The role of anti-vascular endothelial growth factor (anti-VEGF) in the management of proliferative diabetic retinopathy

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Abstract

Diabetes is a major cause of visual impairment among working-age adults in the United States. The proliferative form of diabetic retinopathy is associated with severe vision loss (acuity <5/200). The standard treatment in proliferative diabetic retinopathy (PDR) is panretinal photocoagulation (PRP), which is effective but has established side effects such as peripheral visual-field constraints. Vascular endothelial growth factor (VEGF) is thought to drive the process of vascular proliferation. Drugs targeting VEGF (anti-VEGF) have been studied extensively in diabetic macular edema (DME), and results have shown that diabetic retinopathy regresses with anti-VEGF treatment. Recent studies show that anti-VEGF is not inferior to PRP for PDR while treatment is maintained, though recurrence rate when anti-VEGF treatment is stopped is unclear. In vitreous hemorrhage where PRP cannot be performed, use of anti-VEGF medications can treat underlying PDR and delay or reduce need for vitrectomy. Limitations of anti-VEGF treatment, however, require careful patient selection and monitoring. This review discusses recent clinical trials and guidelines for anti-VEGF use in PDR.

Keywords: aflibercept, angiogenesis inhibitors, antibodies, monoclonal, humanized antibodies, bevacizumab, diabetic retinopathy, intravitreal injection, ranibizumab, vascular endothelial growth factor A.

Citation

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Introduction

Diabetes mellitus is the leading cause of blindness among adults aged 20–64 in the United States.¹ The National Diabetes Statistics Report published by the Centers for Disease Control and Prevention (CDC) estimated that 9.4% of the US population suffers from diabetes.² Of patients with diabetes, 28.5% had evidence of diabetic retinopathy, and 4.4% had vision-threatening retinopathy.³ Proliferative diabetic retinopathy (PDR) is characterized by growth of neovascular vessels, which are prone to leakage, bleeding, and the development of vitreoretinal membranes and tractional retinal detachment. They can also invade the anterior segment to cause neovascular glaucoma or ischemia. Risk of vision loss is significantly higher with PDR than nonproliferative diabetic retinopathy (NPDR), and PDR is more prevalent in type 1 diabetes, where patients present at a younger age.⁴ The Diabetic Retinopathy Study (DRS) found that almost half of eyes with PDR and high-risk characteristics will progress to severe vision loss (visual acuity <5/200) without treatment.⁵

Panretinal photocoagulation has been the standard of care for PDR for decades. However, recent data demonstrate that anti-vascular endothelial growth factor (VEGF) intravitreal injections can be used to treat PDR without photocoagulation of peripheral retina. The purpose of this review is to discuss the encouraging results of anti-VEGF use in PDR and highlight the limitations of anti-VEGF treatment that necessitate careful patient selection and monitoring. A PubMed literature search of articles up to February 2018 was performed using keywords, ‘proliferative diabetic retinopathy’ and ‘anti-VEGF.’ The Diabetic Retinopathy Clinical Research Network (DRCRnet) publications were also reviewed. Articles and their references were assessed and included as deemed relevant by the authors.

VEGF drives PDR

PDR is a microvascular disease, where relative retinal ischemia produces a pro-angiogenic environment. Angiogenesis is mediated in large part by VEGF.⁶ ⁷ The VEGF family consists of
multiple isosforms that are required for the normal development of the vasculature and lymphatics. Molecular studies have shown that VEGF-A promoted vascular permeability and angiogenesis through its interactions with VEGF receptor 2 (VEGFR-2) on vascular endothelial cells. Capillary endothelial tight-junction disruption and development of endothelial cell fenestration weakened blood vessels. VEGF-A also stimulated endothelial cell proliferation and migration — changes associated with early angiogenesis.  

In the eye, the VEGF-A165 splice variant of VEGF-A has been implicated as the cause of pathologic revascularization in the retina. Injection of VEGF into animal eyes produced a phenotype similar to diabetic or ischemic retinopathy with intraretinal hemorrhage, vessel tortuosity, retinal edema, and intraretinal vascular proliferations. In contrast, inhibition of VEGF prevented iris neovascularization. Human studies showed that eyes with PDR had higher levels of VEGF in vitreous or fibrovascular tissues than normal eyes.

