Yes Unclear	Yes Yes (patients, treating physician masked to assigned dose of ranibizumab)  Yes Unclear Partial (outcome assessors, not patients)  Yes Unclear Yes (patient  Yes Yes Yes (patients and technicians assessing	Yes Yes (patients, treating physician masked to assigned dose of ranibizumab)  Yes (2 year study completed by 83.3% of patients in RISE and by 84.6% in RIDE)  Yes Unclear Partial (outcome assessors, not patients)  Yes (97.5% completion)  Yes Yes (patient Yes (100% completion)  Yes Yes Yes (patients and technicians assessing follow-up at	Yes Yes Yes (patients, treating physician masked to assigned dose of ranibizumab)  Yes (2 year yes study completed by 83.3% of patients in RISE and by 84.6% in RIDE)  Yes Unclear Partial (outcome assessors, not patients)  Yes (97.5% Yes completion)  Yes Yes (patient Yes (100% Yes completion)  Yes Yes Yes (patients and Yes (92.3% Yes follow-up at	Yes Yes (patients, treating physician masked to assigned dose of ranibizumab)  Yes Unclear Partial (outcome assessors, not patients)  Yes (patients)  Yes (97.5% Yes (27.5% Yes (27.5% Yes (27.5% Comparison groups similar at baseline) primary endpoint)  Yes Unclear Partial (outcome assessors, not patients)  Yes (97.5% Yes (27.5% Yes (27.5% Comparison groups similar at baseline (except laser group had longer duration of clinically significant DMO); power analysis carried out (power adequate for primary endpoint)  Yes Unclear Yes (patient Yes (100% Yes Comparable groups at baseline)  Yes Yes (patients and technicians assessing BCVA, OCT and IOP)  Bower analysis carried out (power analysis carried out (power analysis carried out (power adequate for VA comparable groups at baseline)  Yes (92.3% Yes Comparable groups at similar at baseline; ITT analysis carried out (power analysis carried out (power adequate for VA comparable groups at similar at baseline)  Yes (92.3% Yes Comparable groups at similar at baseline; ITT analysis carried out (power analysis car

Study (author and year)	Adequate sequence generation	Allocation concealment	Masking	Incomplete outcome data addressed	Free of selective reporting	Free of other bias (eg, similarity at baseline, power assessment)	Funder
Pegaptanib Cunningham <i>et all</i> Adamis <i>et al</i> <sup>99 57</sup>	Yes	Unclear	Yes (patients and outcome assessors)	Yes (95% completion)	Yes	Comparison groups similar at baseline; acknowledge lack of power to detect differences between doses of pegaptanib	Eyetech Pharmaceuticals Inc, New York, and Pfizer Inc, New York
Sultan <i>et al</i> <sup>40</sup>	Yes	Unclear	Yes (patients and outcome assessors)	Yes (69.9–73.8% completion)	Yes	Comparison groups similar at baseline; power analysis carried out (power adequate for VA changes)	Pfizer Inc, New York
Aflibercept Da Vinci et al <sup>80 58</sup>	Unclear (predetermined randomisation scheme)	Unclear	Yes (patients)	Yes (85% completion)	Yes	Comparison groups similar at baseline, power calculation completed	Regeneron Pharmaceuticals, Inc, New York
Steroids Dexamethasone Haller et al <sup>59</sup>	Yes	Unclear	Yes (patients to dexamethasone dose, outcome assessors)	Yes (92% completion)	Yes	Comparison groups similar at baseline; power analysis carried out, but study not powered to detect differences in subgroups	Oculex Pharmaceuticals Inc
FAME Study (Campochiaro <i>et al</i> ) <sup>29 60</sup>	Unclear	Unclear	Partial (patients, masking of outcome assessment not mentioned)	Yes (drop-out rate 19.0–22.7%)	Yes	Comparison groups similar at baseline; power analysis not mentioned	Alimera Sciences Inc, Atlanta, Georgia; Psivida Inc, Watertown, Massachusetts
Pearson <i>et al</i> <sup>43</sup>	Yes	Unclear	Third party masked design (patient and investigator not masked)	No losses to follow-up	Yes	Demographic characteristics were similar between implant and SOC groups; power calculation done, study adequately powered	Bausch & Lomb Inc, Rochester, New York

Ford JA, Lois N, Royle P, et al. BMJ Open 2013;3:e002269. doi:10.1136/bmjopen-2012-002269

Study (author and year)	Adequate sequence generation	Allocation concealment	Masking	Incomplete outcome data addressed	Free of selective reporting	Free of other bias (eg, similarity at baseline, power assessment)	Funder
Triamcinolone DRCR Network 2008 <sup>22 61 63 64</sup>	Yes	Unclear	Partial (patients to triamcinolone dose, outcome assessors not formally masked but generally not aware of participant's study group)	Yes (81–86% completion)	Yes	Comparison groups similar at baseline; power analysis carried out (power adequate for VA changes)	Cooperative agreement from the National Eye Institute, National Institute of Diabetes and Digestive and Kidney Diseases, National Institutes of Health, Department of Health and Human Services
Gillies <i>et al</i> Sutter <i>et al</i> <sup>32</sup> <sup>136–138</sup>	Yes	Yes	Yes (patients, outcome assessors)	Yes (91% completion intervention, 83% control)	Yes	Comparison groups similar at baseline (but limited demographic data); power analysis carried out (power adequate for VA changes)	Sydney Eye Hospital Foundation and Juvenile Diabetes Research Foundation, New York
Gillies <i>et al<sup>33</sup></i>	Yes	Yes	Yes (patients, outcome assessors)	Yes (84.5% completion)	Yes	Power analysis carried out (power adequate for VA changes)	National Health and Medical Research Council, Canberra, Australia, and the Sydney Eye Hospital Foundation Sydney, Australia
Lam <i>et al<sup>64</sup></i>	Yes	Yes	Partial (outcome assessors)	No losses to follow-up	Yes	Comparison groups similar at baseline; power analysis carried out (power adequate for CMT changes)	Action for Vision Foundation, Hong Kong
Ockrim <i>et all</i> Sivaprasad <i>et al</i> <sup>42 62</sup>	Yes	Unclear	Unclear	Yes (94% completion)	Yes	Comparison groups similar at baseline; power analysis carried out (power adequate for VA changes)	Special Trustees of Moorfields Eye Hospital

Study (author and year)	Adequate sequence generation	Allocation concealment	Masking	Incomplete outcome data addressed	Free of selective reporting	Free of other bias (eg, similarity at baseline, power assessment)	Funder
Active comparator trial Ahmadieh et al <sup>β1</sup>	rs Yes	Yes	Yes (patients and outcome assessors)	Unclear	Yes	CMT lower in control group at baseline (p<0.05), other baseline values similar; power analysis carried out (power adequate for CMT changes)	Not reported
DRCR Network <sup>21 46</sup>	Yes	Unclear	Yes (patients, except deferred laser group; outcome assessors); masking discontinued after the first year	Yes (1 year completion for 91–95% of eyes)	Yes	Comparison groups similar at baseline; power analysis carried out (power adequate for VA changes)	Cooperative agreement from the National Eye Institute, National Institute of Diabetes and Digestive and Kidney Diseases, National Institutes of Health and Human Services; Ranibizumab provided by Genentech, triamcinolone provided by Allergan Inc; companies also provided funds to defray the study's clinical site costs
Lim <i>et al<sup>55</sup></i>	Yes	Unclear	Yes (investigators only)	Yes (7.5% drop out after enrolment)	Yes	Groups similar at baseline. The bevacizumab group received more injections	Not reported
Soheilian <i>et al<sup>37 41</sup></i>	Yes	Yes	Yes (patients and outcome assessors)	Unclear (36 week completion for 76–88%)	Yes	CMT significantly lower and VA significantly better in MPC group at baseline, other baseline values similar; power analysis carried out (power adequate for VA changes)	Ophthalmic Research Centre, Labbafinejad Medical Center, Tehran

Study	Participants and baseline values	Intervention	Outcome (change from baseline at study end)		
READ-2 Study (Nguyen et al) <sup>28 47</sup> USA Multicenter Design: 3-arm RCT Follow-up: 6 months, 2-year extension (no relevant outcomes as IVR received by all groups by that time, no safety outcomes for 2-year data)	N: 126 eyes of 126 patients Inclusion criteria: ≥18 years, type 1 or 2 DM, DMO, BCVA 20/40-20/ 320, CMT ≥250 µm, HbA1c ≥6% within 12 months before randomisation; expectation that scatter laser photocoagulation not required for 6 months Exclusion criteria: contributing causes to reduced BCVA other than DMO, focal/grid laser within 3 months, intraocular steroid within	Group 1 (IVR, n=42 eyes): IV injections of 0.5 mg ranibizumab at baseline, 1, 3 and 5 months Group 2 (L, n=42 eyes): focal/grid laser at baseline and 3 months if CMT ≥250 µm Group 3 (IVRL, n=42 eyes): IV injections of 0.5 mg ranibizumab at baseline and 3 months, followed by focal/grid laser treatment 1 week later Regimen for all groups: after	At 6 months BCVA (ETDRS):  IVR L IVRL  IVR L IVR CMT (OCT):	BCVA (letters) +7.24 -0.43 +3.80 Plus ≥3 lines 22% 0 8%	<i>p Value</i> 0.0003 vs L NS vs IVR or L <0.05 vs L
	3 months, intraocular VEGF antagonist within 2 months <i>Age</i> : 62 years <i>Sex</i> : 52–69% female <i>Diabetes type</i> : not reported <i>HbA1c</i> : 7.39–7.77% <i>Baseline VA</i> : ETDRS letter score 24.85–28.35 <i>Baseline CMT</i> : excess foveal	6 months, patients could receive IV injections of ranibizumab no more than every 2 months or focal/grid laser no more than every 3 months if CMT ≥250 µm Laser Modified ETDRS protocol was used	IVR  L IVRL	CMT (µm) -106.3 -82.8 -117.2	p Value All <0.01 vs baseline, NS for elimination of ≥50% excess foveal thickness between groups
READ-3 Study (Do <i>et al</i> ) USA <sup>50</sup> <i>Design</i> : phase 2, 2-arm RCT	thickness 198.75–262.52 µm  Comorbidities: not reported  N: 152 eyes  Inclusion criteria: NR	Group 1 (IVR2.0, n=NR): monthly injections	At 6 months: BCVA		
Follow-up: 6 months	Exclusion criteria: NR Age: NR Sex: NR Diabetes type: NR HbA1c: NR Baseline VA: Mean BCVA Snellen equivalent 20/63 in the 2.0 mg group and 20/80 in the 0.5 mg group Baseline CST (central subfield thickness): 432 µm in the 2.0 mg group and 441 µm in the 0.5 mg group Comorbidities: NR	Group 2 (IVR0.5, n=NR): monthly injections After month 6, eyes evaluated and additional ranibizumab injections given on an as needed basis if DMO still present on OCT.	IVR2.0 IVR0.5 CST IVR2.0 IVR0.5	Mean BCVA letters gain +7.46 +8.69 CST reduction -163.86 µm -169.27 µm	p Value  NR NR NR NR

Study	Participants and baseline values	Intervention	Outcome (change from baseline at study end)		
RESOLVE Study (Massin	N: 151 eyes of 151 patients	Group 1 (IVR0.3, n=51 eyes):	At 12 months		
et al) <sup>36</sup>	Inclusion criteria: >18 years, type 1	0.3 mg (0.05 ml) IV ranibizumab,	BCVA (ETDRS):		
Multicenter international	or 2 DM, clinically significant DMO,	3 monthly injections (dose up to		BCVA (letters)	p Value
<i>Design</i> : 3-arm	BCVA 20/40–20/160, HbA1c <12%,	,	IVR0.3	+11.8 SD6.6	<0.0001 vs C
placebo-controlled RCT	decreased vision attributed to	Group 2 (IVR0.5, n=51 eyes):	IVR0.5	+8.8 SD11.0	<0.0001 vs C
Follow-up: 12 months	foveal thickening from DMO, laser	0.5 mg IV (0.05 ml) ranibizumab,	C	-1.4 SD14.2	
	photocoagulation could be safely	3 monthly injections (dose up to		Change ≥10 lette	
	withheld in the study eye for at	1.0 mg, see below)	IVR0.3	Gain 72.5%	<0.0001 vs C
	least 3 months after randomisation	Group 3 (C, n=49 eyes): sham		loss 0	
	Exclusion criteria: unstable medical	treatment, 3 monthly injections	IVR0.5	Gain 49%	0.001 vs C
	status, panretinal laser	Regimen for all groups: after month		loss 9.8%	
	photocoagulation performed within	1, the injection dose could be	C	Gain 18.4%	
	6 months before study entry,	doubled if CMT remained >300 μm		loss 24.5%	
	previous grid/laser	or was >225 µm and reduction in	CMT (OCT):		
	photocoagulation except patients	retinal oedema from previous		CMT (µm)	p Value
	with only mild laser burns at least	assessment was <50 µm; once	IVR0.3	-200.7 SD122.2	<0.0001 vs C
	1000 µm from the centre of the	injection volume was 0.1 ml it	IVR0.5	-187.6 SD147.8	<0.0001 vs C
	fovea performed >6 months	remained that for subsequent	C	-48.4 SD153.4	
	previously	injections; if treatment had been			
	Age: 63-65 (range 32-85) years	withheld for >45 days, subsequent			
	Sex: 43.1-49% female	injections restarted at 0.05 ml;			
	Diabetes type: 96.1–98% type 2	68.6% of dose doubling with			
	DM	ranibizumab, 91.8% with sham;			
	HbA1c: 7.3-7.6 (range 5.3-11.1) %				
	Baseline VA: ETDRS letter score	photocoagulation in sham group,			
	59.2–61.2 SD9.0–10.2	4.9% in ranibizumab group			
	Baseline CMT: 448.9–459.5				
	SD102.8–120.1 μm				
	Comorbidities: not reported				
RESTORE Study (Mitchell	N: 345 eyes of 345 patients	Group 1 (IVR, n=116 eyes): 0.5 mg	At 12 months		
et al) <sup>24 49</sup>	Inclusion criteria: ≥18 years, type 1	IV ranibizumab plus sham laser	BCVA (ETDRS):		

Ford JA, Lois N, Royle P, et al. BMJ Open 2013;3:e002269. doi:10.1136/bmjopen-2012-002269

Study	Participants and baseline values	Intervention	Outcome (change from baseline at study end)		
Multicenter international	or 2 DM, HbA1c ≤10%, visual	(median injections 7 (range 1–12),		BCVA (letters)	p Value
<i>Design</i> : 3-arm RCT	impairment due to DMO (eligible for	median sham laser treatments 2	IVR	+6.1 SD6.43	<0.0001 vs L
follow-up: 12 months	laser treatment), stable medication	(range 1-5))	IVRL	+5.9 SD7.92	<0.0001 vs L
	for management of diabetes, BCVA		L	+0.8 SD8.56	
	ETDRS letter score 39–78	0.5 mg IV ranibizumab plus active		BCVA change cat	•
	Exclusion criteria: concomitant eye conditions that could affect VA,	laser (median injections 7 (range 2–12), median laser treatments 1	IVR	Plus ≥10: 37.4% Loss ≥10: 3.5%	<0.0001 vs L
	active intraocular inflammation or	(range 1–5))	IVRL	Plus ≥10: 43.2%	<0.0001 vs L
	infection, uncontrolled glaucoma in	Group 3 (L, n=111 eyes): laser		Loss ≥10: 4.2%	
	either eye, panretinal laser	treatment plus sham injections	L	Plus ≥10: 15.5%	
	photocoagulation within 6 months	(median sham injections 7 (range		Loss ≥10: 12.7%	
	or focal/grid laser photocoagulation	1–12), median laser treatments 2	CMT (OCT):		
	within 3 months prior to study entry,	(range 1-4))		CMT (µm)	p Value
	history of stroke, hypertension	Regimen for all groups: 3 initial	IVR	-118.7	0.0002 vs L
	Age: 62.9-64.0 SD8.15-9.29 years	monthly injections, followed by		SD115.07	
	Sex: 37.1–47.7% female	retreatment schedule; 1 injection	IVRL	-128.3	<0.0001 vs L
	Diabetes type: 86.4–88.8% type 2	per month if stable VA not reached;		SD114.34	
	DM	Laser retreatments in accordance	L	-61.3 SD132.29	
	HbA1c: not reported	with ETDRS guidelines at intervals			
	Baseline VA: ETDRS letter score	no shorter than 3 months from			
	62.4–64.8 SD9.99–11.11	previous treatment			
	Baseline CMT: 412.4–426.6				
	SD118.01–123.95				
FVFAL Study (Obii and	Comorbidities: not reported	Crown 1 (IVID 0.5 ) show loos	At 10 months		
EVEAL Study (Ohji and hibashi ) <sup>48</sup>	N: 396 patients Inclusion criteria: NR	Group 1 (IVR 0.5 + sham laser, n=133): day 1, month 1, 2 and	At 12 months BCVA:		
apan Multicenter	Exclusion criteria: NR	pro-renata thereafter based on	DCVA.	Moon avorage	p Value
<i>lesign</i> : phase III	Age: 61.1 years	BCVA		Mean average change from	p value
ouble-masked RCT	Sex: NR	Group 2 (IVR 0.5+ active laser,		baseline to	
ollow-up: 12 months	Diabetes type: 98.7% with type 2	<i>n=132</i> ): day 1, month 1, 2 and		months 1–12	
onow up. 12 months	diabetes	pro-renata thereafter based on	IVR+sham laser	+5.9	vs laser <0.0001
	HbA1c: 7.5%	BCVA	IVR+laser	+5.7	vs laser <0.0001
	Baseline VA: 58.6 letters	Group 3 (sham injection + active	Laser+sham	+1.4	10 10001 10:000
	Baseline CMT: 421.9 µm	laser, n=131): day 1, month 1, 2		Mean change	
	Comorbidities: NR	and pro-renata thereafter based on		from baseline to	
		BCVA		month12 in	
		Active/sham laser photocoagulation		BCVA and CRT	
		performed according to ETDRS	IVR+sham laser	+6.6; -148.0 μm	vs C <0.0001
		guidelines at ≥3 month intervals	IVR+laser	+6.4; –163.8 µm	vs C <0.0001
			Laser+sham	+1.8; -57.1 µm	

Table 3 Continued					
Study	Participants and baseline values	Intervention	Outcome (change from baseline at study end)		
RISE Study (Brown et all	N: 377 eyes of 377 patients	Group 1 (IVR0.3, n=125 eyes):	At 24 months		
Nguyen et al) <sup>38</sup> 139	<i>Inclusion criteria</i> : ≥18 years, type 1	0.3 mg IV ranibizumab	BCVA:		
USA	or 2 diabetes, BCVA 20/40–20/320,	Group 2 (IVR0.5, n=125 eyes):		Plus ≥15 letters	p Value
Multicenter	DMO CMT ≥275 µm	0.5 mg IV ranibizumab	IVR0.3	44.8%	<0.0001 vs C
Design: 3-arm double-blind	Exclusion criteria: prior vitreoretinal	Group 3 (C, n=127 eyes): sham	IVR0.5	39.2%	=0.0002 vs C
sham-controlled RCT	surgery, recent history (within	injection	С	18.1%	
Follow-up: 24 months	3 months of screening) of	Regimen for all groups: monthly		Loss of <15	
	panretinal or macular laser in the	injections; need for macular rescue		letters	
	study eye, intraocular	laser assessed monthly starting at	IVR0.3	97.6%	=0.0086 vs C
	corticosteroids or antiangiogenic	month 3	IVR0.5	97.6%	=0.0126 vs C
	drugs, those with uncontrolled		С	89.8%	
	hypertension, uncontrolled diabetes			Snellen	
	(HbA1c >12%), recent (within			equivalent of	
	3 months) cerebrovascular accident			20/40 or better	
	or myocardial infarction		IVR0.3	60%	<0.0001 vs C
	Age: 61.7–62.8 SD8.9–10.0 (range		IVR0.5	63.2%	<0.0001 vs C
	21–87) years		C	37.8%	
	Sex: 41.6–48% female			Mean BCVA	
	Diabetes type: type 1 or 2		IVR0.3	gain (letters)	.0.0001 va C
	HbA1c: 7.7% SD 1.4–1.5; ≤8%		IVR0.5	+12.5 SD14.1 +11.9 SD12.1	<0.0001 vs C <0.0001 vs C
	(65–68.3%); >8% (31.7%–35%) Baseline VA: Mean ETDRS letter		C	+11.9 SD12.1 +2.6 SD13.9	<0.0001 VS C
	score 54.7–57.2; ≤20/200		CFT:	+2.0 3013.9	
	(7.9–13.6%); >20/200 but		OF I.	Mean change	p Value
	(7.9–13.0%), >20/200 but <20/40 (72.4–72.8%); ≥20/40			from baseline	p value
	(13.6–19.7%)		IVR0.3	-250.6 SD212.2	<0.0001 vs C
	Baseline CMT: 463.8–474.5 μm		IVR0.5	-253.1 SD183.7	<0.0001 vs C
	Comorbidities: History of smoking		C	-133.4 SD209.0	<b>10.0001 10 0</b>
	46.4–51.2%			100.1 02200.0	
RIDE study (Boyer et all	N: 382 eyes	Group 1 (IVR0.3, n=125 eyes):	At 24 months		
Nguyen <i>et al</i> ) <sup>38</sup> 140	Inclusion criteria: ≥18 years, type 1	0.3 mg IV ranibizumab	BCVA:		
USA	or 2 diabetes, BCVA 20/40–20/320	Group 2 (IVR0.5, n=127 eyes):		More than 15	p Value
Multicentre	and DMO CMT ≥275 µm	0.5 mg IV ranibizumab		letters	
Design: 3-arm double-blind	Exclusion criteria: prior vitreoretinal	Group 3 (C, n=130 eyes): sham	IVR0.3	33.6%	<0.0001 vs C
sham-controlled RCT	surgery, recent history (within	injection	IVR0.5	45.7%	<0.0001 vs C
Follow-up: 24 months	3 months of screening) of	Regimen for all groups: Patients	С	12.3%	
					Contin

Study	Participants and baseline values	Intervention	Outcome (change from baseline at study end)		
	panretinal or macular laser in the	were eligible for rescue macular		Less than 15	
	study eye, intraocular	laser starting at month 3		letters	
	corticosteroids or antiangiogenic		IVR0.3	1.6%	>0.05 vs C
	drugs, those with uncontrolled		IVR0.5	3.9%	<0.05 vs C
	hypertension, uncontrolled diabetes		С	8.5%	
	(HbA1c >12%), recent (within			Snellen	
	3 months) cerebrovascular accident			equivalent of	
	or myocardial infarction			20/40 or better	
	<i>Age</i> : 61.8–63.5 (range 22–91)		IVR0.3	54.4%	=0.0002 vs C
	years		IVR0.5	62.2%	<0.0001 vs C
	Sex: 37-49.1% female		С	34.6%	
	Diabetes type: type 1 or 2			Mean BCVA gain	(letters)
	<i>HbA1c</i> : 7.6 SD1.3–1.5; ≤8%		IVR0.3	+10.9 SD10.4	<0.0001vs C
	(65.8–67.5%); >8% (32.5–34.2%)		IVR0.5	+12.0 SD14.9	<0.0001 vs C
	Baseline VA: Mean ETDRS letter		С	+2.3 SD14.2	
	score 56.9–57.5		CMT:		
	<i>Baseline CMT</i> : 447.4–482.6 μm			Mean change	p Value
	Comorbidities: history of smoking			from baseline	
	33.6–51.6%		IVR0.3	-259.8 SD169.3	<0.0001 vs C
			IVR0.5	-270.7 SD201.6	<0.0001 vs C
			С	-125.8 SD198.3	

Injections are intravitreal unless otherwise noted. BCVA, best corrected visual acuity; C, control; CMT, central macular thickness; CSME, clinically significant macular oedema; DDS, dexamethasone; DIL, dexamethasone followed by laser; DM, diabetes mellitus; DMO, diabetic macular oedema; DP, diastolic pressure; DR, diabetic retinopathy; HR QoL, health-related quality of life; IOP, intraocular pressure; IV, intravitreal; IVB, intravitreal bevacizumab; IVP, intravitreal pegaptanib; IVR, intravitreal ranibizumab; IVT, intravitreal triamcinolone; IVTL, intravitreal triamcinolone; IVTL, intravitreal triamcinolone; IVTE, intravitreal veges Trap Eye; L, laser; MLT/MPC, macular laser therapy/macular photocoagulation; NEI VFQ-25, National Eye Institute Visual Function Questionnaire-25; NPDR, non-proliferative diabetic retinopathy; NR, not reported; OCT, optical coherence tomography; PDR, proliferative diabetic retinopathy; PRP, panretinal photocoagulation; RCT, randomised controlled trial; SOC, standard of care; SP, systolic pressure; SRFA, fluocinolone; VA, visual acuity; VEGF, vascular endothelia growth factor.

tudy	Participants and baseline values	Intervention	Outcome (change from baseline at study end)		
OLT Study Michaelides	N: 80 eyes of 80 patients Inclusion criteria: ≥18 years, type 1 or 2 DM, BCVA in	Group 1 (MLT, n=38 eyes): modified ETDRS macular laser	At 24 months BCVA (ETDRS):		
t al/Rajendram t al)) <sup>23 52 85</sup>	the study eye 35–69 ETDRS letters at 4 m (≥6/60 or ≤6/12), center-involving clinically significant DMO with	therapy; reviewed every 4 months up to 52 weeks;		BCVA. mean (SD)	p Value
<	CMT ≥270 µm; media clarity, papillary dilation and	retreatment performed if clinically		-0.5 (10.6)	
e <i>sign</i> : 2-arm CT	cooperation sufficient for adequate fundus imaging; a least 1 prior macular laser therapy; IOP <30 mm Hg;	indicated by ETDRS guidelines (median 4 laser treatments)	IVB	+8.6 (9.1)	0.005 vs MLT
ollow-up:	fellow eye BCVA ≥3/60; fellow eye received no	Group 2 (IVB, n=42 eyes):		BCVA gain	
months	anti-VEGF in past 3 months and no expectation of	1.25 mg (0.05 ml) IV		(letters)	
	such therapy	bevacizumab at baseline, 6 and	MLT	gaining	
	Exclusion criteria: (ocular for study eye) macular ischemia, macular oedema due to causes other than	12 weeks; subsequent IVB injections (up to 52 weeks)		≥10: 7% losing >15:	
	DMO, coexistent ocular disease affecting VA or DMO,	guided by an OCT-based		4%	
	any treatment for DMO in prior 3 months, PRP within	retreatment protocol (median 13	IVB	gaining	0.001 vs
	3 months prior to randomisation or anticipated, PDR, HbA1c >11%, medical history of chronic renal failure;	injections)  Laser modified ETDRS protocol,		≥10: 49% losing >15:	MLT 0.004 vs
	any thromboembolic event within 6 months prior to	retreatment by ETDRS		32%	MLT
	randomisation, unstable angina, evidence of active	guidelines		CMT (µm,	p Value
	ischemia on ECG; major surgery within 28 days of randomisation or planned; participation in an		MLT	<i>quartiles)</i> –118	
	investigational drug trial; systemic anti-VEGF or			SD171	
	pro-VEGF treatment within 3 months of enrolment;		IVB	–146 SD122	0.62 vs MLT
	pregnancy, lactation; intraocular surgery within 3 months of randomisation; aphakia; uncontrolled			SD 122	IVILI
	glaucoma; significant external ocular disease				
	Age: 64.2 SD8.8 years				
	Sex: 31% female Diabetes type: 90% type 2 DM, 10% type 1 DM				
	HbA1c: 7.5–7.6 SD1.2–1.4%				
	Baseline VA: ETDRS letter score 54.6–55.7				
	SD8.6–9.7 Baseline CMT: 481–507 SD121–145 μm				
	Comorbidities: 19% mild NPDR (level 35), 46%				
	moderate NPDR (level 43), 19% moderately severe				
	NPDR (level 47), 13% severe NPDR (level 53), 3% moderate PDR (level 65), 79–88% phakic				
m <i>et al<sup>35</sup></i>	N: 52 eyes of 52 patients	Group 1 (IVB1.25, n=26 eyes):	At 6 months		
ng Kong	Inclusion criteria: ≥18 years, type 1 or 2 DM, clinically	1.25 mg bevacizumab (0.05 ml)	BCVA (ETDRS chart):		
<i>sign</i> : 2-arm T	significant DMO (slit-lamp biomicroscopy, ETDRS criteria; leakage confirmed by fluorescein	Group 2 (IVB2.5, n=26 eyes): 2.5 mg bevacizumab (0.1 ml)			

Study	Participants and baseline values	Intervention	Outcome (change from baseline at study end)		
Follow-up: 6 months	angiography, CMT ≥250 µm on OCT), BCVA ≥1.3 ETDRS logMAR units; only patients with diffuse DMO recruited <i>Exclusion criteria</i> : macular oedema due to reasons other than diabetes, significant media opacities, macular ischemia of ≥1 disk area, vitreomacular traction, PDR, aphakia, glaucoma or ocular hypertension, previous anti-VEGF treatment, intraocular surgery except uncomplicated cataract	Regimen for all groups: 3 monthly IV injections, topical 0.5% levofloxacin 4×/day for up to 2 weeks after each injection	BCVA (logMAR) IVB1.25 IVB2.5	0.11 SD0.31 (+5.5 letters) 0.13 SD0.26 (+6.5 letters)	p Value 0.018 vs baseline, NS vs IVB2.5 0.003 vs baseline
extraction (but > 6 months prior), focal DMO, any lase procedure within previous 4 months, subtenon or intravitreal triamcinolone injection within 6 months, pregnancy  Age: 65.3 SD8.9 years  Sex: 46.2% female		CMT (OCT) IVB1.25	CMT (µm) 96	p Value 0.002 vs baseline, NS vs IVB2.5	
	Diabetes type: not reported  HbA1c: 7.5 SD1%		IVB2.5	74	0.013 vs baseline
	Baseline VA: 0.61 SD0.29 logMAR		Subgroups:		
Baseline CMT: 466 SD127 μm Comorbidities: not reported			► For patients with previous DMO treatment (mainly laser): no significant reduction in CMT at 6 months (452 µm at baseline to 416 µm at		
			6 months, p=0.22); no significant improvement in BCVA (0.66 logMAR at baseline to 0.56 logMAR at		
			6 months (+5 letters), p=0.074)		

Study	Participants and baseline values	Intervention	Outcome (change from baseline at study end)		
Faghihi <i>et al</i> <sup>53</sup>	N: 80 eyes of 40 patients	Group 1 (IVB, n=40 eyes):	At 6 months		
Iran	Inclusion criteria: Bilateral non-tractional CSME,	1.25 mg bevacizumab	Mean change in BCVA (ETDRS		
Design: 2-arm	10/10> V.A≥1/10, Controlled blood pressure.	Group 2 (IVB+MPC, n=40 eyes):	chart):		
RCT Follow-up:	Exclusion criteria: Advanced or advanced active PDR, significant cataract, glaucoma, history of recent	1.25 mg bevacizumab  Regimen for all groups: Eyes		BCVA (logMAR)	p Value
6 months	vascular accident (eg, MI, CVA), Previous treatment of CSME or PDR, or pharmacotherapy for CSME,	examined every 2 months and if evidence of CSME IVB was	IVB	0.138	<0.05 vs baseline
	macular ischemia and uncontrolled hypertension <i>Age</i> : 57.7±8 years	injected. Mean of the number of IVB injections in IVB group and	IVB+MPC	0.179	<0.05 vs baseline
	Sex: 27.5% females Diabetes type: NR	IVB+MPC group were 2.23±1.24 and 2.49±1.09, respectively	▶ no statistically significant difference between the two		
	HbA1c: 8.42±1.82 g/dl Baseline VA: 0.326–0.409 (SD 0.279–0.332)		groups CMT (OCT):		
	Baseline CMT: 277 um-287 um (SD 78-98)			CMT (µm)	p Value
	Comorbidities: not reported		IVB	<b>–39</b>	<0.05 vs baseline
			IVB+MPC	-39	<0.05 vs baseline
			➤ No statistically significant difference between the two		
			groups		

BCVA, best corrected visual acuity; C, control; CMT, central macular thickness; CSME, clinically significant macular oedema; DDS, dexamethasone; DIL, dexamethasone followed by laser; DM, diabetes mellitus; DMO, diabetic macular oedema; DP, diastolic pressure; DR, diabetic retinopathy; HR QoL, health-related quality of life; IOP, intraocular pressure; IV, intravitreal; IVB, intravitreal bevacizumab; IVP, intravitreal pegaptanib; IVP, intravitreal riamcinolone; IVTL, intravitreal triamcinolone plus laser; IVVTE, intravitreal VEGF Trap Eye; L, laser; MLT/MPC, macular laser therapy/macular photocoagulation; NEI VFQ-25, National Eye Institute Visual Function Questionnaire-25; NPDR, non-proliferative diabetic retinopathy; NR, not reported; OCT, optical coherence tomography; PDR, proliferative diabetic retinopathy; PRP, panretinal photocoagulation; RCT, randomised controlled trial; SOC, standard of care; SP, systolic pressure; SRFA, fluocinolone; VA, visual acuity; VEGF, vascular endothelia growth factor.

Continued

	Participants and baseline values	Intervention	Outcome (change from baseline at study end)		
egaptanib					
unningham et all	N: 172 eyes of 172 patients	Group 1 (IVP0.3, n=44	At 36 weeks		
damis <i>et al</i> <sup>39 57</sup>	Inclusion criteria: ≥18 years, type 1 or 2 DM, DMO	eyes): 0.3 mg IV	BCVA:		
SA	involving the center of the macula with corresponding	pegaptanib (90 µI) (median		BCVA (letters)	p Value
esign: 4-arm phase	leakage from microaneurysms, retinal telangiectasis,	5 injections (range 1-6))	IVP0.3	+4.7	0.04 vs C
RCT	or both; clear ocular media, BCVA letter scores	Group 2 (IVP1, n=44 eyes):	IVP1	+4.7	0.05 vs C
follow-up: 36 weeks			IVP3	+1.1	NS vs C
·	the fellow eye; IOP ≤23 mm Hg, focal	(median 6 injections	C	-0.4	
	photocoagulation could be safely deferred for	(range 3–6))		Plus ≥10 letters	
	16 weeks; no ECG abnormalities, no major serological		IVP0.3	34%	0.003 vs C
	abnormalities	3 mg IV pegaptanib (90 µI)	IVP1	30%	
	Exclusion criteria: history of panretinal or focal	(median 6 injections (range	IVP3	14%	
	photocoagulation; neodymium:yttrium-aluminum-	1–6))	C	10%	
	garnet laser or peripheral retinal cryoablation in	Group 4 (C, n=42 eyes):	CMT (OCT):		
	previous 6 months; any ocular abnormality interfering	sham injection (median 5		CMT	p Value
	with VA assessment or fundus photography;	injections (range 1–6))		(μm, 95% CI)	<b>P</b>
	vitreoretinal traction; vitreous incarceration; retinal vein		IVP0.3	-68.0 (-118.9 to	0.02 vs C
	occlusion involving the macula; atrophy/scarring/	injections at baseline, week		-9.88)	
	fibrosis or hard exudates involving the center of the	6 and week 12; thereafter,	IVP1	-22.7 (-76.9 to	NS vs C
	macula; history of intraocular surgery within previous	additional injections		+33.8)	
	12 months, myopia of ≥8 diopters, axial length of	administered every 6 weeks	IVP3	-5.3 (-63.0 to	NS vs C
	≥25 mm, likelihood of requiring panretinal	at the discretion of the	0	+49.5)	
	photocoagulation within following 9 months; cataract	investigators if judged	C	+3.7	
	surgery within 12 months; active ocular or periocular	indicated (maximum of 6	► Subgroups: of 16	10.7	
	infection; previous therapeutic radiation to the eye,	injections up to week 30);	participants with retinal		
	head, or neck; known serious allergies to fluorescein	laser photocoagulation	neovascularisation at		
	dye; HbA1c ≥13%, pregnancy	allowed after week 13 if	baseline, 8 of 13 (62%) in		
	Age: 61.3–64.0 SD9.3–10.1 years	judged indicated by the	the pegaptanib groups and		
	Sex: 45–55% female	study-masked	0 of 3 in the sham group		
	Diabetes type: 5–10% IDDM	ophthalmologist (25% for	had regression of		
	HbA1c: 7.1–7.7 SD1.2–1.6	IVP0.3, 30% for IVP1, 40%	neovascularisation at		
	Baseline VA: letter score 55.0–57.1 SD9.1–11.5	for IVP3, 48% for C)	36 weeks		
	Baseline CMT: 423.2–476.0 μm	101 141 3, 40 /6 101 0/	JU WEEKS		
	Comorbidities: not reported				

Table 5 Continued					
Study	Participants and baseline values	Intervention	Outcome (change from baseline at study end)		
Sultan et af <sup>40</sup> Multicenter international Design: 2-arm placebo-controlled RCT Follow-up: 2 years	N: 260 eyes of 260 patients Inclusion criteria: ≥18 years, type 1 or 2 DM, DMO involving the center of the macula not associated with ischemia, CMT ≥250 μm, BCVA letter score 65–35, IOP ≤21 mm Hg, clear ocular media Exclusion criteria: any abnormality other than DMO affecting VA assessment, vitreomacular traction;	Group 1 (IVP, n=133 eyes): 0.3 mg IV pegaptanib sodium (mean number of injections 12.7 SD4.6) Group 2 (C, n=127 eyes): sham injection (mean number of injections 12.9	At 1 year BCVA (ETDRS): IVP C IVP C	BCVA (letters) +5.2 +1.2 Plus ≥10 letters 36.8% 19.7%	<i>p Value</i> <0.05 vs C 0.0047 vs C
(primary efficacy endpoint at 1 year)	yttrium—aluminium—garnet laser, peripheral retinal cryoablation, laser retinopexy for retinal tears, focal or grid photocoagulation within prior 16 weeks; panretinal photocoagulation <6 months before baseline or likely			Increase in degree 4.1% 12.4%	<i>e by ≥2 steps</i> 0.047 vs C
	to be needed within 9 months; significant media opacities; intraocular surgery in prior 6 months; pathological high myopia; prior radiation in region of	investigator determination (ETDRS criteria), laser photocoagulation could be	IVP C	Decrease in degree 10.2% 3.1%	ee by ≥2 steps NS vs C
	study eye; history of severe cardiac or peripheral vascular disease, stroke in prior 12 months, major surgery in prior 1 month, treatment in prior 90 days with any investigational agent or with bevacizumab for any nonocular condition, HbA1c ≥10% or signs of uncontrolled diabetes, hypertension, known relevant	treatments per year) (laser treatments in 25.2% of IVP	IVP C At 2 years	Decrease ≥25%: 31.7% ≥50%: 14.6% ≥25%: 23.7% ≥50%: 11.9%	e in CMT NS vs C
	allergies; pregnant or lactating  Age: 62.3–62.5 SD9.3–10.2 years  Sex: 39–46% female  Diabetes type: 6.3–7.5% type 1 DM, 92.5–93.7% type 2 DM	group and 45% of C group); in year 2, injections as judged necessary	BCVA (ETDRS):  IVP C	BCVA (letters) +6.1 +1.3 Plus ≥10 letters	p Value <0.01 vs C
	HbA1c: 42.5–45.9% <7.6%, 54.1–57.5% >7.6%  Baseline VA: letter score 57.0–57.5 SD8.1–8.9  Baseline CMT: 441.6–464.6 SD135.5–148.5 μm		IVP C Retinopathy:	38.3% 30%	NS vs C
	Comorbidities: not reported		IVP C	Increase in degre 6.3% 13.8% Decrease in degre	NS vs C
			IVP C CMT (OCT):	16.3% 3.8%	0.03 vs C
			IVP	Decrease in CMT ≥25%: 40.4% ≥50%: 19.2%	NS vs C
			С	≥25%: 44.6% ≥50%: 26.1%	Continued

Table 5 Continued					
Study	Participants and baseline values	Intervention	Outcome (change from baseline at study end)		
			QoL:  ► NEI VFQ-25: between group differences not significant at 54 weeks; at 102 weeks, significantly greater improvement in composite score and subscales distance vision activities, social functioning and mental health with pegaptanib  ► EQ-5D: no significant differences between groups in EQ-5D scores at weeks 54 or 102		
II RCT	N: 221 eyes of 221 patients Inclusion criteria: aged >18 years and diagnosed with type 1 or 2 diabetes mellitus, with DMO involving the central macula defined as CRT (>250 um in the central subfield. Participants were required to have BCVA letter score at 4 m of 73–24. 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; panretinal or macular laser photocoagulation or use of intraocular or periocular corticosteroids or antiangiogenic drugs within 3 months of screening; vision decrease due to causes other than DMO; proliferative diabetic retinopathy (unless regressed and currently inactive); ocular inflammation; cataract or other intraocular surgery within 3 months of screening, laser capsulotomy within 2 months of screening, laser capsulotomy within 2 months of screening, aphakia; spherical equivalent of >8 diopters; or any concurrent disease that would compromise visual acuity or require medical or surgical intervention during the study period: active iris neovascularisation, vitreous hemorrhage, traction retinal detachment, or preretinal fibrosis involving the macula; visually significant vitreomacular traction or epiretinal	Trial of VEGF Trap-Eye (VTE), randomised on a 1:1:1:1:1 basis Group 1 (IVVTE1, n=44 eyes): IVVTE, 0.5 mg every 4 weeks Group 2 (IVVTE2, n=44 eyes): IVVTE, 2 mg every 4 weeks Group 3 (IVVTE3, n=42 eyes): IVVTE, 2 mg for 3 initial months then every 8 weeks Group 4 (IVVTE4, n=45 eyes): IVVTE, 2 mg for 3 initial months then as needed Group 5 (L, n=44 eyes): laser photocoagulation Laser modified ETDRS protocol	At 6 months  IVVTE1 IVVTE2 IVVTE3 IVVTE3 L  IVVTE1 IVVTE2 IVVTE3 IVVTE3 L  IVVTE1 IVVTE2 IVVTE3 IVVTE1 IVVTE2 IVVTE1 IVVTE2 IVVTE1 IVVTE2 IVVTE3 IVVTE3 IVVTE3 IVVTE3 IVVTE3	BCVA (letters) +8.6 +11.4 +8.5 +10.3 +2.5 plus ≥10 letters 50% 64% 43% 58% 32% CMT(um) -144.6 -194.5 -127.3 -153.3 -67.9  BCVA (letters) +11.0 +13.1 +9.7 +12.0	p Value 0.005 vs L <0.0001 vs L 0.008 vs L 0.0004 vs L  NR NR NR NR NR NR CO.0001 vs L <0.0001 vs L <0.0001 vs L ≤0.0001 vs L

Table 5 Co	ontinued				
Study	Participants and baseline values	Intervention	Outcome (change from baseline at study end)		
	membrane evident biomicroscopically or on OCT;		L	-1.3	
	history of idiopathicor autoimmune uveitis; structural			Plus ≥15 letters	
	damage to the center of the macula that is likely to		IVVTE1	40.9%	0.0031 vs L
	preclude improvement in visual acuity after the		IVVTE2	45.5%	0.0007 vs L
	resolution of macular oedema; uncontrolled glaucoma		IVVTE3	23.8%	0.1608 vs L
	or previous filtration surgery; infectious blepharitis,		IVVTE3	42.2%	0.0016 vs L
	keratitis, scleritis, or conjunctivitis; or current treatment		L	11.4%	
	for serious systemic infection: uncontrolled diabetes			Plus ≥10 letters	
	mellitus; uncontrolled hypertension; history of cerebral		IVVTE1	57%	0.0031 vs L
	vascular accident or myocardial infarction within		IVVTE2	71%	0.0007 vs L
	6 months; renal failure requiring dialysis or renal		IVVTE3	45%	0.1608 vs L
	transplant; pregnancy or lactation; history of allergy to		IVVTE3	62%	0.0016 vs L
	fluorescein or povidone iodine; only 1 functional eye		L		
	(even if the eye met all other entry criteria); or an			CMT(µm)	
	ocular condition in the fellow eye with a poorer		IVVTE1	-165.4	<0.0001 vs L
	prognosis than the study eye		IVVTE2	-227.4	<0.0001 vs L
	Age: 60.7–64.0 years (SD 8.1–11.5)		IVVTE3	-187.8	<0.0001 vs L
	Sex: % female 35.6-47.6%		IVVTE3	-180.3	<0.0001 vs L
	Diabetes type: percentage of type 2, 88.6–97.7%		L	-58.4	
	HbA1c: 7.85-8.10 (SD 1.71-1.94)				
	Baseline VA: 57.6-59.9 (SD 10.1-12.5)				
	Baseline CMT: 426.1–456.6 μm (SD 111.8–152.4)				
	Comorbidities: history of any cardiac disease was				
	twice as common in the VEGF Trap-Eye groups				
	compared with the laser group				

BCVA, best corrected visual acuity; C, control; CMT, central macular thickness; CSME, clinically significant macular oedema; DDS, dexamethasone; DIL, dexamethasone followed by laser; DM, diabetic macular oedema; DP, diastolic pressure; DR, diabetic retinopathy; HR QoL, health-related quality of life; IOP, intraocular pressure; IV, intravitreal; IVB, intravitreal bevacizumab; IVP, intravitreal pegaptanib; IVR, intravitreal triamcinolone; IVTL, intravitreal triamcinolone plus laser; IVVTE, intravitreal VEGF Trap Eye; L, laser; MLT/MPC, macular laser therapy/macular photocoagulation; NEI VFQ-25, National Eye Institute Visual Function Questionnaire-25; NPDR, non-proliferative diabetic retinopathy; NR, not reported; OCT, optical coherence tomography; PDR, proliferative diabetic retinopathy; PRP, panretinal photocoagulation; RCT, randomised controlled trial; SOC, standard of care; SP, systolic pressure; SRFA, fluocinolone; VA, visual acuity; VEGF, vascular endothelia growth factor.

Ford JA, Lois N, Royle P, et al. BMJ Open 2013;3:e002269. doi:10.1136/bmjopen-2012-002269

0. 1	<b>5</b>		Outcome (change from		
Study	Participants and baseline values	Intervention	baseline at study end)		
Dexamethasone Callanan et alUSA <sup>44</sup> Design: 2-arm RCT Follow-up: 12 months	N: 253 eyes of 253 patients Inclusion criteria: diffuse DMO, CMT ≥275 μm, BCVA ≥34 and ≤70 letters	Group 1 (DIL, n=126 eyes): dexamethasone IV implant followed by laser	At 12 months BCVA:	Plus ≥10	p Value
	Exclusion criteria: not reported  Age: not reported  Sex: not reported	photocoagulation after 1 month (mean 1.6 implants; 78.6% completion)	DIL L	letters (%) 28 24	NS vs L
	Diabetes type: not reported HbA1c: not reported Baseline VA: not reported Baseline CMT: not reported Comorbidities: not reported	Group 2 (L, n=127 eyes): laser alone (79.5% completion) Regimen for all groups: if needed, patients were retreated with the dexamethasone implant at months 6 or 9, and with laser at months 4, 7 and 10; mean 2.2 laser treatments per patient Laser protocol not reported	<ul> <li>▶ Patients in DIL group had significantly greater increases in BCVA from baseline than patients in the laser group (p&lt;0.05) at months 1–9 only CMT (OCT):</li> <li>▶ Patients in DIL group had significantly greater mean reductions from baseline in CMT at months 1 and 6 only (p&lt;0.001)</li> </ul>		
Haller <i>et al</i> <sup>59</sup> USA Multicenter	N: 171 eyes of 171 patients Inclusion criteria: ≥12 years, DMO persisting for ≥90 days after laser treatment or medical	Group 1 (DDS350, n=57 eyes): 350 μg dexamethasone IV drug delivery system, implanted into	At 90 days BCVA (ETDRS):	Plus ≥10	p Value
Design: 3-arm RCT Follow-up: 6 months (180 days), primary outcome 3 months (90 days)	therapy, BCVA by ETDRS between 20/40 (67 letters) and 20/200 (35 letters) due to clinically detectable DMO; analysis includes only eyes with DMO associated with DR <i>Exclusion criteria</i> : history of vitrectomy in the	the vitreous cavity  Group 2 (DDS700, n=57 eyes):  700 µg dexamethasone IV drug delivery system, implanted into the vitreous cavity	DDS350 DDS700 C CMT (OCT):	letters 21% (graph) 33% 12%	NS vs C 0.007 vs C
(oo dayo)	study eye; use of systemic, periocular, or intraocular steroids within 30 days of enrolment; moderate or severe glaucoma in the study eye; poorly controlled hypertension (SP >160 mm Hg	Group 3 (C, n=57 eyes): no treatment  Regimen for all groups: eyes demonstrating a VA loss of ≥5	DDS350	<i>CMT (µm)</i> -42.57 SD95.96	p Value NS (p=0.07) vs C
	or DP >90 mm Hg); poorly controlled diabetes (HbA1c >13%)  Age: 62.9–63.8 years SD10.2–12.0  Sex: 45.6–49.1% female	letters could be treated with any other therapy (including laser photocoagulation and IV triamcinolone) (n=4 with	DDS700 C	-132.27 SD160.86 +30.21 SD82.12	<0.001 vs C
	Diabetes type: not reported HbA1c: 7.3–7.6% Baseline VA: letter score 54.4–54.7	photocoagulation or IV triamcinolone in the C group, n=2 in the DDS350 group, none	At 180 days BCVA (ETDRS):	Plus ≥10	p Value
	SD9.96–11.88 <i>Baseline CMT</i> : 417.5–446.5 μm SD123.7–155.9	in the DDS700 group)	DDS350	letters 20% (graph)	NS vs C
	Comorbidities: 19–21% prior cataract extraction		DDS700 C	33% (graph) 23% (graph)	NS vs C

Table 6 Continued					
Study	Participants and baseline values	Intervention	Outcome (change from baseline at study end)		
Fluocinolone FAME Study (Campochiaro et all Campochiaro et al) Campochiaro et al) Multicenter international Design: 3-arm placebo-controlled RCT Follow-up: 24 months; abstract with 36 month outcomes	N: 956 eyes of 956 patients Inclusion criteria: DMO, CMT ≥250 μm despite at least 1 prior focal/grid macular laser photocoagulation treatment, BCVA ETDRS letter score between 19 and 68 (20/50–20/400)  Exclusion criteria: glaucoma, ocular hypertension, IOP >21 mm Hg, taking IOP lowering drops; laser treatment for DMO within 12 weeks of screening, any ocular surgery in the study eye within 12 weeks of screening; ocular or systemic steroid therapy; active ocular infection; pregnancy Age: 62.5 SD9.4 years Sex: 40.6% Diabetes type: 6.6% type 1 DM, 92% type 2 DM, 1.4% uncertain HbA1c: 7.8 SD1.59% Baseline VA: ETDRS letter score 53.4 SD12.23 Baseline CMT: 469.0 SD164.78 μm Comorbidities: 47.1% cataract at baseline, 62.7–67.4% phakic	Group 1 (0.5, n=375 eyes): intravitreal insert releasing 0.2 μg/day fluocinolone acetonide (FA) (2, 3, or 4 treatments received by 21.3, 1.9 and 0.3%)  Group 2 (SRFA0.5, n=393 eyes): intravitreal insert releasing 0.5 μg/day fluocinolone acetonide (2, 3, or 4 treatments received by 22.6, 2.5 and 0.3%)  Group 3 (C, n=185 eyes): sham injection (2, 3, or 4 treatments received by 19.5, 2.7 and 1.6%)  Regimen for all groups: patients could receive rescue focal/grid laser therapy any time after the first 6 weeks for persistent oedema (35.2–36.7% in FA groups, 58.9% control group, p<0.001); treatments were allowed every 3 months for persistent or recurrent oedema; patients eligible for another FA insert at 1 year if ≥5 letter	At 24 months BCVA (ETDRS):  SRFA0.2 SRFA0.5 C  SRFA0.5 C Subgroups: ▶ BCVA benefits only in pseudophakic eyes (cataract surgery before or during the study), in phakic eyes, BCVA letter score was reduced by 5 (high dose) and 9 (low dose) from baseline at 24 months  CMT (optical coherence tomography):	BCVA (letters) +4.4 +5.4 +1.7 Plus ≥15 letters (%) 29 29 16	p Value 0.02 vs C 0.017 vs C p Value 0.002 SRFA vs C
		reduction in BCVA or >50 μm CMT increase from best status	SRFA0.2 SRFA0.5  C  ► effect maintained at 36 months	-167.8 -177.1 -111.3	0.005 vs C <0.001 vs C
			At 36 months  SRFA0.2/0.5  C	Plus ≥15 letters 28.7% 18.9%	p Value 0.018 SRFA vs C
				10.9 /0	Continued

Study	Participants and baseline values	Intervention	Outcome (change from baseline at study end)		
Pearson et al <sup>43</sup>	N: 196 patients	Group 1 (SRFA, n=127): 0.5 mg	At 3 years		
USA	Inclusion criteria: persistent or recurrent	sustained release fluocinolone	BCVA:		
Multicenter	unilateral or bilateral DMO with retinal thickening	acetonide intravitreal implant		Gain ≥15	p Value
Design: 2-arm RCT	involving fixation of $\geq 1$ disc area in size, ETDRS	Group 2 (SOC, n=69): standard		letters	
Follow-up: 36 months	visual acuity of ≥20 letters (20/400) to ≤68	of care—either repeat laser or	SRFA	31%	NS
	letters (20/50) and $\geq$ 1 macular laser treatment in		SOC	20%	
	the study eye more than 12 weeks prior to	Laser ETDRS protocol		Loss ≥15	
	enrolment			letters	
	Exclusion criteria: Ocular surgery within		SRFA	17%	NS
	3 months prior to enrolment, uncontrolled IOP		SOC	14%	
	within the past 12 months while on ≥1		CMT:		
	antiglaucoma medication, IOP of ≥22 mm Hg at			Mean change	p Value
	screening while on ≥1 antiglaucoma medication,			in baseline	
	peripheral retinal detachment in the area of		0054	CMT	NO
	implantation or media opacity precluding		SRFA	-86 -410	NS
	diagnosis of status in the study eye		SOC	<b>–110</b>	
	Age: 61.4–62.7 years				
	Sex: 41.7–42% female				
	Diabetes type: 62.3–70% on insulin				
	HbA1c: not reported				
	Baseline VA: not reported Baseline CMT: not reported				
	Comorbidities: not reported				

BCVA, best corrected visual acuity; C, control; CMT, central macular thickness; CSME, clinically significant macular oedema; DDS, dexamethasone; DIL, dexamethasone followed by laser; DM, diabetes mellitus; DMO, diabetic macular oedema; DP, diastolic pressure; DR, diabetic retinopathy; HR QoL, health-related quality of life; IOP, intraocular pressure; IV, intravitreal; IVB, intravitreal bevacizumab; IVP, intravitreal pegaptanib; IVR, intravitreal riamcinolone; IVTL, intravitreal triamcinolone plus laser; IVVTE, intravitreal VEGF Trap Eye; L, laser; MLT/MPC, macular laser therapy/macular photocoagulation; NEI VFQ-25, National Eye Institute Visual Function Questionnaire-25; NPDR, non-proliferative diabetic retinopathy; NR, not reported; OCT, optical coherence tomography; PDR, proliferative diabetic retinopathy; PRP, panretinal photocoagulation; RCT, randomised controlled trial; SOC, standard of care; SP, systolic pressure; SRFA, fluocinolone; VA, visual acuity; VEGF, vascular endothelia growth factor.