**Panretinal photocoagulation in PDR**

The standard of care in PDR has been panretinal photocoagulation (PRP), as the DRS study demonstrated that PRP reduced the rate of severe vision loss by >50%. The thermal destruction of ischemic retina decreases the signal for PRP reduced the rate of severe vision loss by >50%. The thermal destruction of ischemic retina decreases the signal for PRP reduced the rate of severe vision loss by >50%. The thermal destruction of ischemic retina decreases the signal for PRP and regression of PDR was confirmed on exam. However, PRP has well-established limitations. It requires a cooperative patient and a clear view of the retina. Peripheral vision is permanently lost with the destruction of peripheral retina, and nyctalopia and transient decline in visual acuity (VA) have also been reported. Macular edema can develop or worsen after PRP, adversely affecting vision. In a series of 175 eyes treated with PRP, McDonald and Schatz found that 8% of eyes developed chronic macular edema leading to two lines or greater vision loss. More recently, Protocol S found that the cumulative 2-year probability of developing visually significant central diabetic macular edema (DME) was 28% in the PRP-treated group. Additionally, laser-associated breaks in Bruch’s membrane can lead to development of choroidal neovascularization, and damage to the posterior ciliary nerves can lead to mydriasis and loss of accommodation. Other side effects include uveal effusion, angle closure glaucoma, serous retinal detachment, and vitreous hemorrhage. In recent years, the development of molecules that target VEGF itself has changed the approach to diabetic retinopathy. There are four therapies that target VEGF, including three antibodies against VEGF (anti-VEGF).

**Current therapies targeting VEGF**

Pegaptanib (Macugen; Eyetech Inc, Cedar Knolls, NJ, USA) is a 28 nucleotide RNA aptamer that binds to the VEGF-A165 isomer. It was studied and used predominantly in the treatment of DME, and its use has declined in favor of antibody-based treatments. The Macugen Diabetic Retinopathy Study Group included 16 eyes with PDR, of which 13 received pegaptanib. The study noted that eight underwent progression of neovascularization. Once the medication was stopped, neovascularization progression resumed. Bevacizumab (Avastin; Genentech, San Francisco, CA, USA) is a full-length recombinant humanized anti-VEGF monoclonal antibody (IgG) initially Food and Drug Administration (FDA) approved for treatment of metastatic colorectal cancers. It has twice the half-life of ranibizumab; Ophthalmic uses of bevacizumab are not FDA approved; however, its safety and efficacy have been shown in multiple neovascular age-related macular degeneration trials and DME trials, such as DRCR.net protocol T. Protocol T compared mean improvement in VA after intravitreal aflibercept, ranibizumab, or bevacizumab treatment in DME. It found that in eyes with VA of 20/40 or better, there was no significant difference in mean improvement in VA among bevacizumab (7.5±7.4 Early Treatment Diabetic Retinopathy Study (ETDRS) letters), ranibizumab (8.3±6.8), or aflibercept (8.0±7.6). In eyes with vision 20/50 or worse, aflibercept (18.9±11.5 letters) had better outcomes than bevacizumab (11.8±12) or ranibizumab (14.2±10.6) at 1-year follow-up. This difference continued to be significant between aflibercept and bevacizumab at 2-year follow-up. No significant difference in serious adverse reactions, major cardiovascular events, or hospitalization among the three medications was found. Ranibizumab (Lucentis; Genentech, San Francisco, CA, USA/Novartis Ophthalmics, Basel, Switzerland) is a recombinant antibody fragment of the humanized anti-VEGF monoclonal antibody. Based on bevacizumab, ranibizumab underwent affinity maturation to increase binding affinity to all isoforms of VEGF. Its half-life is shorter than other anti-VEGF molecules due to the lack of the antibody Fc domain. The RISE and RIDE studies, discussed later, demonstrated efficacy of ranibizumab in DME. Based on multiple trials, the FDA approved ranibizumab in 2017 for all stages of diabetic retinopathy.

Aflibercept (Eylea; Regeneron, Tarrytown, NY, USA) is a recombinant fusion protein of the binding domains of human VEGF-R1 and VEGF-R2, fused with the Fc domain of human IgG1. It binds endogenous VEGF molecules to prevent their activation of VEGF-R, and it has been shown to bind VEGF with greater affinity than other anti-VEGF agents. Additionally, it can bind placental growth factor (PIGF). It was FDA-approved for the treatment of neovascular age-related macular degeneration (AMD) in 2011. The VISTA and VIVID trials studied aflibercept use in DME.