Table 7 Triamcinolone	studies					
	Participants and baseline		Outcome (change from			
Study	values	Intervention	baseline at study end)			
	N: 840 eyes of 693 patients	Group 1 (IVT1, n=256 eyes):	At 2 years			
et al/Beck et al/Bressler	Inclusion criteria: >18 years, type		BCVA (E-ETDRS):			
et al) <sup>22 61 63 64</sup>	1 or 2 DM, study eye: (1) BCVA	(3.5 treatments)		BCVA (letters)		p Value
USA	(E-ETDRS) between 24 and 73	Group 2 (IVT4, n=254 eyes):	IVT1	–2 SD18		0.02 vs L
Multicenter	(20/320 and 20/40), (2) retinal	4 mg IV triamcinolone				NS vs IVT4
Design: 3-arm RCT	thickening due to DMO involving	(3.1 treatments)	IVT4	-3 SD22		0.002 vs L
Follow-up: 2 years,	the center of the macula main	Group 3 (L, n=330 eyes):	L	+1 SD17		
additional 3 year	cause for visual loss, (3) CMT	focal/grid photocoagulation	W. 477.4	BCVA gain categorie	es	
follow-up	≥250 µm, (4) no expectation of	(2.9 treatments)	IVT1	+10 or more: 25%		0.03 vs L, NS vs
	scatter photocoagulation within	Regimen for all groups:		+9 to -9: 50%		IVT4
	4 months	retreatment protocol: where	N/T4	-10 – more: 26%		0.04
	Exclusion criteria: any prior	indicated, retreatment was	IVT4	+10 or more: 28%		0.01 vs L
	treatment with IV corticosteroids,	performed within 4 weeks		+9 to -9: 44%		
	peribulbar steroid injection within	•	,	-10 or more: 28%		
	prior 6 months, photocoagulation	sooner than 3.5 months from	L	+10 or more: 31%		
	for DMO within prior 15 weeks,	the time of last treatment;		+9 to -9: 50%		
	panretinal scatter	eyes were generally retreated	Cubarauna	-10 or more: 19%		
	photocoagulation within prior	unless: (1) little or no oedema	Subgroups:  Similar results when			
	4 months, pars plana vitrectomy, history of open-angle glaucoma	involving the center of the	considering only pseudophakic			
	or steroid-induced	macula present and CMT	eyes or eyes with minimal			
	IOP elevation requiring	≤225 µm, (2) VA letter score	cataract no substantially			
	IOP-lowering treatment, and IOP	>79 (20/25 or better),	different results based on			
	≥25 mm Hg	(3) substantial improvement in	baseline VA, baseline CMT,			
	Age: 63 SD9 years	macular oedema since last	history of focal/grid			
	Sex: 49% female	treatment (eg, ≥50%	photocoagulation for DMO			
	Diabetes type: 95% type 2 DM,	decrease in CMT), (4)	▶ 3 year results consistent with 2			
	5% type 1 DM	clinically significant adverse	year results for BCVA and			
	HbA1c: 7.9 SD1.8%	effect from prior treatment,	CMT			
	Baseline VA: ETDRS letter score		CMT (OCT):			
	59 SD11 (~20/63)	deemed futile (<5 letter	(22.)	CMT (µm)		p Value
	Baseline CMT: 24 SD130 µm	improvement in VA letter	IVT1	-86 SD167		<0.001 vs L,
	Comorbidities: 21%	score or lack of CMT				NS vs IVT4
	pseudophakic, 2% ocular	reduction) and (6) for laser	IVT4	-77 SD160		<0.001 vs L
	hypertension, 7% mild NPDR,	group, complete focal/grid	L	-139 SD148		
	13% moderate NPDR, 40%	photocoagulation already	Progression of retinopathy:			
	moderately severe NPDR, 11%	given, with no areas identified	-	2 years	3 years	p Value
	severe NPDR, 23.5% mild to	for which additional treatment	IVT1	-	35 <sup>%</sup>	
	moderate, 3% high risk PDR	was indicated	IVT4		30%	<0.05 vs L
	•	Laser Modified ETDRS	L		37%	
						Continued

Study	Participants and baseline values	Intervention	Outcome (change from baseline at study end)		
		protocol as used in prior DRCR.net protocols			
Gillies <i>et al</i>	N: 69 eyes of 43 patients	Group 1 (IVT, n=34 eyes):	At 2 years		
Sutter <i>et al</i> 32 136–138	Inclusion criteria: patients with	4 mg (0.1 ml) IV triamcinolone			
Australia	persistent (≥3 months after	acetonide (mean 2.6	- ( /	BCVA (letters)	p Value
Design: 2-arm	adequate laser treatment) DMO	injections over 2 years)	IVT	+3.1	0.01 vs C
lacebo-controlled RCT	involving the central fovea,	Group 2 (C, n=35 eyes):	C	<b>-2.9</b>	
Follow-up: 2 years,	BCVA in the affected eye ≤6/9	placebo injection		CVA gain categories	
dditional 3-year	Exclusion criteria: uncontrolled	(subconjunctival saline	IVT	+10 or more: 21%	0.013 vs C
ollow-up	glaucoma, loss of vision due to	injection) (mean 1.8 injections		+9 to -9: 70%	
'	other causes, systemic treatment	over 2 years)		-10 or more: 9%	
	with >5 mg prednisolone (or	Regimen for all groups:	C	+10 or more: 12%	
	equivalent) daily, intercurrent	retreatment considered at		+9 to -9: 62%	
	severe systemic disease, any	each visit as long as		-10 or more: 25%	
	condition affecting follow-up or	treatments were at least	CMT (OCT):		
	documentation	6 months apart (retreatment if	, ,	CMT (µm)	p Value
	Age: 62.4–69.6	VA decreased ≥5 letters from	IVT	_125	0.009 vs C,
	SD9.2-12.5 years	previous peak value and			difference
	Sex: 52% female	persistent CMT >250 µm), if			between groups
	Diabetes type: not reported	no improvement after			59 µm (95% ČI
	HbA1c: 7.63-8.28 SD1.12-1.41	4 weeks, further laser			15 to 104)
	Baseline VA: ETDRS letter score	treatment was applied (n=1	C	<b>–7</b> 5	,
	60.5-61.3 SD11.9-13.2	laser treatment in intervention			
	Baseline CMT: 439-444	group, n=16 in placebo group,			
	SD101–125 µm	p=0.0001)			
	Comorbidities: 25%	Laser ETDRS protocol			
	pseudophakic				

tudy	Participants and baseline values	Intervention	Outcome (change from baseline at study end)		
Gillies <i>et al<sup>83</sup></i> Australia <i>Design</i> : 2-arm RCT <i>Follow-up</i> : 24 months	N: 84 eyes of 54 patients Inclusion criteria: DMO involving the central fovea, CMT ≥250 μm, BCVA 17–70 letters (~20/40–20/ 400), laser treatment could be safely delayed for 6 weeks		At 24 months BCVA (ETDRS): ITL L	BCVA (letters) +0.76 -1.49 BCVA gain	<i>p Value</i> NS vs L
	without significant adverse effects  Exclusion criteria: uncontrolled glaucoma, controlled glaucoma but with a glaucomatous visual field defect, loss of vision resulting from other causes, systemic treatment with >5 mg prednisolone (or equivalent) daily, retinal laser treatment within 4 months, intraocular	Group 2 (L, n=42 eyes): sham injection followed by laser treatment (at least 1 retreatment in 2nd year in 45%)  Regimen for all groups: retreatment with injection followed by laser at discretion of chief investigator, with at least 6 weeks between treatments; no retreatment if:	IVTL  Subgroups: ▶ BCVA outcome not significantly affected by cataract surgery	categories +10 or more: 36% +9 to -9: 31% -10 or more: 33% +10 or more: 17% +9 to -9: 59% -10 or more: 24%	0.049 vs L
	surgery within 6 months, concurrent severe systemic disease, any condition affecting follow-up or documentation Age: 65.4–66.9 SD8.9–9.5 years Sex: 38.1–47.6% female Diabetes type: not reported HbA1c: 7.81–8.02 SD1.44–1.63% Baseline VA: letter score 55.2–55.5 SD11.3–12.5 Baseline CMT: 482.1–477.4 SD122.7–155.5 µm Comorbidities: not reported	(1) investigator considered the macula nearly flat and CMT <300 µm; (2) VA was ≥79 letters (20/25) or VA had improved by ≥5 letters compared with the best VA after treatment or baseline acuity; (3) laser treatment was considered by the investigator as inappropriate or had no potential for improvement	~ .	CMT (μm) -137.1 -109.6	p Value NS vs L

Kin et alf s Korea Inclusion criteria: diffuse DMO Explain: 2-arm RCT Follow-up: 3 years Follow-up: 4 years Follow-up: 5 years Follow-up: 5 years Follow-up: 6 years	Study	Participants and baseline values	Intervention	Outcome (change from baseline at study end)		
Design: 2-arm RCT Pollow-up: 3 years Age: not reported Sex: not reported Diabetes type: not reported Baseline CMT: and the protection of the protect Baseline CMT: not reported Baseline CMT: and the protection of the protection	Kim <i>et al</i> <sup>45</sup>	N: 86 eyes of 75 patients	Group 1 (IVT, n=38 eyes):	At 3 years		
Sex. not reported Sex. not reported Diabetes type: not reported Baseline V4: not reported Baseline V2: not reported Baseline V3: not reported Combibilities: not reported Combibilities: not reported Baseline V3: not reported Baseline V3: not reported Combibilities: not	Korea	Inclusion criteria: diffuse DMO	4 mg IV triamcinolone (1.88	BCVA: not reported		
Sex not reported Diabetes type: not reported HbA1c: not reported Baseline CMT: not reported Baseline VMT: not reported CMT: not reported Baseline VMT: not reported Baseline VMT: not reported Baseline CMT: not reported CMT: not reported CMT: not reported Baseline CMT: not reported Baseline CMT: not reported Baseline CMT: 385-424 Baseline VMT: not reported CMT: not reported Baseline CMT: 385-424 Baseline VMT: not reported CMT: not reported Baseline CMT: 385-424 Baseline VMT: not reported CMT: not reported Baseline CMT: 385-424 Baseline VMT: not reported CMT: not reported Baseline CMT: 385-424 Baseline C	Design: 2-arm RCT	Exclusion criteria: not reported	additional treatments,	Outcomes related to DMO:		
Diabetes type: not reported HbA1c: not reported Baseline VA: not reported Baseline VA: not reported Comorbidities: not reported Completion 77:1%)  An In It eyes of 111 patients Inclusion criteria: 18 years, type Jesign: 3-am RCT 1 or 2 bM, clinically significant Comorbidities: not reported Caser protocol not	Follow-up: 3 years	Age: not reported	completion 68.1%)		No DMO	p Value
HbA1c: not reported Baseline VA: not reported Baseline VA: not reported Baseline CMT: not reported Comorbidities: not not not not not not not		Sex: not reported	Group 2 (IVTL, n=48 eyes):		recurrence	
Baseline VX: not reported Baseline CMT: not reported Comorbidities: not reported Como				IVT	3.9%	
Baseline CMT: not reported Comorbidities: not reported Camer at all P <sup>4</sup> Are 111 eyes of 111 patients Corbinate Parameter and the production of the provided Laser protocol not reported Camer at all P <sup>4</sup> Are 111 eyes of 111 patients Corbinate Parameter and the production of the provided Laser protocol not reported Camer at all P <sup>4</sup> Are 111 eyes of 111 patients Corbinate Parameter Pa				IVTL		0.028 vs IVT
Comorbidities: not reported completion 77.1%)			after 4 mg IV triamcinolone		•	
Regimen for all groups: additional treatment possible, criteria not mentioned Laser protocol not reported Group 1 (IVT, n=38 eyes): At 6 months Inclusion criteria: = 18 years, type lesign: 3-arm RCT 1 or 2 DM, clinically significant Follow-up: 6 months 2 years planned) Exclusion criteria: macular oedema due to causes other than diabetic maculopathy, signs of vitreomacular traction, proliferative diabetic retinopathy, aphakia, history of glaucoma or coular hypertension, macular ischemia, any laser procedure within 3 months, significant media opacities Age: 64.7–67.2 SD8.2–10.3 years Age: 64.7–67.2 SD8.2–10.3 years Sex: 42–59% female Diabetes type: not reported Baseline VX: ETDRS logMAR 0.64-0.72 SD0.34-0.36 Baseline CMT: 385-424 SD91-108 μm Comorbidities: 66-84% phakic  Age: 66-84% phakic  Regimen for all groups: At 6 months BCVA (ETDRS): BCVA		Baseline CMT: not reported	(0.92 additional treatments,	IVT	10.33 months	
additional treatment possible, criteria on treentioned Laser protocol not reported Group 1 (IVT, n=38 eyes): At 6 months long Kong Inclusion criteria: >18 years, type of 111 patients of 1 or 2 DM, clinically significant rollow-up: 6 months long Kong Vears planned)  2 years planned)  DMO (ETDRS), CMT ≥250 µm  Codema due to causes other than diabetic maculopathy, signs of vitreomacular traction, proliferative diabetic retinopathy, aphakia, history of glaucoma or ocular hypertension, macular ischemia, any laser procedure within 3 months, ocular surgery within 6 months, significant media opacities  Age: 64.7–67.2 SD8.2–10.3 years  Sex. 42–59% female  Diabetes type: not reported HbA1c: not reported Baseline VA: ETDRS logMAR  0.64–0.72 SD0.34–0.36 Baseline CMT: 385–424 SD91–108 µm Amaditional treatment possible, criterianent one of reproted Case of recurrence or prict of all roll of the protocol of the mentioned Carbor to the protocol of the mentioned Carbor (PLT). The 38 eyes): At 6 months BCVA (ETDRS):  Improvement  Indicate (PLT) = 0.7 SD 10.7 log (PV talled):  Improvement  Improvem		Comorbidities: not reported	completion 77.1%)	IVTL	19.88 months	0.027 vs IVT
criteria not mentioned Laser protocol not reported not no			· · · · · · · · · · · · · · · · · · ·			
Laser protocol not reported $Group \ 1 \ (IVT, n=38 \ eyes)$ : $At 6 \ months$ $Group \ 1 \ (IVT, n=38 \ eyes)$ : $At 6 \ months$ $Group \ 1 \ (IVT, n=38 \ eyes)$ : $At 6 \ months$ $Group \ 1 \ (IVT, n=38 \ eyes)$ : $At 6 \ months$ $Group \ 1 \ (IVT, n=38 \ eyes)$ : $Group \ 2 \ (IVT, n=36 \ eyes)$ : $Group \ 3 \ (IVT, n=36 \ eyes)$ : $Group \ 3 \ (IVT, n=36 \ eyes)$ : $Group \ 3 \ (IVT, n=36 \ eyes)$ : $Group \ 3 \ (IVT, n=36 \ eyes)$ : $Group \ 3 \ (IVT, n=36 \ eyes)$ : $Group \ 3 \ (IVT, n=36 \ eyes)$ : $Group \ 3 \ (IVT, n=36 \ eyes)$ : $Group \ 3 \ (I$			· · · · · · · · · · · · · · · · · · ·			
Am et al β <sup>4</sup> In long Kong Inclusion criteria: -18 years, type Jesign: 3-arm RCT Follow-up: 6 months 2 years planned)  Exclusion criteria: -18 years, type Exclusion criteria: -18 years, type Jesign: 3-arm RCT Follow-up: 6 months 2 years planned)  Exclusion criteria: -18 years, type Exclusion criteria: -18 years, type Level years planned)  Exclusion criteria: -18 years, type Exclusion criteria: -18 years, type Level years planned)  Exclusion criteria: -18 years, type Exclusion criteria: -18 years, type Level years planned)  Exclusion criteria: -18 years, type Level years planned)  Exclusion criteria: -18 years, type Level years planned)  Exclusion criteria: -18 years, type Invited years Invite						
Inclusion criteria: >18 years, type   4 mg IV triamcinolone (no posign: 3-arm RCT   1 or 2 DM, clinically significant retreatments)   2 years planned)   DMO (ETDRS), CMT ≥250 μm   Exclusion criteria: macular oedema due to causes other than diabetic maculopathy, signs of vitreomacular traction, proliferative diabetic retinopathy, aphakia, history of glaucoma or ocular hypertension, macular ischemia, any laser procedure within 6 months, significant media opacities   Age: 64.7-67.2 SD8.2-10.3 years   Sex. 42-59% female   Diabetes type: not reported   Baseline VA: ETDRS logMAR   0.64-0.72 SD0.34-0.36   Baseline CMT: 385-424   SD91-108 μm   Comorbidities: 66-84% phakic   Laser ETDRS protocol   SCM (VITL, n=36 eyes):   improvement   BCVA (ETDRS):   improvement   DIAD (ETDRS):   improvement   PV (IVTL   −0.7 SD 10.7 log   NS between   provided   MAR   groups   IVT   −1.1 SD 10.8 log   MAR   macular oedema had reduced   IVTL   −1.1 SD 10.8 log   MAR   center or at 1 to 2 months   Sw. 250 μm at the foveal   center or at 1 to 2 months   IVTL   −1.6 SD 11.5 log   MAR   center or at 1 to 2 months   IVTL   −1.6 SD 11.5 log   MAR   center or at 1 to 2 months   IVTL   −1.6 SD 11.5 log   MAR   center or at 1 to 2 months   IVTL   −1.6 SD 11.5 log   MAR   center or at 1 to 2 months   IVTL   Sw. 250 tenter   IVTL   Sw.	0.4					
Design: 3-arm RCT 1 or 2 DM, clinically significant Proflow-up: 6 months DMO (ETDRS), CMT ≥250 μm Group 2 (IVTL, n=36 eyes):  2 years planned)  Exclusion criteria: macular oedema due to causes other than diabetic maculopathy, signs of vitreomacular traction, proliferative diabetic retinopathy, aphakia, history of glaucoma or ocular hypertension, macular ischemia, any laser procedure within 3 months, social resurgery within 6 months, significant media opacities  Age: 64.7–67.2 SD8.2–10.3 years  Sex: 42–59% female  Diabetes type: not reported  Baseline VA: ETDRS logMAR  0.64–0.72 SD0.34–0.36 Baseline CMT: 385–424 SD91–108 μm  Comorbidities: 66–84% phakic  Pus ≥15 letters: improvement  At mg IV triamcinolone IVT  IVT  IVT  IVT  IVT  IVT  IVT  IVT						
DMO (ETDRS), CMT $\geq$ 250 µm   Exclusion criteria: macular oedema due to causes other than diabetic maculopathy, signs of vitreomacular traction, proliferative diabetic retinopathy, aphakia, history of glaucoma or ocular hypertension, macular ischemia, any laser procedure within 3 months, ocular surgery within 6 months, significant media opacities   Age: 64.7–67.2 SD8.2–10.3 years   Sex: 42–59% female   Diabetes type: not reported   Baseline VA: ETDRS logMAR   SD91–108 µm   Comorbidities: 66–84% phakic   Laser ETDRS protocol	0 0		· · · · · · · · · · · · · · · · · · ·	BCVA (ETDRS):		
Exclusion criteria: macular oedema due to causes other than diabetic maculopathy, signs of vitreomacular traction, proliferative diabetic retinopathy, aphakia, history of glaucoma or ocular hypertension, macular ischemia, any laser procedure within 3 months, ocular surgery within 6 months, significant media opacities Age: 64.7–67.2 SD8.2–10.3 years  Sex: 42–59% female Diabetes type: not reported HbA1c: not reported Baseline VA: ETDRS logMAR 0.64–0.72 SD0.34–0.36 Baseline CMT: 385–424 SD91–108 µm 4 months SCM: 66–84% phakic  Age: 66–84% phakic  Laser ETDRS protocol			,			p Value
oedema due to causes other than diabetic maculopathy, signs of vitreomacular traction, proliferative diabetic retinopathy, aphakia, history of glaucoma or ocular hypertension, macular ischemia, any laser procedure within 3 months, ocular surgery within 6 months, significant media opacities laser photocoagulation (n=3					•	
than diabetic maculopathy, signs of vitreomacular traction, proliferative diabetic retinopathy, aphakia, history of glaucoma or ocular hypertension, macular ischemia, any laser procedure within 3 months, ocular surgery within 6 months, significant media opacities apears  Age: 64.7–67.2 SD8.2–10.3 retreatments)  Sex: 42–59% female  Diabetes type: not reported  Baseline VA: ETDRS logMAR  0.64–0.72 SD0.34–0.36 according to study group, at ISD91–108 μm  Comorbidities: 66–84% phakic  bi to czeou adate the the condition (ETDRS)  (laser treatment once the condition (ETDRS)  5%  (laser treatment once the condition (ETDRS)  5%  (laser treatment once the condition (and the follows)  5%  macular oedema had reduced IVTL  -1.1 SD 10.8 log  MAR  Plus ≥15 letters:  3%  4MAR  -1.6 SD 11.5 log  MAR  Plus ≥15 letters:  3%  6700 3 (L, n=37 eyes): grid  MAR  Plus ≥15 letters:  5%  FOUT (OCT):  5%  FOUT (OCT):  CMT (OCT):  Sex: 42–59% female  Plus ≥15 letters:  3%  64-10.1 SD 10.8 log  MAR  FOUR > 1.5 log  MAR  FULL > 1.5 log  FULL >	2 years planned)			IVT		
of vitreomacular traction, proliferative diabetic retinopathy, aphakia, history of glaucoma or ocular hypertension, macular center or at 1 to 2 months ischemia, any laser procedure within 3 months, ocular surgery within 6 months, significant media opacities laser photocoagulation (n=3 $Age$ : 64.7–67.2 SD8.2–10.3 retreatments) $CMT$ ( $CCT$ ):  Sex: 42–59% female $Regimen$ for all groups: in $CMT$ ( $CCT$ ):  Sex: 42–59% female $Regimen$ for all groups: in $CMT$ ( $CCT$ ):  Sex: 42–59% female $Regimen$ for all groups: in $CMT$ ( $CCT$ ):  Baseline $VA$ : ETDRS logMAR oedema, retreatment offered $CA$ ( $CCT$ ):  Baseline $CAT$ ( $CCT$ ):  Baseline $CAT$ ( $CCT$ ):  Baseline $CAT$ ( $CCT$ ):  Social Signature $CAT$ ( $CCT$ ):  Social Sig						groups
macular oedema had reduced /VTL			• • • • • • • • • • • • • • • • • • • •			
aphakia, history of glaucoma or ocular hypertension, macular ischemia, any laser procedure after injection, whichever was within 3 months, ocular surgery within 6 months, significant media opacities as reprotocoagulation (n=3)  Age: 64.7-67.2 SD8.2-10.3 retreatments) (no years retreatments)  Sex: 42-59% female Regimen for all groups: in CMT (OCT):  Sex: 42-59% female Regimen for all groups: in CMT (µm) p Value  Diabetes type: not reported case of recurrence or IVT 342 SD124 (-54) NS between HbA1c: not reported persistence of macular oedema, retreatment offered  0.64-0.72 SD0.34-0.36 according to study group, at IVTL 307 SD181 (-116) <0.01 vs Baseline CMT: 385-424 intervals no less than SD91-108 μm 4 months L 350 SD169 (-35)  Comorbidities: 66-84% phakic Laser ETDRS protocol			`	NE		
ocular hypertension, macular ischemia, any laser procedure after injection, whichever was within 3 months, ocular surgery within 6 months, significant media opacities agree photocoagulation (n=3 plus $\geq$ 15 letters: $\geq$ 16 months, significant media opacities alaser photocoagulation (n=3 plus $\geq$ 15 letters: $\geq$ 16 months, significant media opacities alaser photocoagulation (n=3 plus $\geq$ 15 letters: $\geq$ 17 letters: $\geq$ 18 plus $\geq$ 19 letters: $\geq$ 19 letters: $\geq$ 19 letters: $\geq$ 19 letters: $\geq$ 10 months media opacities are retreatments) (no see that the sum of the su				IVIL	_	
ischemia, any laser procedure within 3 months, ocular surgery within 6 months, significant media opacities laser photocoagulation (n=3 $Age$ : 64.7–67.2 SD8.2–10.3 retreatments) (no years (MT ( $\mu$ ) $\mu$ ) $\mu$ 0 $\mu$ 1 $\mu$ 1 $\mu$ 2 $\mu$ 3 $\mu$ 3 $\mu$ 4 $\mu$ 4 $\mu$ 5 $\mu$ 5 $\mu$ 5 $\mu$ 6 $\mu$ 6 $\mu$ 8 $\mu$ 8 $\mu$ 9			· · · · · · · · · · · · · · · · · · ·			
within 3 months, ocular surgery within 6 months, significant Group 3 (L, n=37 eyes): grid media opacities laser photocoagulation (n=3 Plus ≥15 letters: 5% years retreatments) (no years retreatments) CMT (OCT):  Sex: 42–59% female Regimen for all groups: in CMT (µm) p Value Diabetes type: not reported case of recurrence or IVT 342 SD124 (−54) NS between HbA1c: not reported persistence of macular groups, <0.01 Baseline VA: ETDRS logMAR oedema, retreatment offered 0.64–0.72 SD0.34–0.36 according to study group, at IVTL 307 SD181 (−116) <0.01 vs Baseline CMT: 385–424 intervals no less than SD91–108 μm 4 months L 350 SD169 (−35) Comorbidities: 66–84% phakic Laser ETDRS protocol						
within 6 months, significant media opacitiesGroup 3 (L, n=37 eyes): grid laser photocoagulation (n=3MAR Plus ≥15 letters:Age: 64.7–67.2 SD8.2–10.3 yearsretreatments) (no retreatments)5%Sex: 42–59% femaleRegimen for all groups: in Diabetes type: not reported persistence of macular Baseline VA: ETDRS logMAR 0.64–0.72 SD0.34–0.36 Baseline CMT: 385–424 SD91–108 μm SD91–108 μm SD91–108 μm Amonths Amonths Amonths Laser ETDRS protocolIVT VAIVE VAIVE VAIVE AMAR CMT (OCT): CMT (μm) A42 SD124 (-54) A24 SD124 (-54) NS between groups, <0.01 baseline A22 SD124 (-54) NS between groups, <0.01 baseline			The state of the s	,		
media opacities laser photocoagulation (n=3 Plus $\geq$ 15 letters: Age: 64.7–67.2 SD8.2–10.3 retreatments) (no 5% years retreatments) (no 5% years retreatments) CMT (OCT): Sex: 42–59% female Regimen for all groups: in CMT ( $\mu$ m) p Value Diabetes type: not reported case of recurrence or IVT 342 SD124 (–54) NS between HbA1c: not reported persistence of macular groups, <0.01 Baseline VA: ETDRS logMAR oedema, retreatment offered baseline 0.64–0.72 SD0.34–0.36 according to study group, at IVTL 307 SD181 (–116) <0.01 vs Baseline CMT: 385–424 intervals no less than SD91–108 $\mu$ m 4 months L 350 SD169 (–35) Comorbidities: 66–84% phakic Laser ETDRS protocol			,	L		
Age: 64.7–67.2 SD8.2–10.3retreatments) (no5%yearsretreatments)CMT (OCT):Sex: 42–59% femaleRegimen for all groups: inCMT (μm)p ValueDiabetes type: not reportedcase of recurrence orIVT342 SD124 (–54)NS betweenHbA1c: not reportedpersistence of maculargroups, <0.01						
years retreatments) CMT (OCT):  Sex: 42–59% female Regimen for all groups: in CMT (\(\mu\mu\mstrm)\) p Value  Diabetes type: not reported case of recurrence or IVT 342 SD124 (-54) NS between groups, <0.01 between HbA1c: not reported persistence of macular groups, <0.01 between deem and the second secon		•				
Sex: 42–59% female Regimen for all groups: in CMT (µm) p Value Diabetes type: not reported case of recurrence or IVT 342 SD124 (–54) NS between HbA1c: not reported persistence of macular Baseline VA: ETDRS logMAR oedema, retreatment offered 0.64–0.72 SD0.34–0.36 according to study group, at IVTL 307 SD181 (–116) <0.01 vs Baseline CMT: 385–424 intervals no less than SD91–108 µm 4 months L 350 SD169 (–35) Comorbidities: 66–84% phakic Laser ETDRS protocol			, · ·	CMT (OCT):	370	
Diabetes type: not reported HbA1c: not reportedcase of recurrence or persistence of macular oedema, retreatment offeredIVT342 SD124 (-54)NS between groups, <0.01Baseline VA: ETDRS logMAR 0.64-0.72 SD0.34-0.36oedema, retreatment offered according to study group, at intervals no less than SD91-108 μm307 SD181 (-116) 4 months<0.01 vs baselineSD91-108 μm Comorbidities: 66-84% phakic4 months Laser ETDRS protocolL350 SD169 (-35)				CWT (CCT).	CMT (um)	n Value
HbA1c: not reportedpersistence of maculargroups, <0.01Baseline VA: ETDRS logMARoedema, retreatment offeredbaseline0.64–0.72 SD0.34–0.36according to study group, at IVTL307 SD181 (-116)<0.01 vs				IVT		•
Baseline VA: ETDRS logMAR oedema, retreatment offered baseline 0.64–0.72 SD0.34–0.36 according to study group, at IVTL 307 SD181 (–116) <0.01 vs Baseline CMT: 385–424 intervals no less than baseline SD91–108 μm 4 months L 350 SD169 (–35) Comorbidities: 66–84% phakic Laser ETDRS protocol		•			012 02121 ( 01)	
0.64-0.72 SD0.34-0.36 according to study group, at <i>IVTL</i> 307 SD181 (-116) <0.01 vs <i>Baseline CMT</i> : 385-424 intervals no less than baseline  SD91-108 μm 4 months <i>L</i> 350 SD169 (-35) <i>Comorbidities</i> : 66-84% phakic <i>Laser</i> ETDRS protocol		· · · · · · · · · · · · · · · · · · ·	•			•
Baseline CMT: 385–424 intervals no less than baseline SD91–108 μm 4 months L 350 SD169 (–35) Comorbidities: 66–84% phakic Laser ETDRS protocol		——————————————————————————————————————		IVTI	307 SD181 (-116)	
SD91–108 µm 4 months L 350 SD169 (–35)  Comorbidities: 66–84% phakic Laser ETDRS protocol					10. 02.0. ( 1.0)	
Comorbidities: 66–84% phakic Laser ETDRS protocol				L	350 SD169 (-35)	20.0010
		• • • • • • • • • • • • • • • • • • •			(33)	
		eyes	<b>A.</b>			

Study	Participants and baseline values	Intervention	Outcome (change from baseline at study end)		
Ockrim <i>et al</i> /Sivaprasad	N: 88 eyes of 88 patients	Group 1 (IVT, n=43 eyes):	At 12 months		
et al <sup>42 62</sup>	Inclusion criteria: clinically	4 mg IV triamcinolone (mean	BCVA (ETDRS):		
UK	significant DMO persisting	number of IVT injections 1.8		BCVA (letters)	p Value
Design: 2-arm RCT	≥4 months, ≥1 previous laser	(range 1-3))	IVT	-0.2	NS vs L
Follow-up: 1 year	treatment, BCVA 6/12-3/60, VA	Group 2 (L, n=45 eyes):	L	+1.7	
	in fellow eye ≥3/60, duration	ETDRS laser		Plus ≥15 letters	
	visual loss <24 months	photocoagulation (mean	IVT	4.8%	NS vs L
	Exclusion criteria: significant	number of grid laser sessions	L	12.2%	
	macular ischemia, baseline IO	2.1 (range 1–3))	CMT (optical coherence		
	>23 mm Hg, glaucoma,	Regimen for all groups:	tomography):		
	coexistent renal disease, loss of	patients retreated at 4 and		CMT (µm)	p Value
	VA due to other causes, previous		IVT	-91.3	NS vs L
	vitrectomy, intraocular surgery	persistent macular oedema	L	-63.7	
	within 3 months of study entry,	Laser ETDRS protocol			
	previous inclusion in other DR				
	trials, inability to return to				
	follow-up, inability to give informed consent				
	Age: 62.3–64.8 SD7.5–10.1				
	years Sex: 28.9–34.9% female				
	Diabetes type: 97.8–100% type				
	2 DM				
	HbA1c: 7–7.8 IQR6.5–8.7%				
	Baseline VA: ETDRS letter score				
	53.0–54.6 SD13.3–14.2				
	Baseline CMT: 410.4–413.4				
	SD127.8–134.1 µm				
	Comorbidities: 17.8–19.5% PDR,				
	13.3–18.6% pseudophakia,				
	15–17.8% posterior vitreous				
	detachment				

BCVA, best corrected visual acuity; C, control; CMT, central macular thickness; CPL, control plus laser; CSME, clinically significant macular oedema; DDS, dexamethasone; DIL, dexamethasone followed by laser; DM, diabetes mellitus; DMO, diabetic macular oedema; DP, diastolic pressure; DR, diabetic retinopathy; HR QoL, health-related quality of life; IOP, intraocular pressure; IV, intravitreal; IVB, intravitreal bevacizumab; IVP, intravitreal pegaptanib; IVR, intravitreal ranibizumab; IVT, intravitreal triamcinolone; IVTL, intravitreal triamcinolone plus laser; IVVTE, intravitreal VEGF Trap Eye; L, laser; MLT/MPC, macular laser therapy/macular photocoagulation; NEI VFQ-25, National Eye Institute Visual Function Questionnaire-25; NPDR, non-proliferative diabetic retinopathy; NR, not reported; OCT, optical coherence tomography; PDR, proliferative diabetic retinopathy; PRP, panretinal photocoagulation; RCT, randomised controlled trial; RDL, ranibizumab plus deferred laser; RPL, ranibizumab plus laser; SOC, standard of care; SP, systolic pressure; SRFA, fluocinolone; TPL, triamcinoloine plus laser; VA, visual acuity; VEGF, vascular endothelia growth factor.

Table 8 Trials assessing more than one drug

Participants and baseline values	Intervention	Outcome (change from baseline at study end)		
N: 115 eyes of 101 patients Inclusion criteria: eyes with clinically	bevacizumab 1.25 mg	At 24 weeks BCVA (Snellen chart):	BCVA (logMAR)	p Value
macular laser photocoagulation (last	Group 2 (IVB/IVT, n=37		95% CÌ	
Exclusion criteria: visual acuity ≥20/40;	bevacizumab (1.25 mg	IVB	-0.08) (+9 letters	0.01 vs C, NS vs IVB/IVT
6 months; prior intraocular injection or vitrectomy, glaucoma or ocular	triamcinolone (2 mg (0.05 ml)), followed by two	IVB/IVT	-0.21 (-0.30, -0.12) (+10.5	0.006 vs C
characteristics; vitreous hemorrhage; significant media opacity; presence of	alone Group 3 (C, n=37 eyes):	С	-0.03 (-0.08, 0.14) (+1.5 letters	
creatinine ≥3 mg/100 ml; monocular	Regimen for all groups: 3	CMT (OCT):		n Value
Age: 59.7 SD8.3 years (range 39–74)	6-week intervals	IVB	-95.7 <sup>°°</sup>	0.012 vs C, NS vs IVB/IVT
Diabetes type: not reported, 27.6–33.3% on insulin		IVB/IVT	-92.1	0.022 vs C
HbA1c: 9.35–10.06% Baeline VA: not reported Baseline CMT: not reported Comorbidities: (percentage of eyes) 13.9% history of cataract surgery, 81.7% NPDR, 4.3% early PDR, 13.9% regressed PDR; no iris neovascularisation		C	34.9 (7.9, 61.9)	
N: 120 eyes of 120 patients				
400, CMT ≥275 μm  Exclusion criteria: PDR, laser photocoagulation in previous 3 months, no IV corticosteroid or anti-VEGF in previous 3 months  Age: not reported  Sex: not reported  Diabetes type: not reported  HbA1c: not reported  Baseline VA: not reported	bevacizumab  Group 2 (IVT, n=NR eyes): 4 mg (0.1 ml) of IV triamcinolone acetonide Group 3 (IVB/IVT, n=NR eyes): 1.25 mg (0.05 ml) of IV bevacizumab plus 4 mg (0.1 ml) of IV triamcinolone acetonide Regimen for all groups:	<ul> <li>▶ no significant difference be groups (between 1.7 and 2 gained in the different group 2010 report (n=18))</li> <li>CMT (OCT):</li> <li>► CMT reduced in all 3 group (between 17 and 33% reduced different groups in 2010 remarks)</li> </ul>	2.3 lines ups in  os uction in the port (n=18));	
	N: 115 eyes of 101 patients Inclusion criteria: eyes with clinically significant DMO unresponsive to previous macular laser photocoagulation (last session >3 months prior) Exclusion criteria: visual acuity ≥20/40; history of cataract surgery within past 6 months; prior intraocular injection or vitrectomy, glaucoma or ocular hypertension; PDR with high-risk characteristics; vitreous hemorrhage; significant media opacity; presence of traction on the macula; pregnancy; serum creatinine ≥3 mg/100 ml; monocular patients Age: 59.7 SD8.3 years (range 39–74) Sex: 50.5% female Diabetes type: not reported, 27.6–33.3% on insulin HbA1c: 9.35–10.06% Baeline VA: not reported Baseline CMT: not reported Comorbidities: (percentage of eyes) 13.9% history of cataract surgery, 81.7% NPDR, 4.3% early PDR, 13.9% regressed PDR; no iris neovascularisation N: 120 eyes of 120 patients Inclusion criteria: DMO, BCVA 20/40–20/ 400, CMT ≥275 μm Exclusion criteria: PDR, laser photocoagulation in previous 3 months, no IV corticosteroid or anti-VEGF in previous 3 months Age: not reported Diabetes type: not reported HbA1c: not reported Diabetes type: not reported HbA1c: not reported	N: 115 eyes of 101 patients Inclusion criteria: eyes with clinically significant DMO unresponsive to previous macular laser photocoagulation (last session >3 months prior) Exclusion criteria: visual acuity ≥20/40; history of cataract surgery within past 6 months; prior intraocular injection or vitrectomy, glaucoma or ocular hypertension; PDR with high-risk characteristics; vitreous hemorrhage; significant media opacity; presence of traction on the macula; pregnancy; serum creatinine ≥3 mg/100 ml; monocular patients Age: 59.7 SD8.3 years (range 39–74) Sex: 50.5% female Diabetes type: not reported, 27.6–33.3% on insulin HbA1c: 9.35–10.06% Baeline VA: not reported Baseline CMT: not reported Baseline CMT: not reported Comorbidities: (percentage of eyes) 13.9% history of cataract surgery, 81.7% NPDR, 4.3% early PDR, 13.9% regressed PDR; no iris neovascularisation N: 120 eyes of 120 patients Inclusion criteria: DMO, BCVA 20/40–20/ 400, CMT ≥275 μm Exclusion criteria: PDR, laser photocoagulation in previous 3 months Age: not reported Baseline VA: n	Participants and baseline values     Intervention     baseline at study end)       N: 115 eyes of 101 patients Inclusion criteria: eyes with clinically significant DMO unresponsive to previous ession >3 months prior)     Group 1 (IVB, n=41 eyes): bevacizumab 1.25 mg bevacizumab 1.25 mg (0.05 ml)     At 24 weeks       BCVA (Snellen chart): significant DMO unresponsive to previous ession >3 months prior)     Group 2 (IVB/IVT, n=37 eyes): combined triamcinolone (2 mg (0.05 ml)) and triamcinolone (2 mg	Name of the presentation of the protein of

0	5		Outcome (change from		
Study	Participants and baseline values	Intervention	baseline at study end)		
ORCR Network 2010	N: 854 eyes of 691 patients	Group 1 (CPL, n=293	At 1 year		
(Elman <i>et al</i> ) <sup>21 46</sup>	<i>Inclusion criteria</i> : ≥18 years, type 1 or 2	eyes): sham injection plus	BCVA (E-ETDRS Visual		
JSA	DM; study eye: (1) BCVA letter score	prompt (within 3-10 days	Acuity Test):	BCVA (letters)	p Value
Multicenter	78-24 (20/32-20/320), (2) definite retinal	after injection) focal/grid	CPL	+3 SD13	
<i>Design</i> : 4-arm	thickening due to DMO assessed to be	photocoagulation	RPL	+9 SD11	<0.001 vs CPL
lacebo-controlled RCT	main cause of visual loss, (3) retinal	Group 2 (RPL, n=187	RDL	+9 SD12	<0.001 vs CPL
Follow-up: 1-2 years;	thickness measured on time domain OCT	eyes): 0.5 mg IV	TPL	+4 SD13	NS vs CPL
years extension	≥250 µm in central subfield (2 study eyes	ranibizumab plus prompt		BCVA gain categori	es (letters)
Elman) <sup>46</sup> for	per patient could be included if both were	focal/grid photocoagulation	CPL	+10 or more: 28%	
onsenting patients	eligible at study entry)	Group 3 (RDL, n=188		+9 to −9: 59%	
	Exclusion criteria: (1) treatment for DMO	eyes): 0.5 mg IV		-10 or more: 13%	
	within the prior 3 months, (2) panretinal	ranibizumab plus deferred	RPL	+10 or more: 50%	<0.001 vs CPL
	photocoagulation within the prior 4 months	(≥24 weeks) focal/grid		+9 to −9: 45%	
	or anticipated need for panretinal	photocoagulation		-10 or more: 4%	
	photocoagulation within the next	Group 4 (TPL, n=186	RDL	+10 or more: 47%	<0.001 vs CPL
	6 months, (3) major ocular surgery within	eyes): 4 mg IV		+9 to -9: 51%	
	the prior 4 months, (4) history of	triamcinolone plus prompt		-10 or more: 3%	
	open-angle glaucoma or steroid-induced	focal/grid photocoagulation	TPL	+10 or more: 33%	NS vs CPL
	IOP elevation, requiring IOP-lowering	Regimen for all groups:		+9 to -9: 52%	
	treatment, (5) IOP ≥25 mm Hg; systolic	Baseline treatment 0.5 mg		-10 or more: 14%	
	pressure >180 mm Hg, diastolic pressure	IV ranibizumab and 4 mg	Subgroups:		
	>110 mm Hg; myocardial infarction, other	preservative free	▶ BCVA results in TPL group		
	cardiac event requiring hospitalisation,	triamcinolone; study	substantially better for		
	cerebrovascular accident, transient	treatment every 4 weeks	pseudophakic eyes than for		
	ischemic attack, treatment for acute	up to 12 weeks, then	phakic eyes (comparable to		
	congestive heart failure within 4 months	retreatment algorithm: 16	results for RPL and RDL		
	before randomisation	to 20 weeks, monthly	groups) (p not reported)		
	Age: median 62-64 years (25th, 75th	retreatment unless	▶ No difference in results		
	centile 55–58, 69–70)	'success' criteria were met	according to prior treatment		
	Sex: 41–46% female	(visual acuity letter score	for DMO, baseline VA,		
	Diabetes type: 6-9% type 1 DM, 89-92%	≥84 (20/20) or OCT	baseline CMT, baseline		
	type 2 DM, 2–3% uncertain	central subfield thickness	level of retinopathy, focal or		
	HbA1c: median 7.3–7.5% (25th, 75th	<250 μm); 24–48 weeks,	diffuse oedema		
	centile 6.5–6.7, 8.3–8.6)	patients subdivided	CMT (OCT):		
	Baseline VA: letter score 63 SD12	(according to predefined	(201).	CMT (µm)	p Value
	(~20/63 SD2.4 lines)	criteria) into 'success',	CPL	–102 SD151	p . a.a.o
	Baseline CMT: 405 SD134 µm	'improvement', 'no	RPL	-131 SD129	<0.001 vs CPL
	Comorbidities: 60–67% prior treatment for		RDL	-137 SD136	<0.001 vs CPL
	Committee Co Cr /o prior trodument for	provenient or idialo;		.07 02 100	Contin

Ford JA, Lois N, Royle P, et al. BMJ Open 2013;3:e002269. doi:10.1136/bmjopen-2012-002269

hudu.	Portioinanta and becaling values	Intervention	Outcome (change from baseline at study end)		
tudy	Participants and baseline values				
	DMO; 61–68% with NPDR, 26–36% with	'improvement' group	TPL	-127 SD140	<0.001 vs CPL
	PDR or PDR scars	continued treatment, other	Subgroups:		
		groups treated at	▶ pattern of CMT decrease		
		investigator discretion;	similar for groups with CMT		
		alternative treatment	<400 and ≥400 µm at		
		permitted if eye met criteria			
		for 'failure' or 'futility'. In	► Significantly more patients		
		the case of retreatment,	with severe NPDR or worse		
		ranibizumab could be	improved by 2 levels or		
		given as often as every	more in the ranibizumab		
		4 weeks, and	groups (28%, no significant		
		triamcinolone every	change in the other groups)		
		16 weeks (with sham	At 2 years (expanded results,		
		injections as often as	Elman 2011) BCVA (E-ETDRS Visual		
		every 4 weeks). Retreatment for focal/grid			
		laser (after ≥13 weeks	Acuity Test):	BCVA (letters)	p Value
			CPL (n=211)	+3 SD15	p value
		there was oedema	RPL (n=136)	+7 SD13	0.03 vs CPL
		involving or threatening the		+9 SD14	<0.001 vs CP
		center of the macula and if		+2 SD19	NS vs CPL
		complete laser had not	BCVA gain categories (letters)	12 05 10	140 40 01 2
		been given; retreatment	CPL	+10 or more: 36%	
		algorithms facilitated by	0. 2	+9 to -9: 52%	
		web-based real-time data		-10 or more: 13%	
		entry system. Median	RPL	+10 or more: 44%	NS vs CPL
		number of drug injections		+9 to -9: 49%	
		before 1 year visit was 8-9		-10 or more: 7%	
		for ranibizumab, 3 for	RDL	+10 or more: 49%	0.01 vs CPL
		triamcinolone, and 5 sham		+9 to −9: 48%	
		injections. Retreatment		-10 or more: 3%	
		between 1 and 2 years	TPL	+10 or more: 41%	NS vs CPL
		(Elman 2011): median		+9 to −9: 40%	
		injections 2 in RPL group,		-10 or more: 19%	
		3 in RDL group; in TPL	CMT (OCT):		
		group 68% of eyes		CMT (µm)	p Value
		received at least 1	CPL	-138 SD149	
		injection; at least one focal/	RPL	-141 SD155	0.003 vs CPL
		grid laser sessions	RDL	-150 SD143	0.01 vs CPL
		between 1 and 2 years:	TPL	-107 SD145	NS vs CPL
		51% CPL, 40% RPL, 29%			
		RDL, 52% TPL			

Study	Participants and baseline values	Intervention	Outcome (change from baseline at study end)		
<u> </u>	Taranipanio ana baccinio valuec	Laser Modified ETDRS protocol as used in prior DRCR.net protocols	bassime at staay snay		
Jorge <i>et al<sup>51</sup></i> Brazil	N: 63 eyes of 47 patients Inclusion criteria: Refractory	Group 1 (IVB 1.5 mg, n=NR): injections at	At 48 weeks BCVA		
Design: Prospective RCT	cener-involving DMO  Exclusion criteria: NR	baseline and monthly if CSFT (central subfield thickness) measured by		Mean BCVA reduction from	p Value
Follow-up: 24 and 48 weeks (to date, 73% and 56% of patients completed 24 and 48 weeks, respectively)	Age: NR Sex: NR Diabetes type: NR HbA1c: NR Baseline VA: NR Baseline CMT: NR Comorbidities: NR	SDOCT (spectral domain OCT) >275 µm  Group 2 (IVR 0.5 mg, n=NR): injections at baseline and monthly if CSFT >275 µm	IVB1.5	baseline (logMAR) -0.21	vs baseline <0.05 at all-time points vs IVR0.5: no significant difference at all time-points
	Comorbidities. NIT	001 1 >273 μm	IVR0.5	-0.21	vs baseline <0.05 at all time-points vs IVB1.5: no significant difference at all time-points
			CSFT		uno pomo
				Mean CSFT reduction from baseline	p Value
			IVB1.5	−129.6 µm	vs baseline <0.05 at all-time points vs IVR0.5 no significant different at all-time points
			IVR0.5	–137.9 μm	vs baseline <0.05 at all-time points vs IVB1.5 no significant different at all-time points
			At 12 months		•

Study	Participants and baseline values	Intervention	Outcome (change from baseline at study end)		
Study Lim et al <sup>55</sup> Korea Design: 3-arm RCT Follow-up: 12 months	Participants and baseline values  N: 111 eyes of 105 patients Inclusion criteria: eyes with clinically significant DMO based on ETDRS and DMO with central macular thickness of at least 300 μm by optical coherence tomography (OCT)  Exclusion criteria: unstable medical status, including glycemic control and blood pressure; any previous treatment for DMO, including intravitreal, sub-Tenon injection or macular photocoagulation, history of vitreoretinal surgery, uncontrolled glaucoma; proliferative diabetic retinopathy with active neovascularisation, previous panretinal photocoagulation, presence of vitreomacular traction, history of systemic corticosteroids within 6 months, contraindications for bevacizumab or triamcinolone acetonide  Age: 60.4 SD 7.4 (range 48–70) years Sex: 52% female  Diabetes type: NR  HbA1c: 7.2 SD 1.2–7.4 SD1.2  Baseline VA: 0.62 SD 0.23–0.65 SD 0.28 logMAR  Baseline CMT: 447 SD 110–458 SD	Group 1 (IVB/IVT, n=36): IV injection of 1.25 mg (0.05 ml) IVB at 0 and 6 weeks and IV injection of 2 mg (0.05 ml) IVT at 0 weeks. Mean number of addition injection 1.28 Group 2 (IVB, n=38): IV injection of 1.25 mg	IVB/IVT IVB	BCVA (logMAR) -0.15 -0.16 -0.16 CMT (μm) -199 -17s9 -200	p Value 0.088 (between groups)  p Value 0.132 (between groups)
	92 µm  Comorbidities: NR				

Study	Participants and baseline values	Intervention	Outcome (change from baseline at study end)		
Soheilian et al <sup>β7 41 54 141</sup>	N: 150 eyes of 129 patients Inclusion criteria: eyes with clinically	Group 1 (IVB, n=50 eyes): IV injection of	At 36 weeks BCVA (Snellen chart):		
Iran <i>Design</i> : 3-arm RCT	significant DMO (ETDRS criteria)  Exclusion criteria: previous panretinal of	bevacizumab 1.25 mg (0.05 ml) (retreatment IVB		BCVA (logMAR), SD	p Value
Follow-up: 36 weeks (Soheilian 2007 reports	focal laser photocoagulation, prior ocular surgery or injection, history of glaucoma	14 eyes) Group 2 (IVB/IVT, n=50	IVB	-0.28 SD0.25 (+14 SD12.5 letters)	0.053 vs IVB/IVT o
12 week results of the	or ocular hypertension, VA ≥20/40 or <20/ 300, iris neovascularisation, high risk		IVB/IVT	-0.04 SD0.33 (+2	
same trial, these were not considered here)	PDR, significant media opacity,	(1.25 mg (0.05 ml)) and	MPC	SD16.5 letters)	
	monocularity, pregnancy, serum creatinine ≥3 mg/dL, uncontrolled DM	(0.05 ml)), followed by two			
	Age: 61.2 SD6.1 years Sex: 47.3% female	injections of bevacizumab alone (retreatment IVB/IVT		+0.01 SD0.27 (-0.5 SD13.5	
	Diabetes type: not reported  HbA1c: not reported	10 eyes) Group 3 (MPC, n=50		letters) Snellen line	
	Baseline VA: 0.55-0.73 SD0.26-0.28	eyes): focal or modified	11/0	changes	NO
	logMAR  Baseline CMT: 300–359 SD118–149 μm	grid laser (retreatment MPC 3 eyes)	IVB	+2 lines or more: 37%	NS between groups
	Comorbidities: 94% NPDR, 6% early PDR	Regimen for all groups: Retreatments performed at		stable within 2 lines: 59.3%	
		12 week intervals as required		<ul><li>–2 lines or more:</li><li>3.7%</li></ul>	
			IVB/IVT	+2 lines or more: 25%	
				stable within 2	
				lines: 54.2% –2 lines or more:	
			MPC	20.8% +2 lines or more:	
				14.8% stable within 2	
				lines: 66.7%	
				<ul><li>–2 lines or more:</li><li>18.5%</li></ul>	
			CMT (OCT):		
			IVB	<i>CMT (μm), SD</i> –56 SD140	p Value 0.044 vs baseline, NS between
			IVB/IVT	-5 SD113	groups
			MPC	-8 SD67	

Study	Participants and baseline values	Intervention	Outcome (change from baseline at study end)
			Subgroups:    Indicate
			subgroup with ≥400 µm at
			baseline (36 weeks: 1VB -27.2 SD34.8%, IVB/IVT -
			8.8 SD35.9%, MPC -15.1
			SD14.6%, p<0.001 vs
			baseline in IVB and MPC
			groups only)
CVA, best corrected w M, diabetes mellitus; I travitreal bevacizumab ye; L, laser; MLT/MPC R, not reported; OCT, P, systolic pressure; SI	BCVA, best corrected visual acuity; C, control; CMT, central macular thick DM, diabetes mellitus; DMO, diabetic macular oedema; DP, diastolic pres intravitreal bevacizumab; IVP, intravitreal pegaptanib; IVR, intravitreal rani Eye; L, laser; MLT/MPC, macular laser therapy/macular photocoagulation NR, not reported; OCT, optical coherence tomography; PDR, proliferative SP, systolic pressure; SRFA, fluocinolone; VA, visual acuity; VEGF, vasc.	r thickness; CSME, clinically signific c pressure; DR, diabetic retinopathy al ranibizumab; IVT, intravitreal trian llation; NEI VFQ-25, National Eye Ir rative diabetic retinopathy; PRP, pa vascular endothelia growth factor.	BCV4, best corrected visual acuity; C, control; CMT, central macular thickness; CSME, clinically significant macular oedema; DDS, dexamethasone; DIL, dexamethasone followed by laser; DM, diabetes mellitus; DMO, diabetic macular oedema; DP, diastolic pressure; DR, diabetic retinopathy; HR QoL, health-related quality of life; IOP, intraorular pressure; IV, intravitreal; IVB, intravitreal ranibizumab; IVT, intravitreal triamcinolone; IVTL, intravitreal triamcinolone plus laser; IVVTE, intravitreal VEGF Trap Eye; L, laser; MLT/MPC, macular laser therapy/macular photocoagulation; NEI VFQ-25, National Eye Institute Visual Function Questionnaire-25; NPDR, non-proliferative diabetic retinopathy; NR, not reported; OCT, optical coherence tomography; PDR, proliferative diabetic retinopathy; PRP, panretinal photocoagulation; RCT, randomised controlled trial; SOC, standard of care; Systolic pressure; SRFA, fluocinolone; VA, visual acuity; VEGF, vascular endothelia growth factor.

Adverse events are shown in tables 9 and 16. Conjunctival haemorrhages were higher in the ranibizumab arms compared with laser (RESTORE) or no treatment (RESOLVE). In the RESOLVE, RISE and RIDE studies, a considerably higher incidence of intraocular pressure (IOP) increase was reported in the ranibizumab arm compared to control. This increase in IOP was not demonstrated in the RESTORE study. There were no consistent differences in systemic adverse events between ranibizumab and laser or placebo.

#### Bevacizumab

Eight RCTs investigating the use of bevacizumab in DMO were identified (tables 4 and 8). One RCT, the BOLT study (n=80), randomised patients to laser therapy or 1.25 mg intravitreal bevacizumab. The 24 months, the mean changes in BCVA and the proportion of patients who gained 10 ETDRS letters or more was statistically significantly higher in the bevacizumab arm than in the laser arm. Faghihi *et at*  $t^{53}$  (n=80) compared 1.25 mg bevacizumab (average 2.23 injections per patient) with 1.25 mg bevacizumab plus a single laser treatment (average 2.49 injections per patient). After 6 months, the authors found both treatments to be effective at improving BCVA, but neither treatment was found to result in a greater benefit.

Lam et  $at^{55}$  (n=52) compared two doses of bevacizumab (1.25 and 2.5 mg) in patients with diffuse DMO. Patients with focal DMO associated with localised retinal thickening were excluded. At 6 months, following 3 initial monthly injections (no treatment in the remaining 3 months), both groups showed a statistically significantly increased mean BCVA compared with baseline vision, but there was no difference between doses.

Four trials have investigated the combination of bevacizumab and triamcinolone. Ahmadieh *et al*<sup>31</sup> (n=115) compared combined bevacizumab (three 1.25 mg injections at 6 week intervals) plus triamcinolone (2 mg baseline injection only, Triamhexal) with bevacizumab alone (three 1.25 mg at 6 week intervals) and sham injection in patients who had DMO unresponsive (definition not reported) to previous laser (last session more than 3 months previously). The combination arm and bevacizumab alone arm improved mean BCVA more than the sham injection. For BCVA, the combination of bevacizumab plus triamcinolone was non-statistically significantly better than bevacizumab alone.

Soheilian *et al*<sup>67</sup> <sup>41</sup> (n=150) compared combined bevacizumab (1.25 mg) plus triamcinolone (2 mg) with bevacizumab alone and laser alone in patients who were laser naïve. At 36 weeks, bevacizumab alone improved BCVA more than either combination therapy or laser, although the difference was not statistically significant. Extended follow-up at 24 months showed that there was no statistically significant difference between groups for BCVA; however, the direction of effect favours the bevacizumab and combination arms more than the laser.<sup>54</sup>

Downloaded from http://bmjopen.bmj.com/ on June 12, 2017 - Published by group.bmj.com

	READ-2 study <sup>28 47</sup>	RESOLVE study <sup>36</sup>	RESTORE study <sup>24</sup>	RISE study <sup>38</sup>	RIDE study <sup>38</sup>
Retinal haemorrhage	NR	NR	NR	IVR0.3: 12.8%; IVR0.5: 12.7%; C: 20.3%	IVR0.3: 15.2%; IVR0.5: 22.6%; C: 18.9%
Cataract	NR	NR	NR	IVR0.3: 16.8%; IVR0.5: 11.9%; C: 14.6%	IVR0.3: 20%; IVR0.5: 23.4%; C: 23.6%
Vitreous detachment	NR	NR	NR	IVR0.3: 13.6%; IVR0.5: 11.1%; C: 15.4%	IVR0.3: 8.8%; IVR0.5: 12.9%; C: 15%
Ocular hyperemia	NR	NR	NR	IVR0.3: 15.2%; IVR0.5: 11.1%; C: 10.6%	IVR0.3: 3.2%; IVR0.5: 3.2%; C: 7.9%
Vitreous floaters	NR	NR	NR	IVR0.3: 12.8%; IVR0.5: 14.3%; C: 5.7%	IVR0.3: 7.2%; IVR0.5: 8.1%; C: 3.1%
Eye irritation	NR	NR	NR	IVR0.3: 10.4%; IVR0.5: 9.5%; C: 6.5%	IVR0.3: 5.6%; IVR0.5: 5.6%; C: 3.1%
Foreign body sensation in eyes estematic adverse eve	NR	NR	NR	IVR0.3: 12.8%; IVR0.5: 7.1%; C: 4.1%	IVR0.3: 8%; IVR0.5: 2.4%; C: 5.5%
Arterial	Stroke in 1 pt (2%) in	IVR0.3: n=0; IVR0.5:	IVR: n=6 (5%); IVRL:	IVR0.3: 3.2% (n=1 stroke);	IVR0.3: 1.6% (stroke), 5.6% (heart
thromboembolic events	IVRL group- not related to study drug	n=3 (6%); C: n=2 (4%)	n=1 (<1%); L: n=1 (<1%)	IVR0.5: 7.9% (n=5 strokes); C: 7.3% (n=2 strokes)	attack); IVR0.5: 2.4% (stroke), 2.4% (heart attack); C: 1.6% (stroke), 5.6% (heart attack)
Hypertension	NR	IVR0.3: n=4 (8%); IVR0.5: n=5 ((10%); C: n=5 (10%)	IVR: n=9 (8%); IVRL: n=6 (5%); L: n=9 (8%)	Serious IVR0.3: 0.8%; IVR0.5: 3.2%; C: 0.8%	Serious IVR0.3: 1.6%; IVR0.5: 1.6%; C: 0%
Non-ocular haemorrhage	NR	IVR0.3: n=1 (2%); IVR0.5: n=1 (2%); C: n=0	IVR: n=1 (<1%); IVRL: n=0; L: n=1 (<1%)	NR	NR
Proteinuria	NR	NR	IVR: n=1 (<1%); IVRL: n=1 (<1%); L: n=0	NR	NR
Deaths	1 (2%) due to CVA in IVRL group	NR	IVR: n=2 (2%); IVRL: n=2 (2%); L: n=2 (2%)	IVR0.3: 2.4%; IVR0.5: 4%; C: 0.8%	IVR0.3: 3.2%; IVR0.5: 4.8%; C: 1.6%

Downloaded from http://bmjopen.bmj.com/ on June 12, 2017 - Published by group.bmj.com

	BOLT study <sup>23 52</sup>	Lam <i>et al<sup>35</sup></i>	Faghihi <i>et al<sup>53</sup></i>
Number of patients	MLT: n=38; IVB: n=42	IVB1.25, n=26; IVB2.5, n=26	IVB1.25 n=40 IVB 1.25 plus MLT n=40
Ocular adverse events			Not reported
Loss of _15 or _30 ETDRS letters	MLT: n=1 transient, 3 at 24 month analysis;	No significant ocular events (IOP increase, retinal	
	IVB: n=4 transient	tear, retinal detachment, endophthalmitis); no	
Vitreous haemorrhage	MLT: n=1; IVB: n=0	significant difference in change in cataract scores	
Eye pain/irritation/watering during or after injection	MLT:n=0; IVB: n=8	between groups	
Red eye after injection	MLT: n=0; IVB: n=8		
Endophthalmitis	NR		
Transient IOP increase	≥30 mm Hg—MLT: 0; IVB:		
	n=4≥45 mm Hg—MLT: n=1; IVB: n=1		
Floaters after injection	MLT: n=0; IVB: n=2		
Corneal epithelial defect	MLT:n=0; IVB:n=1		
Vitreomacular traction with macular	MLT: n=1; IVB: n=0		
oedema			
Systematic adverse events			
Anaemia	MLT: n=1; IVB: n=0	No systematic adverse effects (1 patient in 1.25 mg	
Vomiting after FFA	MLT: n=1; IVB: n=0	group with foot gangrene requiring amputation due to	
Uncontrolled hypertension	MLT:n=0; IVB: n=1	worsening diabetic neuropathy, considered unrelated	
Polymyalgia rheumatica	MLT:n=0; IVB: n=1	to treatment)	
Intermittent claudication	MLT:n=0; IVB: n=1		
Gastroenteritis	MLT:n=0; IVB: n=1		
Fall	MLT:n=2; IVB: n=0		
Urinary tract infection	MLT:n=0; IVB: n=1		
Chest infection	MLT:n=0; IVB: n=1		
Headaches, dizziness, tiredness	MLT:n=1; IVB: n=0		
Bell palsy	MLT:n=1; IVB: n=0		
Admission for diabetic foot ulcer	MLT:n=1; IVB: n=1		
Admission for cholecystectomy	MLT:n=0; IVB: n=1		
Admission for fall/loss of consciousness	MLT:n=1; IVB: n=0		
Angina—hospital admission	MLT:n=1; IVB: n=0		
Cerebrovascular accident	MLT:n=1; IVB: n=0		
Myocardial infarction	MLT:n=0; IVB: n=2		
Coronary artery bypass graft	MLT:n=0; IVB: n=1		
Dyspnea, chest pain-admitted for hospital observation	MLT:n=0; IVB: n=1		
Death .	NR		

Table 11 Pegaptanib safety		
	Cunningham et al / Adamis et al <sup>39 57</sup>	Sultan et al <sup>40</sup>
Number of patients	IVP0.3, n=44 eyes; IVP1, n=44 eyes; IVP3, n=42 eyes	IVP, n=133 eyes; C, n=127 eyes
Ocular adverse events		
Eye pain	Pegaptanib: 31%; C: 17%	IVP: 11.1%; C: 7%
Vitreous haemorrhage	Pegaptanib: 22%; C: 7%	IVP: 6.3%; C: 7.7%
Punctuate keratitis	Pegaptanib: 18%; C: 17%	IVP: 11.8%; C: 6.3%
Cataract	Pegaptanib: 13%; C: 10%	IVP: 8.3%; C: 9.2%
Eye discharge	Pegaptanib: 11%; C: 10%	NR
Conjunctival haemorrhage	Pegaptanib: 10%; C: 0%	IVP: 22.2%; C: 14.1%
Vitreous opacities	Pegaptanib: 9%; C: 5%	NR
Blurred vision	Pegaptanib: 7%; C: 5%	NR
Other vitreous disorder	Pegaptanib: 7%; C: 0%	NR
Other visual disturbance	Pegaptanib: 7%; C: 0%	NR
Culture-negative endophthalmitis	Pegaptanib: n=1	NR
IOP increase	NR	IVP: 17.4%; C: 6.3%
Retinal haemorrhage	NR	IVP: 6.3%; C: 10.6%
Retinal exudates	NR	IVP: 6.3%; C: 5.6%
Conjunctivitis	NR	IVP: 5.6%; C: 4.2%
Lacrimation increased	NR	IVP: 5.6%; C: 2.8%
Diabetic retinal oedema	NR	IVP: 11.1%; C: 17.6%
Macular oedema	NR	IVP: 9.7%; C: 11.6%
Systemic adverse events		
Non-ocular hypertension	NR	IVP: 13.9%; C: 9.9%
Cardiac disorders	NR	IVP: 6.9%; C: 5.6%
Deaths	NR	IVP: n=4
IOP, intraocular pressure; IVP, intravitrea	l pegaptanib; NR, not reported.	

Lim *et al*<sup>55</sup> (n=111) also evaluated the combination of bevacizumab plus triamcinolone when compared with bevacizumab alone or triamacinolone alone. At 12 months, the authors found no statistically significant difference between groups for BCVA or CMT.

The Efficacy Study of Triamcinolone and Bevacizumab Intravitreal for Treatment of Diabetic Macular Oedema (ATEMD) study, currently only published in abstract form, compared combined therapy with bevacizumab (1.25 mg) and triamcinolone (4 mg) with each of these alone. <sup>56</sup> At 6 months, they found no statistically significant difference between groups. One study comparing bevacizumab with ranibizumab is discussed above. <sup>51</sup> No bevacizumab trials were suitable for meta-analysis because treatment arms were not comparable among included studies.

Adverse events are shown in tables 10 and 16. There was a low frequency of adverse events reported in the included trials. A higher incidence of mild anterior chamber reaction was reported in bevacizumab groups compared with controls. The incidence of IOP increase was comparable between bevacizumab and laser. Soheilian  $et\ at^{87-41}$  were the only authors to report the incidence of lens opacity. No patients in the bevacizumab alone group were found to have lens opacities but in four patients (8%) in the bevacizumab plus triamcinolone group, this finding was observed over the 36-week follow-up period.

# Pegaptanib

Two studies have evaluated pegaptanib in DMO and both compared it with sham injection (table 5). Cunningham *et al*<sup>69</sup>  $^{57}$  compare three doses of pegaptanib (0.3, 1 and 3 mg) and sham injection in laser-naive patients (n=172). At 6 months, patients in the 0.3 and 1 mg groups performed statistically significantly better than those in either the 3 mg or sham groups. Six injections (median) were administered in the 0.3 and 1 mg groups, whereas only five (median) injections were administered in the 3 mg group.

The second trial (n=260), reported by Sultan and colleagues in 2011, compared pegaptanib (0.3 mg) and sham injection. At 2 years, the pegaptanib group showed a statistically significantly greater improvement in mean BCVA compared with sham. 40 However, there was no statistically significant difference in the proportion of patients with an improvement of 10 letters or more. Patients were allowed rescue laser at the assessors' discretion (25.2% of patients in the pegaptanib group and 45% of patients in the sham group received rescue treatment). In regard to meta-analysis, data were only available to combine these trials for the proportion of patients with more than 15 letter gain. Although neither trial individually demonstrated a statistically significant difference favouring pegaptanib over sham (figure 5), when pooled together in meta-analysis, a statistically significant difference was found in favour of pegaptanib (OR 1.94, 95% CI 1.01 to 3.71).

	DA VINCI 2010 <sup>30 58</sup>
Number of patients	IVVTE (all doses) n=175, laser n=44
Ocular adverse events	
Conjunctival hemorrhage	At 6 months: Laser 18.2%, IVVTE 18.9%
	At 12 months: Laser 18.2%, IVVTE 26.9%
IOP increase	At 6 months: Laser 2.3%, IVVTE 9.7%
	At 12 months: Laser 2.3%, IVVTE 9.7%
Eye pain	At 6 months: Laser 4.5%, IVVTE 8.6%
	At 12 months: Laser 4.5%, IVVTE 13.7%
Ocular hyperaemia	At 6 months: Laser 4.5%, IVVTE 6.3%
•	At 12 months: Laser 4.5%, IVVTE 7.4%
Vitreous floaters	At 6 months: Laser 4.5%, IVVTE 5.1%
	At 12 months: Laser 4.5%, IVVTE 6.9%
Endophthalmitis	At 6 months: Laser 0%, IVVTE 1.1%
	At 12 months: Laser 0%, IVVTE 1.1%
Uveitis	At 6 months: Laser 0%, IVVTE 0.6%
	At 12 months: Laser 0%, IVVTE 0.6%
Diabetic retinal oedema	At 6 months: Laser 2.3%, IVVTE 0%
	At 12 months: Laser 2.3%, IVVTE 4.6%
Visual acuity reduced	At 6 months: Laser 2.3%, IVVTE 0%
	At 12 months: Laser 2.3%, IVVTE 0%
Vitreous hemorrhage	At 6 months: Laser 2.3%, IVVTE 0%
	At 12 months: Laser 6.8%, IVVTE 0%
Corneal abrasion	At 6 months: Laser 0%, IVVTE 0.6%
	At 12 months: Laser 0%, IVVTE 4.6%
Retinal tear	At 6 months: Laser 0%, IVVTE 0.6%
	At 12 months: NR
Systematic events	
Hypertension	At 6 months: Laser 6.8%, IVVTE 9.7%
	At 12 months: Laser 0%, IVVTE 1.7%
Myocardial infarction	At 6 months: Laser 0%, IVVTE 1.1%
	At 12 months: Laser 0%, IVVTE 1.7%
Cerebrovascular event	At 6 months: Laser 0%, IVVTE1.1%
	At 12 months: Laser 2.3%, IVVTE 1.7%
Death	At 6 months: Laser 0%, IVVTE 1.7%
	At 12 months: Laser 2.3%, IVVTE 4%

Adverse events for pegaptanib are shown in table 11. There was a higher incidence of eye pain compared to control (31% vs 17%).<sup>39</sup> <sup>57</sup> Cataract formation was similar between the pegaptanib and control groups. There was a higher incidence of IOP increase in the pegaptanib arm compared to control (17.4% vs 6.3%).<sup>40</sup>

#### Other anti-VEGF

Aflibercept has been evaluated in the Da Vinci study (n=219)<sup>30</sup> <sup>58</sup> (table 5). Four regimens of aflibercept (0.5 mg 4 weekly, 2 mg 4 weekly, 2 mg monthly for 3 months, then every 8 weeks, and 2 mg monthly for 3 months followed by treatment as required) were compared with laser. At 6 months, all aflibercept arms had a statistically better BCVA and CMT change than the laser arm. The regimen that resulted in the greatest BCVA gain and CMT reduction was 2 mg every 4 weeks; however, statistical significance between aflibercept arms was not reported. One year extended follow-up showed

that all aflibercept arms were found to have a statistically significantly better BCVA compared to laser.<sup>58</sup>

Adverse events are shown in table 12. There was a higher incidence of IOP increase and eye pain in the aflibercept group compared with laser. Other adverse events were too infrequent to draw meaningful conclusions. The incidence of cataracts was not reported.

#### **Steroids**

# Dexamethasone

Two included trials assessed the use of dexamethasone to treat DMO (table 6): Haller 2010 (full text available) and Callanan (available to date only in an abstract form). Haller 2010 (n=171) compared two doses of dexamethasone, administered as an intravitreal implant (350 and 700  $\mu m$ ) through a 20-gauge transceleral incision, with no treatment. At 90 days only, the 700  $\mu m$  group showed a statistically significantly higher proportion of patients with 10 or more letter gain

	Callanan <i>et al</i> <sup>44</sup>	Haller et al <sup>59</sup>
Number of patients		
Ocular adverse even	ts	
IOP elevation	DIL: 20% (p<0.001); 1%	
	≥10 mm HgL: 1.6% ; 0% ≥10 mm Hg	
Cataract	NR	NR
Anterior chamber	NR	DDS350: 29.1%; DDS700: 26.4%; C: 1.8%
cells		
Anterior chamber	NR	DDS350: 27.3%; DDS700: 20.8%; C: 8.8%
flare		
Vitreous	NR	DDS350: 20%; DDS700: 22.6%; C: 5.3%
haemorrhage		
Eye pain	NR	DDS350: 18.2%; DDS700: 9.4%; C: 3.5%
Vitreous disorder	NR	DDS350: 20%; DDS700: 15.1%; C: 3.5%
Increased IOP	NR	DDS350: 14.5%; DDS700: 9.4%; C: 0%
Conjunctival	NR	DDS350: 14.5%; DDS700: 7.5%; C: 0%
haemorrhage		
Vitreous floaters	NR	DDS350: 7.3%; DDS700: 17%; C: 0%
		No significant differences in: reduced VA, eye irritation,
		abnormal sensation in eye, macular oedema, eye pruritus,
		retinal hemorrhage, DR, nonocular events

compared to no treatment (33% compared with 12%, p=0.007). The 350  $\mu$ m group showed a non-statistically significant improvement compared with laser alone (21% compared with 12%). At 180 days, there was no statistically significant difference between either the dexamethasone group or no treatment group. The treatment effect appeared to peak at 3 months.

The second trial, by Callanan and colleagues (n=253), compared dexamethasone (dose not reported) plus laser with laser alone. Although a greater improvement in mean BCVA was seen at 1–9 months in the dexamethasone plus laser group compared with laser alone, there was no statistically significant difference at 12 months. A mean of 1.6 implants were used over the 12 month period.

These trials were not suitable for meta-analysis since one study is only available in abstract form.

Adverse events are shown in table 13. In the 350 and 700 µm groups compared with no treatment, there was a higher incidence of anterior chamber cells (29.1/26.4% compared with 1.8%), anterior chamber flare (27.3/20.8% compared with 8.8%), vitreous haemorrhage (20/22.6% compared with 5.3%) and increased IOP (14.5/9.4% compared with 0%). However, there was no statistically significant difference in cataract formation between groups at 12 months.<sup>59</sup> Callanan *et al*<sup>44</sup> reported an increase in IOP in the dexamethasone plus laser group compared with laser alone (20% compared with 1.6%).

#### Fluocinolone

Two trials assessed fluocinolone implant for DMO (table 6). The FAME study (n=956) compared two doses of fluocinolone (0.2 and 0.5  $\mu$ g/day) with sham injection in patients with at least one prior laser treatment.<sup>29</sup>

Approximately 25% of patients in each group had more than one prior laser treatment. At 24 months, both doses of fluocinolone showed a statistically significant improvement in mean BCVA compared to sham. There was a modest difference between fluocinolone groups. Rescue laser was given after the first 6 weeks for persistent oedema and was allowed every 3 months. A range of 35–37% of patients in the fluocinolone group and 59% in the sham injection group required rescue laser. Extended follow-up at 36 months showed that both the fluocinolone arms continued to result in a statistically significant benefit compared with sham. <sup>60</sup>

Pearson *et al*<sup>43</sup> (n=196) compared fluocinolone (0.59 mg) with standard of care, either laser or no treatment. At 3 years, there was no statistically significant difference in the proportion of patients with 15 letter gain or more (31% fluocinolone compared with 20% standard of care) between groups and the proportion of patients losing 15 letters or more in the fluocinolone group (17% compared with 14%). Increased incidence of cataracts may have contributed to this difference.

These trials were not suitable for meta-analysis.

Adverse events are shown in table 14. Pearson and colleagues reported a higher incidence of cataracts at 3 years in the fluocinolone group compared with standard of care (55.9% compared with 21.7%). In the extended report of the FAME study, there was a considerably higher incidence of cataract surgery in phakic eyes in the 0.2 and 0.5  $\mu$ g/day fluocinolone groups (80% and 87.2% compared with 27.3%) and increased IOP at any point (37% and 46% compared with 12%).

Following the demonstration in the FAME trial that a lower dose was about as good as higher ones, the higher doses are unlikely to be used.

Table 14 Fluocinolone safety		
	FAME study (Campochiaro <i>et al</i> ) <sup>29 60</sup>	Pearson et al <sup>43</sup>
Number of patients		
Ocular adverse events		
IOP at 12 months	NR	NR
Progression of cataract	NR	NR
Cataract	NR	SRFA: 55.9%;
		SOC: 21.7%
Transient vitreous floaters	NR	NR
Transient subconjunctival	NR	NR
haemorrhage		
Cataract surgery	SRFA0.2: 41.1% (74.9% of those without cataract surgery at baseline,	NR
, , , , , , , , , , , , , , , , , , ,	80% at 36 months); SRFA0.5: 50.9% (84.5% of those without cataract	
	surgery at baseline, 87.2% at 36 months); C: 7% (23.1% of those	
	without cataract surgery at baseline, 27.3% at 36 months)	
Glaucoma	SRFA0.2: 1.6%; SRFA0.5: 2.3%; C: 0.5%	NR
Increased IOP	SRFA0.2: 3.2%; SRFA0.5: 3.3%; C: 0%	SRFA: 69.3%;
increased for	31 II A0.2. 3.2 /0, 31 II A0.3. 3.0 /0, 0. 0 /0	SOC: 11.6%
IOB > 20 mm Ha at any point	SDEAD 2: 10 40/ : SDEAD 5: 22 00/ : C: 4 20/	NR
IOP >30 mm Hg at any point	SRFA0.2: 18.4%; SRFA0.5: 22.9%; C: 4.3%	INU
during 36 months	CDEA0 0: 0 10/ : CDEA0 5: 4 00/ : O: 00/	ND
Trabeculectomy	SRFA0.2: 2.1%; SRFA0.5: 4.8%; C: 0%	NR
Other glaucoma surgery	SRFA0.2: 1.3%; SRFA0.5: 1.3%; C: 0.5%	NR
Trabeculoplasty	SRFA0.2: 0.8%; SRFA0.5: 2.3%; C: 0%	NR
Vitreous haemorrhage	NR	SRFA: 40.2%;
		SOC: 18.8%
Abnormal sensation in eye	NR	SRFA: 37%;
		SOC: 11.6%
Macular oedema	NR	SRFA: 34.6%
Eye pain	NR	SRFA: 26.8%;
		SOC: 15.9%
Eye irritation	NR	SRFA: 22%;
		SOC: 10.1%
Increased lacrimation	NR	SRFA: 22%;
		SOC: 8.7%
Photophobia	NR	SRFA: 21.3%;
		SOC: 21.7%
Blurred vision	NR	SRFA: 21.3%;
		SOC: 15.9%
Vitreous floaters	NR	SRFA: 21.3%;
		SOC: 8.7%
Systemic adverse events		220.0.,70
Serious cardiovascular events	SRFA0.2: 12%; SRFA0.5: 13.2%; C: 10.3%	
Pruritus	NR	SRFA: 38.6%;
Tantao		SOC: 21.7%
Deaths	NR	NR
		TVIII
IOP, intraocular pressure; NR, not rep	ported; SOC, standard of care; SRFA, fluocinolone.	

# Triamcinolone

Ten trials evaluating triamcinolone were identified (tables 7 and 8). All trials evaluated intravitreal administration of triamcinolone, but there were no trials evaluating posterior or anterior subtenon injections. Two trials used Trivaris, <sup>21</sup> <sup>61</sup> two trials used Kenacort, <sup>32</sup> <sup>33</sup> one trial used Kenalog, <sup>62</sup> one trial used Trimahexal <sup>31</sup> and four trials did not report the type of triamcinolone used. <sup>34</sup> <sup>3745</sup> <sup>56</sup> Three doses were assessed in the included studies (1, 4 and 8 mg) and triamcinolone has been combined with laser or bevacizumab.

Ip and colleagues (n=840) were the only authors to evaluate triamcinolone 1 mg (Trivaris).  $^{22}$   $^{61}$   $^{63}$   $^{64}$  They found a statistically significant improvement in mean BCVA at 2 years in the laser group compared with the triamcinolone group and no significant difference between 1 compared with 4 mg.

Several trials compared 4 mg intravitreal triamcinolone. Ip and colleagues (n=840) found that laser therapy resulted in a greater improvement in mean BCVA at 2 years compared to 4 mg triamcinolone (Trivaris).  $^{22}$   $^{61}$   $^{63}$   $^{64}$  Lam *et al*<sup>34</sup> (n=111) found no

	DRCR Network 2008					
	(lp et al/Beck et al/ Bressler et al) <sup>22 61 63 64</sup>	Gillies et al/Sutter et al <sup>32 136–138</sup>	Gillies <i>et al<sup>β3</sup></i>	Kim <i>et al</i> <sup>45</sup>	Lam <i>et al</i> <sup>34</sup>	Ockrim <i>et all</i> Sivaprasad <i>et al</i> <sup>42 62</sup>
umber of patients						
cular adverse events						
	At 2 years (or 3 years when indicated)	At 2 years	-	Not reported	-	At 12 months
IOP ≥30 mm Hg	IVT1: n=22; IVT4: n=53; L: n=3	NR	NR	·	NR	IVT: IOP significantly higher than in L group (18.2 mm Hg, range 12–26 mm Hg); no cases of glaucoma
IOP >22 mm Hg	NR	NR	NR		IVT: 37% (p=0.002 vs L); IVTL: 36% (p=0.002 vs L); L: 5%	NR
IOP ≥10 mm Hg from baseline	IVT1: n=41; IVT4: n=85; L: n=12	NR	NR		NR	NR
IOP ≥5 mm Hg	NR	IVT: 68% (p=0.007 vs C); C: 10%	NR		NR	NR
IOP lowering medication used	IVT1: n=31; IVT4: n=76; L: n=25	IVT: 44% (p=0.0002 vs C); C: 3%	IVTL: 64% (p<0.001); L: 24%		NR	NR
Cataract surgery	IVT1: 23% (of those phakic at baseline, 46% by 3 years (p<0.001 between all groups); IVT4: 51% (of those phakic at baseline, 83% by 3 years); L: 13% (of those phakic at baseline, 31% by 3 years)	IVT: 56% (of phakic eyes over 3 years, p<0.001 vs C); C: 8% (of phakic eyes over 3 years)			NR	NR
Ptosis	NR	NR	NR		NR	NR
Retinal detachment	IVT1: n=2; IVT4: n=4; L: n=2	NR	NR		None	NR
Retinal vein occlusion	IVT1: n=1; IVT4: n=2; L: n=3	NR	NR		NR	NR
Retinal artery occlusion	IVT1: n=0; IVT4: n=0; L: n=1	NR	NR		NR	NR
Anterior ischemic optic neuropathy	IVT1: n=1; IVT4: n=0; L: n=0	NR	NR		NR	NR
Vitrectomy	IVT1: n=26; IVT4: n=19; L: n=31	NR	NR		NR	NR
Open angle glaucoma	IVT1: n=2; IVT4: n=7; L: n=2	NR	NR		NR	NR
Glaucoma filtering surgery	IVT1: n=0; IVT4: n=2; L: n=0	NR	NR		NR	NR
Laser trabeculoplasty	IVT1: n=0; IVT4: n=1; L: n=0	IVT: n=2; C: n=0	IVTL: n=1		NR	NR
	IVT1: n=0; IVT4: n=1; L: n=0	NR	NR		NR	NR

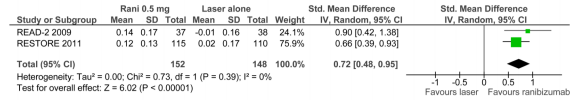
	DRCR Network 2008 (Ip et al/Beck et al/ Bressler et al) <sup>22 61 63 64</sup>	Gillies et al/Sutter et al <sup>32 136–138</sup>	Gillies <i>et al</i> <sup>33</sup>	Kim et al <sup>45</sup>	Lam <i>et ai</i> <sup>34</sup>	Ockrim <i>et all</i> Sivaprasad <i>et al</i> <sup>42 62</sup>
Endophthalmitis	IVT1: n=0; IVT4: n=;0 L: n=0	(Infectious) IVT: n=1; C: NR	(Culture-negative) IVTL: n=1		None	(sterile) IVT: n=1
Pseudoendophthalmitis	IVT1: n=0; IVT4: n=;0 L: n=0	NR	NR		NR	NR
Chemosis	NR	NR	NR		NR	NR
Percentage of increase in cataract scores	NR	NR	NR		IVT:+1.0 SD1.1 (p=NS vs L); IVTL:+1.3 SD1.9 (p=NS vs L); L: +0.5 SD0.9	NR
Ocular hypertension (>21 mm Hg)	NR	NR	NR		NR	NR
Cataract progression	NR	NR	Phakic eyes, progression by ≥2 AREDS grade, IVTL: 64% (p<0.001); L: 11% (p<0.001)		NR	NR
Corneal decompensation	NR	IVT: NR; C: n=1	NR		NR	NR
Cataract surgery	NR	NR	IVTL: 61% (p<0.001); L: 0%		NR	IVT: n=2; L: n=1
Vitreous haemorrhage	NR	NR	NR		IVTL: n=1	
Lens opacity	NR	NR	NR		NR	Significantly greater change in lens opacity IVT group than in L gro (1.9)
Deaths	N=33, unrelated to study treatment	IVT: n=1; C: n=2	IVTL: n=2; L: n=1		NR	NR

	Ahmadieh <sup>31</sup>	ATEMD 2011 (Oliveira Neto <i>et al</i> ) <sup>56</sup>	DRCR Network 2010 (Elman <i>et al</i> , ) <sup>21 46</sup>	Lim et al <sup>55</sup>	Soheilian <i>et al<sup>37 41</sup></i>
Number of patients					
Ocular adverse events Mild anterior chamber reaction	IVB: 19.5% (n=8 eyes), resolved after 1 week of no treatment; IVB/IVT: 18.9% (n=7 eyes), resolved after 1 week of no treatment	NR	NR	NR	IVB: 20% (n=10 eyes), resolved after 1 week; IVB/IVT: 18% (n=9 eyes), resolved after 1 week
Marked anterior chamber reaction	IVB: n=1 (topical corticosteroid and cycloplegic drops)	NR	NR	NR	IVB: n=1 (topical corticosteroids and cycloplegic drops);
Progression of fibrous proliferation	IVB: n=1 with no sign of retinal traction	NR	NR	NR	IVB: n=1 with no sign of retinal traction;
Vitreous haemorrhage	IVB/IVT: n=1 after third injection (excluded from study)	NR	NR	NR	NR
IOP rise	IVB: 23, 22 and 28 mm Hg at 6, 12 and 18 weeks (anti-glaucoma drops)	NR	IOP elevation more frequent with triamcinolone + PL	IVB/ IVT: 8.3% IVT: 10.8%	NR
IOP ≥10 mm Hg from baseline	NR	NR	CPL: n=16; RPL: n=10; RDL: n=5; TPL: n=70	NR	NR
IOP ≥30 mm Hg from baseline	NR	NR	CPL: n=3; RPL: n=2; RDL: n=4; TPL: n=46	NR	NR
Initiation of IOP lowering treatment at any visit	NR	NR	CPL: n=9; RPL: n=5; RDL: n=4; TPL: n=41	NR	NR
Iris neovascularisation	None	NR	NR	NR	NR
Lens opactiy	None	NR	NR	NR	Severe lens opacity IVB/IVT: n=4 eyes; MPC: n=1 eye
Endophthalmitis	NR	NR	CPL: n=1; RPL: n=1; RDL: n=1; TPL: n=0	NR	None
Pseudoendophthalmitis	NR	NR	CPL: n=1; RPL: n=0; RDL: n=0; TPL: n=1	NR	NR
Ocular vascular event	NR	NR	CPL: n=1; RPL: n=1; RDL: n=0; TPL: n=2	NR	NR
Retinal detachment	NR	NR	CPL: n=0; RPL: n=0; RDL: n=1; TPL: n=0	NR	None
Vitrectomy	NR	NR	CPL: n=7; RPL: n=0; RDL: n=3; TPL: n=0	NR	NR
Vitreous haemorrhage	NR	NR	CPL: n=15; RPL: n=3; RDL: n=4; TPL: n=2	NR	None

	Ahmadieh <sup>31</sup>	ATEMD 2011 (Oliveira Neto <i>et al</i> ) <sup>56</sup>	DRCR Network 2010 (Elman <i>et al</i> , ) <sup>21 46</sup>	Lim et al <sup>55</sup>	Soheilian <i>et al<sup>37 41</sup></i>
Cataract surgery	NR	NR	CPL: n=11 (of those phakic at baseline); RPL: n=6 (of those phakic at baseline); RDL: n=8 (of those phakic at baseline); TPL: n=19 (of those phakic at baseline)	NR	NR
Glaucoma surgery	NR	NR	NR	NR	NR
Retinal neovascularisation	NR	NR	NR	NR	IVB: n=4 (all resolved); MPC: n=3 eyes (2 resolved)
Development of early PDR	NR	NR	NR	NR	IVB: n=1; IVB/IVT: n=4; MPC: n=3
Progression to high-risk PDR	NR	NR	NR	NR	IVB: n=4; IVB/IVT: n=3; MP: n=3
Ocular hypertension (≥23 mm HG)	NR	NR	NR	NR	IVB/IVT: 16% (n=8 of eyes), controlled medically in all except 1 that progressed to neovascular glaucoma
Systemic adverse events					The state of the s
Acute myocardial		N=1, considered	No specific systemic adverse events		No significant blood pressure
infarction		not to be related to the study drug	that could be attributed to chance		increase, no thromboembolic events
Deaths	C: n=1	N=1, considered not to be related to the study drug	CPL: n=8; RPL: n=5; RDL: n=3; TPL: n=2		IVB/IVT: n=2; MPC: n=2

C, control; CPL, control plus laser; DMO, diabetic macular oedema; IOP, intraocular pressure; IVB, intravitreal bevacizumab; IVR, intravitreal ranibizumab; IVRL, intravitreal ranibizumab; IV

# 2.1 Mean change in BCVA



#### 2.2 Proportion with >15 letter gain

	Rani 0.5	mg	Laser a	lone		Odds Ratio	Odds Ratio
Study or Subgroup	Events	Total	<b>Events</b>	Total	Weight	M-H, Random, 95% Cl	M-H, Random, 95% CI
READ-2 2009	9	37	0	38	28.0%	25.67 [1.43, 459.42]	
RESTORE 2011	26	115	9	110	72.0%	3.28 [1.46, 7.37]	<del></del>
Total (95% CI)		152		148	100.0%	5.83 [0.90, 37.86]	
Total events	35		9				
Heterogeneity: Tau <sup>2</sup> = Test for overall effect				= 0.16);	I <sup>2</sup> = 48%		0.001 0.1 1 10 1000
. cct .c. Svoran oncot		5.00	,				Favours laser Favours ranibizuma

#### 2.3 CMT

	Rani 0.5 mg			Laser alone			Std. Mean Difference		Std. Mean Difference
Study or Subgroup	Mean	SD	Total	Mean	SD	Total	Weight	IV, Random, 95% CI	IV, Random, 95% CI
READ-2 2009	-103.73	126.76	37	-144.76	109.22	38	47.2%	0.34 [-0.11, 0.80]	<b>+■</b> -
RESTORE 2011	-118.7	115.07	115	-61.3	132.29	110	52.8%	-0.46 [-0.73, -0.20]	-
Total (95% CI)			152			148	100.0%	-0.08 [-0.87, 0.71]	•
Heterogeneity: Tau <sup>2</sup> = 0.29; Chi <sup>2</sup> = 8.96, df = 1 (P = 0.003); $I^2$ = 89% Test for overall effect: Z = 0.20 (P = 0.84)									-2 -1 0 1 2 Favours laser Favours ranibizumab

Figure 2 Ranibizumab 0.5 mg alone versus laser alone. (A) Mean change in best corrected visual acuity. (B) Proportion with >15 letter gain. (C) central macular thickness.

statistically significant difference between laser and triamcinolone at 6 months (triamcinolone type not reported). When these two trials were pooled through meta-analysis, the treatment effect favoured laser but the differences were not statistically significant (figure 6). Ockrim *et al*<sup>62</sup> (n=88) compared 4 mg intravitreal triamcinolone (Kenalog) with laser alone. At 12 months, they

found no statistically significant BCVA improvement between the triamcinolone and laser groups. Gillies *et al*<sup>32</sup> (n=69) compared 4 mg of triamcinolone (Kenacort) with sham injection. Mean BCVA improved statistically significantly with triamcinolone at 24 months compared with sham injection (3.1 letter gain compared with 2.9 letter loss, p=0.01).

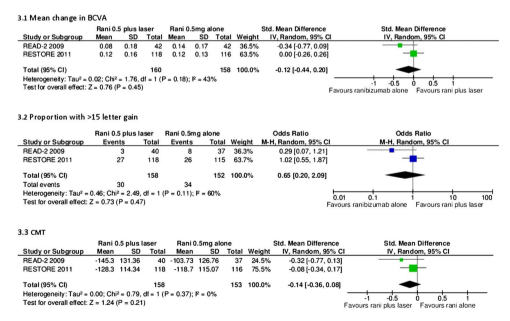


Figure 3 Ranibizumab 0.5 mg plus laser versus ranibizumab 0.5 mg alone. (A) Mean change in best corrected visual acuity. (B) Proportion with >15 letter gain. (C) central macular thickness.

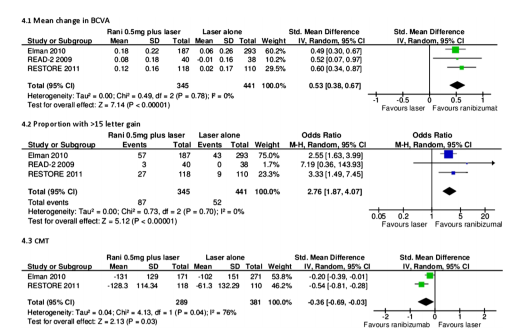


Figure 4 Ranibizumab 0.5 mg plus laser versus laser alone. (A) Mean change in best corrected visual acuity. (B) Proportion with >15 letter gain. (C) Central macular thickness.

Lam et  $al^{64}$  (n=111) compared triamcinolone 4 mg alone with 4 mg of triamcinolone plus laser or laser alone. At 6 months, the authors found no difference in BCVA between any of the groups. Elman *et al*<sup>21</sup> (n=854) compared 4 mg of triamcinolone (Trivaris) plus laser with ranibizumab plus prompt (within 3-10 days) or deferred (more than 24 week) laser and laser alone. At 2 years, they found a statistically significant difference in mean BCVA between ranibizumab plus prompt/ deferred laser compared with laser alone (7 letter gain/ 9 letter gain compared with 3 letter gain), but no difference with triamcinolone plus laser compared with laser alone (2 letter gain compared with 3 letter gain). Neto et al<sup>56</sup> (n=120) compared 4 mg triamcinolone alone (triamcinolone type not reported) with 4 plus 1.25 mg bevacizumab. At 6 months, they found no statistically significant difference between groups.

The Elman and Lam studies were suitable for meta-analysis, which showed non-statistically significant improvements in mean BCVA and the proportions of patients with more or equal than 15 letter gain in the triamcinolone plus laser group compared with laser alone (figure 7).

Adverse events are shown in tables 15 and 16. Triamcinolone was associated with consistently higher incidences of IOP increase and cataracts. Gilles and colleagues reported a cataract rate of over 50% by 3 years in patients treated with triamcinolone.

#### Other pertinent studies

Only one study in abstract form directly compared bevacizumab with ranibizumab.<sup>51</sup> Bevacizumab and ranibizumab have been compared through an indirect comparison of five trials.<sup>65</sup> There was no evidence of a difference between the drugs; however, wide credible intervals meant that the superiority of either drug could not be excluded.

Two-year results of the CATT (Comparison of AMD Treatment Trials) and 1 year results of the IVAN (Inhibit VEGF in Age-related choroidal Neovascularisation), recently published, have demonstrated a good safety profile of anti-VEGF therapies when used to treat patients with age-related macular degeneration. <sup>66</sup> <sup>67</sup> The CATT study randomised 1208 patients with AMD to monthly or as required injection of either ranibizumab or bevacizumab. At 1 year, the mean BCVA was similar in

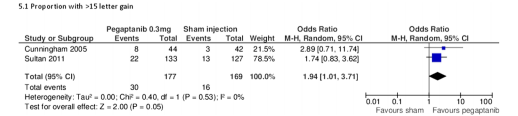


Figure 5 Pegaptanib 0.3 mg versus sham injection. (A) Proportion with >15 letter gain.

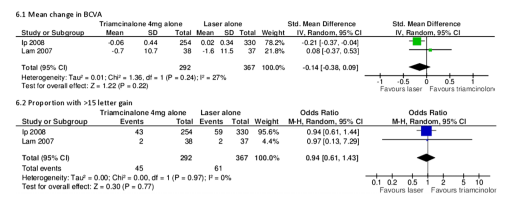


Figure 6 Triamcinolone 4 mg versus laser alone. (A) Mean change in best corrected visual acuity. (B) Proportion with >15 letter gain.

both groups (8 letter gain in bevacizumab and 8.5 in ranibizumab). Over 2 years, the rates of deaths, myocardial infarction and stroke did not differ between the ranibizumab and bevacizumab treatment groups. However, there was a higher rate of serious adverse events in the bevacizumab group compared with the ranibizumab group. This increased event rate was driven mainly by hospitalisations (RR 1.29, 95% CI 1.01 to 1.66). However, the hospitalisations were not caused by known adverse events of bevacizumab. Arteriothrombotic events and heart failure occurred in less than 2% of participants in the IVAN, and they were more often observed in the ranibizumab group than in the bevacizumab group (p=0.03). Further data from other ongoing clinical trials may provide more insight on the safety or anti-VEGF treatment and possible differences on this respect among available drugs.

Campbell et  $al^{68}$  conducted a population-based nested case—control study of 91 378 older adults with a history of physician-diagnosed retinal disease. The authors found that neither ranibizumab nor bevacizumab was associated with significant risks of ischaemic stroke, acute myocardial infarction, congestive heart failure or venous thromboembolism.

A recent systematic review specifically assessing adverse events in anti-VEGF drugs found a low incidence of serious (below 1 in 100) and non-serious ocular events (below 1 in 500) from ranibizumab, bevacizumab and pegaptanib.  $^{69}$ 

Fung et  $a\bar{l}^{70}$  used an internet-based survey of clinicians to assess the safety of bevacizumab. The survey covered over 5000 patients and found that bevacizumab was associated with an infrequent incidence of adverse events (all less than 0.21%).

One study, which assessed diclofenac, did not meet the inclusion criteria (follow-up for only 12 weeks).<sup>71</sup> The authors randomised 32 patients to either intravitreal diclofenac or triamcinolone and found that both diclofenac and triamcinolone reduced CMT, but a statistically significant visual improvement was observed only in the triamcinolone group.

Sfikakis *et al*<sup>72</sup> undertook a 30-week randomised crossover trial comparing infliximab and placebo. The study failed to meet our inclusion criteria (only 11 patients included). The authors found that infliximab resulted in a 28.6% improvement in vision compared with 4.3% with placebo. The improvement seen with placebo could be due to a 'carry over effect', seen in cross-over trials.

The Fenofibrate Intervention and Event Lowering in Diabetes (FIELD) trial was primarily a study to see if the lipid-lowering agent fenofibrate could reduce macrovascular and microvascular events in type 2 diabetes.<sup>73</sup>

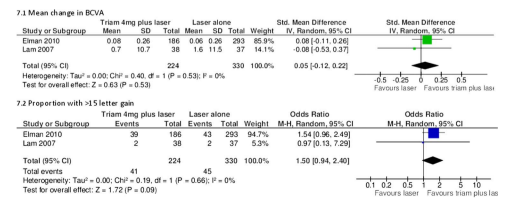


Figure 7 Triamcinolone 4 mg plus laser versus laser alone. (A) Mean change in best corrected visual acuity. (B) Proportion with >15 letter gain.

However, a substudy within FIELD recruited 1012 patients to a retinopathy study. The primary outcome in the main study was need for laser therapy (3.4% on fenofibrate vs 4.9% on placebo), but the substudy used retinal photography to assess progression of retinopathy or development of macular oedema. The HR at 6 years for DMO was 0.69 (95% CI 0.54 to 0.87) in the fenofibrate group compared to placebo.

Ruboxistaurin is another oral agent which has been assessed for the treatment of DMO. Aiello and colleagues randomised 686 patients to receive placebo or one of three doses of ruboxistaurin. There was no statistically significant difference in delay to sight-threatening DMO in any ruboxistaurin group compared to placebo. The authors suggest that differences in laser treatment between groups may have contributed to the non-significant finding.

# Assessment of heterogeneity within meta-analysis

Heterogeneity was assessed methodologically and statistically. Methodological heterogeneity was assessed by comparing the study population, interventions, outcome measures and follow-up. Studies that were not methodologically comparable were excluded from the meta-analysis. For example, bevacizumab trials were not pooled because Soheilian  $\operatorname{et} \operatorname{at}^{37}$  included patients who were laser naïve and Ahmadieh  $\operatorname{et} \operatorname{at}^{31}$  included patients who were unresponsive to laser. Some analyses were also excluded because sufficient details were not reported in the studies. For example, several studies failed to report SDs.  $^{35\ 39}$ 

Statistical heterogeneity was assessed through I<sup>2</sup> scores. High statistical heterogeneity was found in two analyses (2.3 and 4.3). Therefore, these results should be interpreted with due caution. Moderate heterogeneity was found in three analyses (2.2, 3.1 and 3.2). Low heterogeneity was found in the remaining eight analyses.

# **Ongoing trials**

There are numerous ongoing studies listed in appendix 2. The most salient studies include a study to compare ranibizumab and bevacizumab (Schmidt-Erfurth), a study investigating rescue ranibizumab treatment for patients who have failed on bevacizumab (Chaudhry), a study evaluating two algorithms for ranibizumab, 'treat and extend' and 'as required' (RETAIN), further studies of Trap-eye (VIVID and VISTA) and trials which are examining the use of NSAIDs, such as diclofenac and nepafenac (NEVANAC and Soheilian).

# **DISCUSSION**

It appears that anti-VEGF treatment is effective in DMO, especially ranibizumab and bevacizumab. Meta-analysis of available short-term data (up to 2 years) suggests that ranibizumab is superior to laser and that adding laser to ranibizumab treatment does not confer additional benefit. Steroid treatment has demonstrated mixed success and, almost uniformly, increased the incidence

of cataracts and IOP. The licence for fluocinolone takes note of this and it is positioned as a treatment when others have failed.

# Strengths and limitations of the review

There are a number of strengths of this review. A robust systematic review methodology was used. Reliability was improved by excluding trials with small sample sizes or short follow-up. Since a number of trials included similar intervention arms, consistent treatment effects further improve reliability. Validity was improved by assessing the quality of trials using the Cochrane risk of bias tables. Including abstracts from ARVO provided up-to-date results. Pooling results through meta-analysis provided further evidence. The random effects model was used throughout to allow for heterogeneity among studies.

This review, however, has limitations. Although the inclusion of abstracts provides more up-to-date results, the studies contained in these abstracts could not be assessed for risk of bias and should therefore be interpreted with caution. In addition, reporting of quality assessment criteria was variable. Allocation concealment was especially poorly reported. There was only one study which compared different anti-VEGFs<sup>51</sup> and none that compared steroids (fluocinolone vs dexamethasone vs triamcinolone). Therefore, it is difficult to assess the effectiveness within drug classes. As with any meta-analysis, questions of heterogeneity arise. Follow-up periods varied among studies. A difference of 6 months was allowed for studies to be pooled for meta-analysis, but this could have still resulted in heterogeneity. High statistical heterogeneity was found in a quarter of the analyses. Furthermore, because of the low number of trials included, publication bias could not be assessed by funnel plot analysis. The manufacturers funded most of the trials for ranibizumab, pegaptanib, dexamethasone and fluocinolone, whereas trials for bevacizumab and triamcinolone were generally funded by non-pharmaceutical organisations. Generally, the noncommercial studies had smaller numbers, perhaps because of the funding restraints.

It is important to note that there may be differences in laser treatment protocol between studies. This applies to trials which combine drug treatments with laser or include laser as a comparator. All studies referred to the ETDRS protocol<sup>19 20</sup> or a modified version of it. In the ETDRS, once a diagnosis of clinically significant macular oedema was made, an angiogram was obtained to identified 'treatable lesions'. 'Treatable lesions' included discrete points of retinal hyperfluorescence or leakage (most of these are often microaneurysms), areas of diffuse leakage within the retina related to microaneurysms, intraretinal microvascular abnormalities, diffusely leaking retinal capillary bed and retinal avascular zones. In the ETDRS protocol, treatment of lesions closer than 500 microns from the centre of the macula was not required initially; however, if vision was less than 20/40 and the oedema and leakage persisted, treatment up to 300 microns from the centre of the macula was

recommended unless there was capillary dropout; in the latter case, treatment was not recommended as it may lead to further loss of perifoveal capillaries.

However, in routine clinical practice, clinicians generally use lighter and less intense treatment than specified in the ETDRS protocol. <sup>76</sup> In addition, some centres do not use fluorescein angiography (unlike the ETDRS study<sup>19</sup>) to guide treatment. The exact adherence to the ETDRS protocol within studies is unclear. For example, in the BOLT study, a modified ETDRS protocol was used. One of the aims of the protocol was 'not darkening/whitening of microaneurysms', which is not consistent with the ETDRS protocol.

# Interpretation of the results

The anti-VEGF drugs appear to be clinically effective in treating DMO in short-term studies (up to 2 years). Ranibizumab has the most robust evidence base and has shown superiority compared to laser and sham injection in all trials and meta-analyses, except for the proportion of patients with 10 or more letter gain in the DRCR.net study published by Elman *et al*<sup>46</sup> at 2 years follow-up. Adding laser to ranibizumab conferred no benefit. Bevacizumab has also been shown to be superior to laser. Three doses have been used (1.25, 1.5 and 2.5). The higher dose does not appear to add further benefit, and most studies in the literature use 1.25 mg. The addition of triamcinolone to bevacizumab did not provide further benefits. Pegaptanib has only been compared to sham injection. Mean change in BCVA favoured pegaptanib, but only through meta-analysis did the proportion of patients with more than 15 letter gain favour pegaptanib. Further published data are required before drawing conclusions on aflibercept. However, although the anti-VEGF drugs are a significant advance, they fail to improve BCVA by 10 or more letters in half or more patients, and so they do not provide a complete answer to DMO.

Steroid treatments have inconsistent results and are undoubtedly associated with increased IOP and cataract. The effects of dexamethasone appear to peak at 3 months. At 6 months, there was no significant difference compared with laser. This might imply that earlier retreatment is needed if the beneficial effect is to be maintained, but increasing the number of treatments would very likely increase the associated complications, especially with the relatively large needle size. The addition of laser did not appear to add further benefit. There was no significant difference in cataract formation at 6 months with dexamethasone compared to observation, but it is likely that a higher incidence of cataracts would be seen with longer follow-up. Significantly more patients suffered increased IOP in the dexamethasone group compared with observation. Fluocinolone has been shown to be effective compared with sham injection (FAME);<sup>29 60</sup> however, when compared to standard of care (laser or observation at clinician's discretion), there was no significant difference in the proportion of patients with a 15 letter or more gain. Both studies reported higher incidence of cataract formation in the fluocinolone group, with over 80% at 3 years at the higher dose. Results for triamcinolone are inconsistent. Ip *et al*<sup>61</sup> found that laser was more effective, while others have found no statistically significant difference. Triamcinolone combined with laser, however, seemed to have similar efficacy as ranibizumab combined with laser in pseudophakic eyes. <sup>21 46</sup> Triamcinolone is more effective than sham injection. Triamcinolone has consistently been associated with increased incidence of cataract and raised IOP.

Steroids and laser therapy may affect CMT in a different manner from anti-VEGF drugs. For example, when ranibizumab alone is compared with ranibizumab plus laser, it appears to be more effective in terms of mean change in BCVA and proportion of patients with more than 15 letter gain. However, ranibizumab plus laser is more effective at reducing CMT. Furthermore, when triamcinolone plus laser is compared with ranibizumab plus laser, the latter appears to be more effective in terms of change in BCVA and proportion of patients with more than 15 letter gain, but triamcinolone plus laser is more effective at reducing CMT. The reasons for this are unclear. There is a weak correlation between CMT and BCVA. However, the long-term benefits of reducing CMT are currently unknown.

No large observational studies were identified that compared anti-VEGF drugs. Using an internet-based survey, Fung et  $al^{70}$  found the incidence of adverse events in bevacizumab to be low. One small outbreak of sterile endophthalmitis was reported with a single batch of bevacizumab in Canada, emphasising the need for sterility when preparing aliquots.  $^{77}$  Curtis et al $^{78}$  carried out a very large retrospective cohort study in 146 942 patients aged 65 and over with age-related macular degeneration (AMD). Their aim was to examine cardiovascular outcomes in patients treated with the four options: photodynamic therapy (PDT), pegaptanib, bevacizumab and ranibizumab. The authors reported that one of their comparisons showed an increase in overall mortality and stroke risk with bevacizumab compared to ranibizumab, with HRs of 0.86 (95% CI 0.75 to 0.98) and 0.78 (0.64 to 0.96), respectively. However, owing to the very large cost differences between bevacizumab and ranibizumab, the authors noted that selection bias might be operating, with poorer people (with poorer health) more likely to be treated with bevacizumab. They therefore carried out another analysis using only ophthalmological clinics which used only one drug, to avoid selection bias. This analysis showed no significant difference: overall mortality HR for ranibizumab 1.10 (95% CI 0.85 to 1.141); MI 0.87 (0.53 to 1.14); stroke 0.87 (0.61 to 1.24).

Gower *et al*<sup>79</sup> analysed 77 886 anti-VEGF injections from Medicare data (46% ranibizumab and 54% bevacizumab). Results have only been published in abstract form. The authors found an increased risk of overall mortality and cerebrovascular events in the bevacizumab

group (HR 1.11 99% CI 1.01 to 1.23 and 1.57, 1.04 to 2.37, respectively). There was no statistically significantly increased risk in the ranibizumab group. The authors acknowledge that a limitation of the study is a failure to adjust for important confounding factors (such as smoking, hypertension and hyperlipidaemia). Considering the cost difference, it is likely that patients treated with bevacizumab would have been in a lower socioeconomic class and therefore at high risk of mortality and vascular disease.

# Implications for clinicians

The anti-VEGF drugs appear to be a significant advance in the treatment of DMO and are regarded now as the treatment of choice for patients affected by this condition. Studies assessing the effectiveness of steroids have reported mixed results. The high rates of cataract and increased IOP are a drawback. Triamcinolone combined with laser may be a good option for pseudophakic patients and may be more cost-effective than treatment with ranibizumab. However, the need for fewer administrations, potentially one every 3 years with fluorinolone, is advantageous. From an administration perspective, some patients might prefer infrequent steroid injections with a sizeable risk of cataract, and a small, but existent, risk of glaucoma, to frequent anti-VEGF injections, even if the potential gain may not be fully comparable. Steroids may also be considered for patients who do not adequately respond to anti-VEGFs. Currently, the role of laser in the treatment of DMO is debatable. Short-term data from available trials have demonstrated the superiority of anti-VEGF with regard to laser treatment but have failed to demonstrate a benefit of combining both treatment approaches. It is possible that some ophthalmologists may still opt to offer laser treatment to patients with very focal areas of leakage.

Currently, there is more evidence for the effectiveness of ranibizumab and bevacizumab than for pegaptanib and VEGF-trap eye. The results of direct head to head trials of ranibizumab and bevacizumab are awaited. Bevacizumab is not licensed for intraocular use but costs considerably less than other forms of therapy. Ranibizumab is licensed and more expensive, but its use is supported by large manufacturer-funded trials demonstrating its clinical effectiveness. In the UK, the General Medical Council recommends that unlicensed medications should only be prescribed if 'an alternative, licensed medicine would not meet the patient's needs' and there is 'a sufficient evidence base and/or experience of using the medication to demonstrate its safety and efficacy. 80 The FDA says that when using a drug 'off-label', clinicians 'have the responsibility to be well informed about the product, to base its use on firm scientific rationale and on sounded medical evidence, and to maintain records of the product's use and effects'.81 Patients should be fully aware of the use of any unlicensed medication and consent to any safety or efficacy uncertainties.

The place of intravitreal steroids needs consideration now that we have the anti-VEGFs drugs, as does the role of laser. The anti-VEGFs drugs may now be the first-line treatment in place of laser, with laser being used selectively for focal lesions, and in sequence after anti-VEGF therapy once the retinal thickness has been reduced. However, it should be noted that about half of the patients do not get good results with anti-VEGFs. In RESTORE, only 50% of patients had gains in VA of 10 or more letters. So the anti-VEGFs are 'game-changers', but their impact should not be overestimated.

In those who do not respond to anti-VEGFs or laser, there remains a place for steroids, despite their high adverse effect rates. The European licence for fluocinolone recognises this, by stating that it should be used when other therapies have not had sufficient effect. The commonest adverse effect is cataract, but that is very common in people with diabetes, and many are already pseudophakic when treatment of DMO is required.

Vitreoretinal surgery for the treatment of DMO was not included in our review. Laidlaw reviewed the literature and only found evidence for vitrectomy when there were signs of clinical or OCT traction. 83 However, even in these cases, the evidence was not strong.

# Implications for policy makers

In the UK, the National Institute of Health and Clinical Excellence (NICE) has recently made the decision not to recommend ranibizumab for the treatment of DMO.<sup>84</sup> NICE concluded that ranibizumab, although clinically effective, was not cost-effective compared to laser therapy. Bevacizumab is less than a tenth of the cost of ranibizumab but is unlikely to be licensed. This beckons the question as to whether policy makers should recommend cheaper unlicensed medications over a more expensive licensed alternative when their efficacy and side effects appear to be similar.

# **Unanswered questions**

Several unanswered questions remain. Studies evaluating the effectiveness of ranibizumab compared with bevacizumab are needed. Although the anti-VEGFs are clinically effective and a major step forward in the management of DMO, it has to be noted that they have little effect in a large number of patients. Generally speaking, the proportion of patients who have demonstrated 10 or more letter gain using anti-VEGFs is between 30% to 50% in the trials that demonstrate the greatest effectiveness. Most of these patients would not achieve the 20/40 visual acuity required for driving. More effective treatments, or combinations of treatments, are required.

There is a lack of specific evidence for the use of anti-VEGF drugs or steroids in patients with macular ischaemia secondary to DMO. A number of trials excluded patients with macular ischaemia. <sup>23</sup> <sup>34</sup> <sup>35</sup> <sup>40</sup> <sup>53</sup> <sup>62</sup> The RESTORE trial included patients with macular

ischaemia and undertook a subgroup analysis.<sup>24</sup> The authors compared patients with (n=34) and without (n=35) macular ischaemia at baseline. They found that those without macular ischaemia responded better to ranibizumab (mean average change in BCVA at 12 months 7.2 letters gain compared with 6.3 letters). Larger trials are needed to assess the use of anti-VEGF drugs and steroids in patients with macular ischaemia.

The duration of treatment is as yet uncertain. Most of the included studies use a retreatment protocol based on clinical need or OCT results. For example, in the BOLT study, patients received a median of nine injections of bevacizumab over 24 months. <sup>23</sup> <sup>85</sup> However, it is not yet known for how frequent long-term maintenance injections will be needed and whether laser treatment in sequence could potentially reduce the number of anti-VEGF injections required. Other treatment strategies to apply laser, such as using laser power at subthreshold levels, may prove more effective. <sup>86</sup> Future trials should use active comparators which are used in routine clinical practice and avoid placebo-controlled trials.

#### CONCLUSION

This review evaluated current treatments for DMO. Undoubtedly, the use of anti-VEGFs heralds a new era for patients who suffer from DMO. Currently, the anti-VEGFs ranibizumab and bevacizumab have consistently shown good clinical effectiveness without major unwanted side effects. Steroid results have been mixed and are usually associated with cataract formation and IOP increase. Based on the short-term data available, adding laser therapy to anti-VEGFs does not appear to confer additional benefit.

Despite the current wider spectrum of treatments for DMO, only a small proportion of patients recover good vision (≥20/40), and thus the search for new therapies to prevent and manage DMO needs to be continued.

Contributors JAF screened titles, checked data extraction, performed the meta-analysis and drafted the manuscript. NL conceived the idea, interpreted the results and provided clinical expertise throughout. PR performed the literature search, updated the searches, screened the titles and managed the references. CC extracted data from the studies. DS screened the titles and checked the meta-analysis. NW designed the review and supervised the running of the study. All authors contributed to the final draft.

# Competing interests None.

Provenance and peer review Not commissioned; externally peer reviewed.

**Disclosure** The authors report no proprietary or commercial interest in any product mentioned or concept discussed in this article.

Data sharing statement No additional data are available.

**Protocol** This review was built upon several technology appraisals for NICE, and therefore no protocol exists.

#### **REFERENCES**

 Holman N, Forouhi NG, Goyder E, et al. The Association of Public Health Observatories (APHO) Diabetes Prevalence Model: estimates of total diabetes prevalence for England, 2010–2030. Diabet Med 2011;28:575–82.

- Williams R, Airey M, Baxter H, et al. Epidemiology of diabetic retinopathy and macular oedema: a systematic review. Eye (London) 2004;18:963–83.
- Henricsson M, Sellman A, Tyrberg M, et al. Progression to proliferative retinopathy and macular oedema requiring treatment. Assessment of the alternative classification of the Wisconsin Study. Acta Ophthalmol Scand 1999;77:218–23.
- Chen É, Looman M, Laouri M, et al. Burden of illness of diabetic macular edema: literature review. Curr Med Res Opin 2010;26:1587–97.
- Hirai FE, Knudtson MD, Klein BE, et al. Clinically significant macular edema and survival in type 1 and type 2 diabetes. Am J Ophthalmol 2008;145:700–6.
- Klein R, Klein BE, Moss SE, et al. The Wisconsin Epidemiologic Study of Diabetic Retinopathy: XVII. The 14-year incidence and progression of diabetic retinopathy and associated risk factors in type 1 diabetes. Ophthalmology 1998;105:1801–15.
- Klein R, Knudtson MD, Lee KE, et al. The Wisconsin Epidemiologic Study of Diabetic Retinopathy XXIII: the twenty-five-year incidence of macular edema in persons with type 1 diabetes. Ophthalmology 2009:116:497–503.
- Knudsen LL, Lervang HH, Lundbye-Christensen S, et al. The North Jutland County Diabetic Retinopathy Study (NCDRS)
   Non-ophthalmic parameters and clinically significant macular oedema. Br J Ophthalmol 2007;91:1593–5.
- Thomas RL, Dunstan F, Luzio SD, et al. Incidence of diabetic retinopathy in people with type 2 diabetes mellitus attending the Diabetic Retinopathy Screening Service for Wales: retrospective analysis. BMJ 2012;344:e874.
- Wong TY, Mwamburi M, Klein R, et al. Rates of progression in diabetic retinopathy during different time periods: a systematic review and meta-analysis. *Diabetes Care* 2009;32:2307–13.
- Huang ES, Brown SE, Ewigman BG, et al. Patient perceptions of quality of life with diabetes-related complications and treatments. Diabetes Care 2007;30:2478–83.
- Shea AM, Curtis LH, Hammill BG, et al. Resource use and costs associated with diabetic macular edema in elderly persons. Arch Ophthalmol 2008;126:1748–54.
- Minassian DC, Owens DR, Reidy A. Prevalence of diabetic macular oedema and related health and social care resource use in England. Br J Ophthalmol 2012;96:345–9.
- Happich M, Reitberger U, Breitscheidel L, et al. The economic burden of diabetic retinopathy in Germany in 2002. Graefes Arch Clin Exp Ophthalmol 2008;246:151–9.
- Bhagat N, Grigorian RA, Tutela A, et al. Diabetic macular edema: pathogenesis and treatment. Surv Ophthalmol 2009;54:1–32.
- Murata T, Ishibashi T, Khalil A, et al. Vascular endothelial growth factor plays a role in hyperpermeability of diabetic retinal vessels. Ophthalmic Res 1995;27:48–52.
- Barile GR, Pachydaki SI, Tari SR, et al. The RAGE axis in early diabetic retinopathy. Invest Ophthalmol Vis Sci 2005;46:2916–24.
- The Diabetic Retinopathy Study Research Group. Preliminary report on effects of photocoagulation therapy. Am J Ophthalmol 1976;81:383–96.
- Early Treatment Diabetic Retinopathy Study research group. Photocoagulation for diabetic macular edema. Early Treatment Diabetic Retinopathy Study report number 1. Arch Ophthalmol 1985:103:1796–806
- Early Treatment Diabetic Retinopathy Study research group.
   Treatment techniques and clinical guidelines for photocoagulation of diabetic macular edema. Early Treatment Diabetic Retinopathy Study report number 2. Ophthalmology 1987;94:761–74.
- Elman MJ, Aiello LP, Beck RW, et al. Randomized trial evaluating ranibizumab plus prompt or deferred laser or triamcinolone plus prompt laser for diabetic macular edema. Ophthalmology 2010;117:1064–77.
- Ip MS, Bressler SB, Antoszyk AN, et al. A randomized trial comparing intravitreal triamcinolone and focal/grid photocoagulation for diabetic macular edema: baseline features. Retina 2008:28:919–30.
- Michaelides M, Kaines A, Hamilton RD, et al. A prospective randomized trial of intravitreal bevacizumab or laser therapy in the management of diabetic macular edema (BOLT study) 12-month data: report 2. Ophthalmology 2010;117:1078–86.
- Mitchell P, Bandello F, Schmidt-Erfurth U, et al. The RESTORE study: ranibizumab monotherapy or combined with laser versus laser monotherapy for diabetic macular edema. Ophthalmology 2011;118:615–25.
- Lovestam-Adrian M, Agardh E. Photocoagulation of diabetic macular oedema—complications and visual outcome. Acta Ophthalmol Scand 2000;78:667–71.

- Lee CM, Olk RJ. Modified grid laser photocoagulation for diffuse diabetic macular edema. Long-term visual results. *Ophthalmology* 1991;98:1594–602.
- Nwanze CC, Akinwale A, Adelman RA. Bevacizumab vs.
   Ranibizumab in preserving or improving vision in patients with wet,
   age-related macular degeneration: a cost-effectiveness review. Clin
   Med Insights: Ther 2012;4:29–38.
- Nguyen QD, Shah SM, Khwaja AA, et al. Two-year outcomes of the ranibizumab for edema of the mAcula in diabetes (READ-2) study. Ophthalmology 2010:117:2146–51.
- study. Ophthalmology 2010;117:2146–51.
   Campochiaro PA, Brown DM, Pearson A, et al. Long-term benefit of sustained-delivery fluocinolone acetonide vitreous inserts for diabetic macular edema. Ophthalmology 2011;118:626–35.
- Do DV, Schmidt-Erfurth U, Gonzalez VH, et al. The DA VINCI Study: phase 2 primary results of VEGF Trap-Eye in patients with diabetic macular edema. Ophthalmology 2011;118:1819–26.
- 31. Ahmadieh H, Ramezani A, Shoeibi N, *et al.* Intravitreal bevacizumab with or without triamcinolone for refractory diabetic macular edema; a placebo-controlled, randomized clinical trial. *Graefes Arch Clin Exp Ophthalmol* 2008;246:483–9.
- Gillies MC, Sutter FK, Simpson JM, et al. Intravitreal triamcinolone for refractory diabetic macular edema: two-year results of a double-masked, placebo-controlled, randomized clinical trial. Ophthalmology 2006;113:1533–8.
- Gillies MC, McAllister IL, Zhu M, et al. Intravitreal triamcinolone prior to laser treatment of diabetic macular edema: 24-month results of a randomized controlled trial. Ophthalmology 2011;118:866–72.
- Lam DS, Chan CK, Mohamed S, et al. Intravitreal triamcinolone plus sequential grid laser versus triamcinolone or laser alone for treating diabetic macular edema: six-month outcomes.
   Ophthalmology 2007;114:2162–7.

   Lam DS, Lai TY, Lee VY, et al. Efficacy of 1.25 MG versus 2.5 MG
- Lam DS, Lai TY, Lee VY, et al. Efficacy of 1.25 MG versus 2.5 MG intravitreal bevacizumab for diabetic macular edema: six-month results of a randomized controlled trial. Retina 2009;29:292–9.
- Massin P, Bandello F, Garweg JG, et al. Safety and efficacy of ranibizumab in diabetic macular edema (RESOLVE Study): a 12-month, randomized, controlled, double-masked, multicenter phase II study. *Diabetes Care* 2010;33:2399–405.
- Soheilian M, Ramezani A, Bijanzadeh B, et al. Intravitreal bevacizumab (avastin) injection alone or combined with triamcinolone versus macular photocoagulation as primary treatment of diabetic macular edema. Retina 2007;27:1187–95.
- Nguyen QD, Brown DM, Marcus DM, et al. Ranibizumab for diabetic macular edema: results from 2 phase III randomized trials: BISE and BIDE. Ophthalmology 2012;119:789–801
- RISE and RIDE. *Ophthalmology* 2012;119:789–801.

  39. Cunningham ET Jr, Adamis AP, Altaweel M, *et al.* A phase II randomized double-masked trial of pegaptanib, an anti-vascular endothelial growth factor aptamer, for diabetic macular edema. *Ophthalmology* 2005;112:1747–57.
- Sultan MB, Zhou D, Loftus J, et al. A phase 2/3, multicenter, randomized, double-masked, 2-year trial of pegaptanib sodium for the treatment of diabetic macular edema. Ophthalmology 2011;118:1107–18.
- Soheilian M, Ramezani A, Obudi A, et al. Randomized trial of intravitreal bevacizumab alone or combined with triamcinolone versus macular photocoagulation in diabetic macular edema. Ophthalmology 2009;116:1142–50.
- Sivaprasad S, Ockrim Z, Massaoutis P, et al. Posterior hyaloid changes following intravitreal triamcinolone and macular laser for diffuse diabetic macular edema. Retina 2008;28:1435–42.
- Pearson PA, Comstock TL, Ip M, et al. Fluocinolone acetonide intravitreal implant for diabetic macular edema: a 3-year multicenter, randomized, controlled clinical trial. Ophthalmology 2011;118:1580–7.
- Callanan D, Gupta S, Ciulla TA, et al. Efficacy and safety of combination therapy with dexamethasone intravitreal implant (DEX Implant) plus laser photocoagulation versus monotherapy with laser for treatment of diffuse diabetic macular edema (DDME) [abstract]. Invest Ophthalmol Vis Sci 2011;52:E-Abstract 3968.
- Kim Y, Kang S, Yi CH. Three-year follow-up of intravitreal triamcinolone acetonide injection and macular laser photocoagulation for diffuse diabetic macular edema [abstract]. *Invest Ophthalmol Vis Sci* 2010;51:E-Abstract 4260.
- Elman MJ, Bressler NM, Qin H, et al. Expanded 2-year follow-up of ranibizumab plus prompt or deferred laser or triamcinolone plus prompt laser for diabetic macular edema. Ophthalmology 2011;118:609–14.
- Nguyen QD, Shah SM, Heier JS, et al. Primary end point (six months) results of the ranibizumab for edema of the mAcula in diabetes (READ-2) study. Ophthalmology 2009;116:2175–81.

- Ohji M, Ishibashi T Sr, REVEAL study group. Efficacy and safety of ranibizumab 0.5 mg as monotherapy or adjunctive to laser versus laser monotherapy in Asian patients with visual impairment due to diabetic macular edema: 12-month results of the REVEAL Study labstractl. *Invest Ophthalmol Vis Sci* 2012:53:E-Abstract 4664.
- 49. Mitchell P, RESTORE extension study group. 2-Year safety and efficacy outcome of ranibizumab 0.5 mg in patients with visual impairment due to diabetic macular edema (DME): an interim analysis of the RESTORE extension study [abstract]. *Invest Ophthalmol Vis Sci* 2012;53:E-Abstract 4667.
- Do DV, Campochiaro PA, Boyer DS, et al. 6 month results of the READ 3 Study: ranibizumab for edema of the mAcula in diabetes [abstract]. Invest Ophthalmol Vis Sci 2012;53:E-Abstract 5282.
- Jorge R, Nepomuceno AB, Takaki E, et al. A prospective randomized trial of intravitreal bevacizumab versus ranibizumab for the management of refractory diabetic macular edema [abstract]. *Invest Ophthalmol Vis Sci* 2012;53:E-Abstract 347.
- Michaelides M, Fraser-Bell S, Hamilton R, et al. Macular perfusion determined by fundus fluorescein angiography at the 4-month time point in a prospective randomized trial of intravitreal bevacizumab or laser therapy in the management of diabetic macular edema (Bolt Study): report 1. Retina 2010;30:781–6.
- Faghihi H, Esfahani MR, Harandi ZA, et al. Intravitreal bevacizumab vs. combination of intravitreal bevacizumab plus macular photocoagulation in clinically significant diabetic macular edema: 6 months results of a randomized clinical trial. *Iranian J Ophthalmol* 2010;22:21–6.
- Soheilian M, Garíami KH, Ramezani A, et al. Two-year results of a randomized trial of intravitreal bevacizumab alone or combined with triamcinolone versus laser in diabetic macular edema. Retina 2012;32:314–21.
- 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;227:100–6.
- Oliveira Neto HL, Andrade RE, Casella M, et al. A randomized clinical trial to compare the efficacy and safety of isolated or combined intravitreal injection of triamcinolone acetonide and bevacizumab for diabetic macular edema ATEMD protocol—a Brazilian clinical trial [abstract]. Invest Ophthalmol Vis Sci 2011;52:E-Abstract 5331.
- Adamis AP, Altaweel M, Bressler NM, et al. Changes in retinal neovascularization after pegaptanib (Macugen) therapy in diabetic individuals. Ophthalmology 2006;113:23–8.
- Do DV, Nguyen QD, Boyer D, et al. One-year outcomes of the DA VINCI Study of VEGF Trap-Eye in eyes with diabetic macular edema. Ophthalmology 2012;119:1658–65.
- Haller JA, Kuppermann BD, Blumenkranz MS, et al. Randomized controlled trial of an intravitreous dexamethasone drug delivery system in patients with diabetic macular edema. Arch Ophthalmol 2010;128:289–96.
- Campochiaro PA, Brown DM, Pearson A, et al. Sustained delivery fluocinolone acetonide vitreous inserts provide benefit for at least 3 Years in patients with diabetic macular edema. Ophthalmology 2012;119:2125–32.
- Beck RW, Edwards AR, Aiello LP, et al. Diabetic Retinopathy Clinical Research Network (DRCR.net). Three-year follow-up of a randomized trial comparing focal/grid photocoagulation and intravitreal triamcinolone for diabetic macular edema. Arch Ophthalmol 2009;127:245–51.
- Ockrim ZK, Sivaprasad S, Falk S, et al. Intravitreal triamcinolone versus laser photocoagulation for persistent diabetic macular oedema. Br J Ophthalmol 2008;92:795–9.
- Bressler NM, Edwards AR, Beck RW, et al. Exploratory analysis
  of diabetic retinopathy progression through 3 years in a
  randomized clinical trial that compares intravitreal triamcinolone
  acetonide with focal/grid photocoagulation. Arch Ophthalmol
  2009;127:1566–71.
- Diabetic Retinopathy Clinical Research Network. A randomized trial comparing intravitreal triamcinolone acetonide and focal/grid photocoagulation for diabetic macular edema. *Ophthalmology* 2008;115:1447–9.
- Ford JA, Elders A, Shyangdan D, et al. The relative clinical effectiveness of ranibizumab and bevacizumab in diabetic macular oedema: an indirect comparison. BMJ 2012;345:e5182.
- Chakravarthy U, Harding SP, Rogers CA, et al. Ranibizumab versus bevacizumab to treat neovascular age-related macular degeneration: one-year findings from the IVAN randomized trial. Ophthalmology 2012;119:1399–411.
- Martin DF, Maguire MG, Fine SL, et al. Ranibizumab and bevacizumab for treatment of neovascular age-related macular degeneration: two-year results. Ophthalmology 2012;119:1388–98.

- Campbell RJ, Gill SS, Bronskill SE, et al. Adverse events with intravitreal injection of vascular endothelial growth factor inhibitors: nested case-control study. BMJ 2012;345:e4203.
- Van der Reis MI, La Heij EC, De Jong-Hesse Y, et al. A systematic review of the adverse events of intravitreal anti-vascular endothelial growth factor injections. Retina 2011;31:1449–69.
- Fung AE, Rosenfeld PJ, Reichel E. The International Intravitreal Bevacizumab Safety Survey: using the internet to assess drug safety worldwide. Br J Ophthalmol 2006;90:1344–9.
- Elbendary AM, Shahin MM. Intravitreal diclofenac versus intravitreal triamcinolone acetonide in the treatment of diabetic macular edema. *Retina* 2011;31:2058–64.
- Sfikakis PP, Grigoropoulos V, Emfietzoglou I, et al. Infliximab for diabetic macular edema refractory to laser photocoagulation: a randomized, double-blind, placebo-controlled, crossover, 32-week study. Diabetes Care 2010;33:1523

  –8.
- Keech AC, Mitchell P, Summanen PA, et al. Effect of fenofibrate on the need for laser treatment for diabetic retinopathy (FIELD study): a randomised controlled trial. Lancet 2007;370:1687–97.
- PKC-DMES Study Group. Effect of ruboxistaurin in patients with diabetic macular edema: thirty-month results of the randomized PKC-DMES clinical trial. Arch Ophthalmol 2007;125:318–24.
- PKC-DMES Study Group. Effect of ruboxistaurin on visual loss in patients with diabetic retinopathy. *Ophthalmology* 2006:113:2221–30.
- Fong DS, Strauber SF, Aiello LP, et al. Comparison of the modified Early Treatment Diabetic Retinopathy Study and mild macular grid laser photocoagulation strategies for diabetic macular edema. Arch Ophthalmol 2007;125:469–80.
- Health Canada. Reports of eye inflammation, endophthalmitis, and Toxic Anterior Segment Syndrome (TASS) following off-label intravitreal use of Avastin (bevacizumab). 2008 [cited 24 Oct 2012]; http://www.hc-sc.gc.ca/dhp-mps/medeff/advisories-avis/prof/\_2008/ avastin\_4\_hpc-cps-eng.php
- Curtis LH, Hammill BG, Schulman KA, et al. Risks of mortality, myocardial infarction, bleeding, and stroke associated with therapies for age-related macular degeneration. Arch Ophthalmol 2010;128:1273–9.
- Gower EW, Cassard S, Chu L, et al. Adverse event rates following intravitreal injection of Avastin or Lucentis for treating age-related macular degeneration [abstract]. Invest Ophthalmol Vis Sci 2011;52:E-Abstract 6644.
- General Medical Council. Prescribing medicines for use outside the terms of their licence (off-label). 2012 [cited 24 Oct 2012];http:// www.gmc-uk.org/guidance/ethical\_guidance/prescriptions\_faqs. asn#10
- U.S.Food and Drug Administration. Off-Label" and investigational use of marketed drugs, biologics, and medical devices— Information Sheet. 2011 [cited 24 Oct 2012];http://www.fda.gov/ RegulatoryInformation/Guidances/ucm126486.htm
- Alimera Sciences. Alimera Sciences' ILUVIEN Receives Marketing Authorization in France for the Treatment of Chronic Diabetic Macular Edema. 2012 [cited 24 Oct 2012];http://investor. alimerasciences.com/releasedetail.cfm?ReleaseID=692876
- Laidlaw DA. Vitrectomy for diabetic macular oedema. Eye 2008:22:1337–41.
- 84. National Institute for Health and Clinical Excellence. Ranibizumab for the treatment of diabetic macular oedema:TA237. 2011 [cited 24 Oct 2012];http://publications.nice.org.uk/ranibizumab-for-the-treatment-of-diabetic-macular-oedema-ta237
- Rajendram R, Fraser-Bell S, Kaines A, et al. A 2-year prospective randomized controlled trial of intravitreal bevacizumab or laser therapy (BOLT) in the management of diabetic macular edema: 24-month data: report 3. Arch Ophthalmol 2012;130:972–9.
- Sivaprasad S, Dorin G. Subthreshold diode laser micropulse photocoagulation for the treatment of diabetic macular edema. Expert Rev Medical Devices 2012;9:189–97.
- Cho WB, Moon JW, Kim HC. Intravitreal triamcinolone and bevacizumab as adjunctive treatments to panretinal photocoagulation in diabetic retinopathy. *Br J Ophthalmol* 2010:94:858–63.
- Googe J, Brucker AJ, Bressler NM, et al. Randomized trial evaluating short-term effects of intravitreal ranibizumab or triamcinolone acetonide on macular edema after focal/grid laser for diabetic macular edema in eyes also receiving panretinal photocoagulation. Retina 2011;31:1009–27.
- Faghihi H, Roohipoor R, Mohammadi SF, et al. Intravitreal bevacizumab versus combined bevacizumab-triamcinolone versus macular laser photocoagulation in diabetic macular edema. Eur J Ophthalmol 2008;18:941–8.

- Figueroa MS, Contreras I, Noval S. Surgical and anatomical outcomes of pars plana vitrectomy for diffuse nontractional diabetic macular edema. *Retina* 2008;28:420–6.
- Isaac DL, Abud MB, Frantz KA, et al. Comparing intravitreal triamcinolone acetonide and bevacizumab injections for the treatment of diabetic macular oedema: a randomized double-blind study. Acta Ophthalmol 2012;90:56–60.
- Paccola L, Costa RA, Folgosa MS, et al. Intravitreal triamcinolone versus bevacizumab for treatment of refractory diabetic macular oedema (IBEME study). Br J Ophthalmol 2008;92:76–80.
- Prager SG, Kriechbaum K, Mylonas G, et al. Comparison of intravitreally applied bevacizumab and triamcinolone on diabetic macular edema [abstract]. Invest Ophthalmol Vis Sci 2010;51: E-Abstract 4262.
- Ozturk BT, Kerimoglu H, Bozkurt B, et al. Comparison of intravitreal bevacizumab and ranibizumab treatment for diabetic macular edema. J Ocul Pharmacol Ther 2011;27:373–7.
- Marey HM, Ellakwa AF. Intravitreal bevacizumab alone or combined with triamcinolone acetonide as the primary treatment for diabetic macular edema. Clin Ophthalmol 2011;5:1011–16.
- Shahin MM, El-Lakkany RS. A prospective, randomized comparison of intravitreal triamcinolone acetonide versus intravitreal bevacizumab (avastin) in diffuse diabetic macular edema. *Middle East Afr J Ophthalmol* 2010;17:250–3.
- Loftus JV, Sultan MB, Pleil AM, et al. Changes in vision- and health-related quality of life in patients with diabetic macular edema treated with pegaptanib sodium or sham. *Invest Ophthalmol Vis Sci* 2011;52:7498–505.
- Ferrone PJ, Jonisch J. Ranibizumab dose comparison for the treatment of diabetic macular edema [abstract]. *Invest Ophthalmol Vis Sci* 2011;52:E-Abstract 5333.
- Solaiman KA, Diab MM, Abo-Elenin M. Intravitreal bevacizumab and/or macular photocoagulation as a primary treatment for diffuse diabetic macular edema. *Retina* 2010;30:1638–45.
- Scott IU, Edwards AR, Beck RW, et al. Diabetic Retinopathy Clinical Research Network. A phase II randomized clinical trial of intravitreal bevacizumab for diabetic macular edema. Ophthalmology 2007;114:1860–7.
- Lee SJ, Kim ET, Moon YS. Intravitreal bevacizumab alone versus combined with macular photocoagulation in diabetic macular edema. Korean J Ophthalmol 2011;25:299–304.
- Audren F, Lecleire-Collet A, Erginay A, et al. Intravitreal triamcinolone acetonide for diffuse diabetic macular edema: phase 2 trial comparing 4 mg vs 2 mg. Am J Ophthalmol 2006;142:794–9.
- Audren F, Erginay A, Haouchine B, et al. Intravitreal triamcinolone acetonide for diffuse diabetic macular oedema: 6-month results of a prospective controlled trial. Acta Ophthalmol Scand 2006;84:624–30.
- Avitabile T, Longo A, Reibaldi A. Intravitreal triamcinolone compared with macular laser grid photocoagulation for the treatment of cystoid macular edema. Am J Ophthalmol 2005;140:695–702.
- Bandello F, Pognuz DR, Pirracchio A, et al. Intravitreal triamcinolone acetonide for florid proliferative diabetic retinopathy. Graefes Arch Clin Exp Ophthalmol 2004;242:1024–7.
- Bonini MA, Jorge R, Barbosa JC, et al. Intravitreal injection versus sub-Tenon's infusion of triamcinolone acetonide for refractory diabetic macular edema: a randomized clinical trial. Invest Ophthalmol Vis Sci 2005;46:3845–9.
- Cellini M, Pazzaglia A, Zamparini E, et al. Intravitreal vs. subtenon triamcinolone acetonide for the treatment of diabetic cystoid macular edema. BMC Ophthalmol 2008;8:5TN.
- Cardillo JA, Melo LA Jr, Costa RA, et al. Comparison of intravitreal versus posterior sub-Tenon's capsule injection of triamcinolone acetonide for diffuse diabetic macular edema. Ophthalmology 2005;112:1557–63.
- 109. Chung EJ, Freeman WR, Azen SP, et al. Comparison of combination posterior sub-tenon triamcinolone and modified grid laser treatment with intravitreal triamcinolone treatment in patients with diffuse diabetic macular edema. Yonsei Medi J 2008;49:955–64.
- Dehghan MH, Ahmadieh H, Ramezani A, et al. A randomized, placebo-controlled clinical trial of intravitreal triamcinolone for refractory diabetic macular edema. Int Ophthalmol 2008;28:7–17.
- 111. Chew E, Strauber S, Beck RW, et al. Diabetic Retinopathy Clinical Research Network. Randomized trial of peribulbar triamcinolone acetonide with and without focal photocoagulation for mild diabetic macular edema: a pilot study. Ophthalmology 2007;114:1190–6.
- 112. Gil AL, Azevedo MJ, Tomasetto GG, et al. Treatment of diffuse diabetic maculopathy with intravitreal triamcinolone and laser photocoagulation: randomized clinical trial with morphological and

- functional evaluation. Arq Bras *Arquivos Brasileiros de Oftalmol* 2011:74:343–7.
- Entezari M, Ahmadieh H, Dehghan MH, et al. Posterior sub-tenon triamcinolone for refractory diabetic macular edema: a randomized clinical trial. Eur J Ophthalmol 2005:15:746–50.
- Hauser D, Bukelman A, Pokroy R, et al. Intravitreal triamcinolone for diabetic macular edema: comparison of 1, 2, and 4 mg. Retina 2008;28:825–30.
- Jonas JB, Kamppeter BA, Harder B, et al. Intravitreal triamcinolone acetonide for diabetic macular edema: a prospective, randomized study. J Ocul Pharmacol Ther 2006;22:200–7.
- Joussen AM, Weiss C, Bauer D, et al. Triamcinolone versus inner-limiting membrane peeling in persistent diabetic macular edema (TIME study): design issues and implications. Graefes Arch Clin Exp Ophthalmol 2007:245:1781–7.
- Kaderli B, Avci R. Comparison of topical and subconjunctival anesthesia in intravitreal injection administrations. Eur J Ophthalmol 2006;16:718–21.
- Kang SW, Sa HS, Cho HY, et al. Macular grid photocoagulation after intravitreal triamcinolone acetonide for diffuse diabetic macular edema. Arch Ophthalmol 2006;124:653–8.
- Kim JE, Pollack JS, Miller DG, et al. ISIS-DME: a prospective, randomized, dose-escalation intravitreal steroid injection study for refractory diabetic macular edema. Retina 2008;28:735–40.
- Lam DS, Chan CK, Mohamed S, et al. A prospective randomised trial of different doses of intravitreal triamcinolone for diabetic macular oedema. Br J Ophthalmol 2007;91:199–203.
- Lee HY, Lee SY, Park JS. Comparison of photocoagulation with combined intravitreal triamcinolone for diabetic macular edema. Korean J Ophthalmol 2009;23:153–8.
- 122. Maia OO Jr, Takahashi BS, Costa RA, et al. Combined laser and intravitreal triamcinolone for proliferative diabetic retinopathy and macular edema: one-year results of a randomized clinical trial. Am J Ophthalmol 2009;147:291–7.
- 123. Massin P, Audren F, Haouchine B, et al. Intravitreal triamcinolone acetonide for diabetic diffuse macular edema: preliminary results of a prospective controlled trial. Ophthalmology 2004;111:218–24.
- Mohamed S, Leung GM, Chan CK, et al. Factors associated with variability in response of diabetic macular oedema after intravitreal triamcinolone. Clin Experiment Ophthalmol 2009;37:602–8.
- Nakamura A, Shimada Y, Horio N, et al. Vitrectomy for diabetic macular edema with posterior subtenon injection of triamcinolone acetonide. [Japanese]. Folia Ophthalmol Japonica 2004;55:958–62.
- Spandau UH, Derse M, Schmitz-Valckenberg P, et al. Dosage dependency of intravitreal triamcinolone acetonide as treatment for diabetic macular oedema. Br J Ophthalmol 2005;89:999–1003.
- Tunc M, Onder HI, Kaya M. Posterior sub-Tenon's capsule triamcinolone injection combined with focal laser photocoagulation for diabetic macular edema. *Ophthalmology* 2005;112:1086–91.
- 128. Verma LK, Vivek MB, Kumar A, et al. A prospective controlled trial to evaluate the adjunctive role of posterior subtenon triamcinolone in the treatment of diffuse diabetic macular edema. J Ocul Pharmacol Ther 2004;20:277–84.
- Wickremasinghe SS, Rogers SL, Gillies MC, et al. Retinal vascular caliber changes after intravitreal triamcinolone treatment for diabetic macular edema. *Invest Ophthalmol Vis Sci* 2008;49:4707–11.
- Yalcinbayir O, Gelisken O, Kaderli B, et al. Intravitreal versus sub-Tenon posterior triamcinolone injection in bilateral diffuse diabetic macular edema. Ophthalmologica 2011;225:222–7.
- Haller JA, Bandello F, Belfort R Jr, et al. Randomized, sham-controlled trial of dexamethasone intravitreal implant in patients with macular edema due to retinal vein occlusion. Ophthalmology 2010;117:1134–46.
- 132. Haller JA, Dugel P, Weinberg DV, et al. Evaluation of the safety and performance of an applicator for a novel intravitreal dexamethasone drug delivery system for the treatment of macular edema. Retina 2009;29:46–51.
- Kuppermann BD, Blumenkranz MS, Haller JA, et al. Randomized controlled study of an intravitreous dexamethasone drug delivery system in patients with persistent macular edema. Arch Ophthalmol 2007;125:309–17.
- Boyer DS, Faber D, Gupta S, et al. Dexamethasone intravitreal implant for treatment of diabetic macular edema in vitrectomized patients. Retina 2011;31:915–23.
- Campochiaro PA, Hafiz G, Shah SM, et al. Sustained ocular delivery of fluocinolone acetonide by an intravitreal insert. Ophthalmology 2010;117:1393–9.

- Gillies MC, Islam FM, Zhu M, et al. Efficacy and safety of multiple intravitreal triamcinolone injections for refractory diabetic macular oedema. Br J Ophthalmol 2007;91:1323–6.
- Gillies MC, Simpson JM, Gaston C, et al. Five-year results of a randomized trial with open-label extension of triamcinolone acetonide for refractory diabetic macular edema. Ophthalmology 2009;116:2182–7.
- Sutter FK, Simpson JM, Gillies MC. Intravitreal triamcinolone for diabetic macular edema that persists after laser treatment: three-month efficacy and safety results of a prospective, randomized, double-masked, placebo-controlled clinical trial. *Ophthalmology* 2004:111:2044–9
- 139. Brown DM, Nguyen QD, Rubio RG, et al. Ranibizumab for Diabetic Macular Edema (DME): 24-Month Efficacy and Safety Results of RISE—a phase 3 randomized controlled trial [abstract]. Invest Ophthalmol Vis Sci 2011;52:E-Abstract 6647.
- 140. Boyer D, Sy J, Rundle AC, et al. Ranibizumab (Anti-VEGF) for vision loss due to diabetic macular edema—results of two phase III randomized trials [abstract]. 71st Scientific Sessions June 24–28, 2011, San Diego Convention Center—San Diego, California, USA2011;Abstract No, 133-LBOR.
- Soheilian M, Ramezani A, Yaseri M, et al. Initial macular thickness and response to treatment in diabetic macular edema. Retina 2011;31:1564–73.

# APPENDIX 1: METHODS OF THE LITERATURE SEARCH Searches for clinical trials

Ovid MEDLINE 1948-week 2 July 2012 and Ovid MEDLINE(R) In-Process and Other Non-Indexed Citations 11 July 2012

- 1. Diabetic Retinopathy/dt (Drug Therapy)
- 2. Macular Edema/dt (Drug Therapy)
- 3. (diabet\* adj2 macular adj (edema or oedema)).tw.
- 4. (diabet\* adj2 maculopathy).tw.
- 5. (diabet\* adj2 retinopathy).tw.
- 6. 1 or 2 or 3 or 4 or 5
- (ranibizumab or lucentis or bevacizumab or avastin or pegaptanib or macugen or aflibercept or vegf trap-eye or steroid\* or corticosteroid\* or dexamethasone or fluocinolone or triamcinolone or anti-VEGF\* or anti-vascular endothelial growth factor\*).tw.
- 8. exp Vascular Endothelial Growth Factor A/
- 9. exp Fluocinolone Acetonide/
- 10. exp Triamcinolone/
- 11. 7 or 8 or 9 or 10
- 12. 6 and 11
- 13. randomised controlled trial.pt.
- 14. controlled clinical trial.pt.
- 15. (masked or sham or placebo or control group or random\*).tw.
- 16. 13 or 14 or 15
- 17. 12 and 16
- 18. (case reports or editorial or letter or review).pt.
- 19. 17 not 18
- 20. limit 19 to humans

EMBASE 1947-2012 week 27

- (ranibizumab or lucentis or bevacizumab or avastin or pegaptanib or macugen or aflibercept or vegf trap-eye or dexamethasone or fluocinolone or triamcinolone or anti-VEGF\* or anti-vascular endothelial growth factor\*).m\_titl.
- (diabetic macular edema or diabetic macular oedema or diabetic retinopathy or diabetic maculopathy).m\_titl.
- 3. 1 and 2
- 4. random\*.tw.
- 5. 3 and 4

Cochrane Central Register of Controlled Trials, Issue 7 of 12, July 2012

Ranibizumab or lucentis or bevacizumab or avastin or pegaptanib or macugen or aflibercept or vegf trap-eye or steroid\* or corticosteroid\* or dexamethasone or fluocinolone or triamcinolone or anti-VEGF\* or anti-vascular endothelial growth factor\* in Record Title and diabetic macular edema or diabetic macular oedema or diabetic retinopathy or diabetic maculopathy in Record Title

Web of Science—with Conference Proceedings (updated 12 July 2012)

Title=(ranibizumab or lucentis or bevacizumab or avastin or pegaptanib or macugen or aflibercept or vegf trap-eye or steroid\* or

corticosteroid\* or dexamethasone or fluocinolone or triamcinolone or anti-VEGF\* or anti-vascular endothelial growth factor\*) AND Title=(diabetic macular edema or diabetic macular oedema or diabetic retinopathy or diabetic maculopathy) AND Title=(random\*)

# Searches for systematic reviews

Ovid MEDLINE(R) Daily Update 11 July 2012, Ovid MEDLINE(R) In-Process and Other Non-Indexed Citations 11 July 2012

- 1. Diabetic Retinopathy/dt (Drug Therapy)
- 2. Macular Edema/dt (Drug Therapy)
- 3. (diabet\* adj2 macular adj (edema or oedema)).tw.
- 4. (diabet\* adj2 maculopathy).tw.
- 5. (diabet\* adj2 retinopathy).tw.
- 6. 1 or 2 or 3 or 4 or 5
- (ranibizumab or lucentis or bevacizumab or avastin or pegaptanib or macugen or aflibercept or vegf trap-eye or steroid\* or corticosteroid\* or dexamethasone or fluocinolone or triamcinolone or anti-VEGF\* or anti-vascular endothelial growth factor\*).tw.
- 8. exp Vascular Endothelial Growth Factor A/
- 9. exp Fluocinolone Acetonide/
- 10. exp Triamcinolone/
- 11. 7 or 8 or 9 or 10
- 12. 6 and 11
- 13. (systematic review or meta-analysis or pubmed or medline).tw.
- 14. meta-analysis.pt.
- 15. cochrane.af.
- 16. 13 or 14 or 15
- 17. 12 and 16

Cochrane Database of Systematic Reviews and Technology Assessments Database, Cochrane Library July Issue, 2012

Ranibizumab or lucentis or bevacizumab or avastin or pegaptanib or macugen or aflibercept or vegf trap-eye or steroid\* or corticosteroid\* or dexamethasone or fluocinolone or triamcinolone or anti-VEGF\* or anti-vascular endothelial growth factor\* in Record Title and diabetic macular edema or diabetic macular oedema or diabetic retinopathy or diabetic maculopathy in Record Title

# Searches for safety and adverse events

Ovid MEDLINE(R) Daily Update 11 July 2012, Ovid MEDLINE(R) In-Process and Other Non-Indexed Citations 11 July 2012; EMBASE 1980–2012 week 27

- (ranibizumab or lucentis or bevacizumab or avastin or pegaptanib or aflibercept or vegf trap-eye or macugen or dexamethasone or fluocinolone or triamcinolone or anti-VEGF\* or anti-vascular endothelial growth factor\*).m\_titl.
- 2. (diabetic macular edema or diabetic macular oedema or diabetic retinopathy or diabetic maculopathy).m\_titl.
- 3. 1 and 2
- 4. (risk or safety or adverse or harm or pharmacovigilance).tw.
- (side-effect\* or precaution\* or warning\* or contraindication\$ or contra-indication\* or tolerability or toxic\*).tw.
- 6. 4 or 5
- 7. 3 and 6

# Searches of the annual meeting abstracts (for trials, reviews and safety studies)

- ARVO (Association for Research in Vision and Ophthalmology) (2002–2012)
- ▶ ADA (American Diabetes Association) (2002–2012)
- ► EASD (European Association for the Study of Diabetes) (2002–2012)

# Other searches

Web sites of the following

Drugs@FDA: FDA Approved Drug Products

- ► European Medicines Association
- ▶ ClinicalTrials.gov
- EU Clinical Trials Register
   National Institute for Health and Clinical Excellence

#### APPENDIX 2: ONGOING TRIALS IN CLINICALTRIALS.GOV

- Schmidt-Erfurth and colleagues are comparing ranibizumab and bevacizumab in DME (NCT00545870)
- ► TRIASTIN study is comparing ranibizumab, triamcinolone and sham injection (NCT00682539)
- ► Maturi and colleagues are comparing bevacizumab plus dexamethasone with bevacizumab alone (NCT01309451)
- ▶ IBeTA study (Jorge and colleagues) is comparing bevacizumab (1.5 mg) plus laser, triamcinolone (4 mg) plus laser with laser alone (NCT00997191)
- ► Chaudhry and colleagues are evaluating ranibizumab in patients who have failed with 3–6 injections of bevacizumab (NCT01253694)
- ► MIDME study (Pfizer) is comparing pegaptanib 0.3 mg with sham injection (NCT01175070)
- ► Figueira and colleagues are comparing pegaptanib plus laser with laser alone (NCT01281098)
- ▶ RESPOND (Novartis) is comparing ranibizumab (0.5 mg) alone with ranibizumab plus laser or laser alone (NCT01135914)
- RETAIN (Novartis) study is comparing two different ranibizumab algorithms; 'treat and extend' versus as needed (NCT01171976)
- ► RED-ES (Novartis) is comparing ranibizumab with laser in patients with visual impairment due to DME (NCT00901186)
- READ 3 study (Do and colleagues) are comparing two doses of ranibizumab 0.5 and 2 mg (NCT01077401)
- VIVID-DME and VISTA DME studies (Bayer) are comparing aflibercept with laser. (NCT01331681 and NCT01363440)
- ▶ Gillies and colleagues are comparing bevacizumab with dexamethasone (NCT01298076)
- Soheilian and colleagues are performing a phase I study looking at the use of diclofenac compared with bevacizumab in DME (NCT00999791)
- López-Miranda and colleagues are comparing the use of bevacizumab before and after laser therapy (NCT00804206)
- NEVANAC study is comparing triamcinolone alone with triamcinolone plus nepafenac (NSAID) (NCT00780780)
- ▶ Elman and colleagues are comparing laser alone, laser combined with an intravitreal injection of triamcinolone, laser combined with an intravitreal injection of ranibizumab, or intravitreal injection of ranibizumab alone (NCT00444600)
- BRDME (Schlingemann and collagues) study is comparing the use of bevacizumab and ranibizumab in the treatment of patients with DME (OCT central area thickness > 275 μm) (NCT01635790)
- Wiley and colleagues are comparing bevacizumab and ranibizumab in patients with DME in at least one eye (NCT01610557)
- Protocol T study (Wells and colleagues) is comparing effectiveness of a aflibercept, bevacizumab and ranibizumab for DME (NCT01627249)
- Allergan-funded study comparing safety and efficacy of 700 μg dexamethasone implant against 0.5 mg ranibizumab in patients with DME (NCT01492400)
- Pfizer-funded study comparing effectiveness of 0.3 mg pegaptanib against sham injection (NCT01100307)
- Allergan-funded study comparing safety and efficacy of an intravitreal dexamethasone implant (700 and 350 μg) against sham in patients with DME (NCT00168389)
- Allergan-funded study comparing safety and efficacy of an intravitreal dexamethasone implant (700 and 350 μg) against sham in patients with DME (NCT00168337)



# Current treatments in diabetic macular oedema: systematic review and meta-analysis

John Alexander Ford, Noemi Lois, Pamela Royle, Christine Clar, Deepson Shyangdan and Norman Waugh

BMJ Open 2013 3:

doi: 10.1136/bmjopen-2012-002269

Updated information and services can be found at: http://bmjopen.bmj.com/content/3/3/e002269

These include:

**References** This article cites 134 articles, 19 of which you can access for free at:

http://bmjopen.bmj.com/content/3/3/e002269#BIBL

**Open Access** this is an open-access article distributed under the terms of the creative

commons attribution non-commercial license, which permits use, distribution, and reproduction in any medium, provided the original work is properly cited, the use is non commercial and is otherwise in compliance with the license. see: http://creativecommons.org/licenses/by-nc/2.0/ and

http://creativecommons.org/licenses/by-nc/2.0/legalcode.

Email alerting service Receive free email alerts when new articles cite this article. Sign up in the

box at the top right corner of the online article.

Topic Collections

Articles on similar topics can be found in the following collections

Ophthalmology (105)

Pharmacology and therapeutics (430)

# **Notes**

To request permissions go to: http://group.bmj.com/group/rights-licensing/permissions

To order reprints go to: http://journals.bmj.com/cgi/reprintform

To subscribe to BMJ go to: http://group.bmj.com/subscribe/