**Clinical trials of anti-VEGF medications for DME**

The use of anti-VEGF medication in diabetes was first established in macular edema trials. During the studies, investigators noted that eyes randomized to the anti-VEGF group were more likely to have regression of diabetic retinopathy than in the control group. The use of anti-VEGF therapy was first established in the Macugen Diabetic Retinopathy Study Group included 16 eyes with PDR, of which 13 received pegaptanib. The study noted that eight underwent regression of neovascularization. Once the medication was stopped, neovascularization progression resumed. Bevacizumab (Avastin; Genentech, San Francisco, CA, USA) is a full-length recombinant humanized anti-VEGF monoclonal antibody (IgG) initially Food and Drug Administration (FDA) approved for treatment of metastatic colorectal cancers. It has twice the half-life of ranibizumab; Ophthalmic uses of bevacizumab are not FDA approved; however, its safety and efficacy have been shown in multiple neovascular age-related macular degeneration trials and DME trials, such as DRCR.net protocol T. Protocol T compared mean improvement in VA after intravitreal aflibercept, ranibizumab, or bevacizumab treatment in DME. It found that in eyes with VA of 20/40 or better, there was no significant difference in mean improvement in VA among bevacizumab (7.5±7.4 Early Treatment Diabetic Retinopathy Study (ETDRS) letters), ranibizumab (8.3±6.8), or aflibercept (8.0±7.6). In eyes with vision 20/50 or worse, aflibercept (18.9±11.5 letters) had better outcomes than bevacizumab (11.8±12) or ranibizumab (14.2±10.6) at 1-year follow-up. This difference continued to be significant between aflibercept and bevacizumab at 2-year follow-up. No significant difference in serious adverse reactions, major cardiovascular events, or hospitalization among the three medications was found. Ranibizumab (Lucentis; Genentech, San Francisco, CA, USA/Novartis Ophthalmics, Basel, Switzerland) is a recombinant antibody fragment of the humanized anti-VEGF monoclonal antibody. Based on bevacizumab, ranibizumab underwent affinity maturation to increase binding affinity to all isoforms of VEGF. Its half-life is shorter than other anti-VEGF molecules due to the lack of the antibody Fc domain. The RISE and RIDE studies, discussed later, demonstrated efficacy of ranibizumab in DME. Based on multiple trials, the FDA approved ranibizumab in 2017 for all stages of diabetic retinopathy.

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The VIVID/VISTA studies

The effect of anti-VEGF medications on diabetic retinopathy severity was also noted in aflibercept studies. The VIVID and VISTA trials compared intravitreal aflibercept, 2 mg, every 4 or 8 weeks with macular laser for DME.\textsuperscript{33} The primary outcome was change in mean best corrected visual acuity (BCVA) over 52 weeks. Mean BCVA gain from baseline in VISTA was 12.5 letters in aflibercept every 4 weeks, and 10.7 in aflibercept every 8 weeks, compared to 0.2 letters with laser (\textit{p}<0.0001). VIVID had similar results with greater VA gains in aflibercept-treated eyes (10.5 letters, every 4 weeks; 10.7 letters, every 8 weeks) when compared to laser (1.2 letters, \textit{p}<0.0001). A larger proportion of aflibercept-treated eyes also gained >15 or more ETDRS letters (\textit{p}<0.0001). A greater proportion of aflibercept-treated eyes were likely to experience a two-step or greater regression in DRSS in VISTA (33.8% every 4 weeks and 29.1% every 8 weeks versus 7.5% with laser, \textit{p}<0.01) and VIVID (33.3% every 4 weeks and 27.7% every 8 weeks versus 7.5%, \textit{p}<0.001). These results were maintained through 148 weeks of follow-up.

Protocol T comparison of bevacizumab, ranibizumab, and aflibercept

Comparison studies of anti-VEGF also showed that diabetic retinopathy regression was observed after treatment with all three anti-VEGF therapies. Protocol T was a comparison of bevacizumab, ranibizumab, and aflibercept for DME.\textsuperscript{29} A total of 660 patients (423 eyes with NPDR and 93 eyes with PDR) were randomized in a 1:1:1 fashion to intravitreal injection every 4 weeks for the first 6 months, then treatment following a predefined algorithm. Focal macular laser could be performed at 24 weeks for persistent DME. Eyes were followed for 2 years, and macular edema results were previously discussed.

Table 1. Percentage of eyes with ≥ 2 levels of DRSS improvement in all eyes and highest-risk NPDR eyes at 3 and 12 months of the RISE/RIDE study. A significantly high proportion of eyes treated with ranibizumab had improvement of diabetic retinopathy by two levels or more by the DRSS scale than with sham injection. This was true for all eyes with diabetic retinopathy (\textit{p}<0.001), but more so for the subset of eyes with the highest-risk NPDR at baseline. Thirty-two percent (32%) showed two or more levels of improvement after 3 months of treatment and 76% after 12 months of treatment, significantly higher than proportion of eyes receiving sham injection (0% at 3 months, 2% at 12 months, \textit{p}<0.001).

| Month | Sham injection | Ranibizumab, 0.3 mg | \textit{p}-value |
|-------|----------------|---------------------|-----------------|
|       | All eyes       |                     |                 |
| Month 3 | 0%             | 18%                 | \textit{p}<0.001|
| Highest-risk NPDR | 3%             | 32%                 | \textit{p}<0.001|
| Month 12 | 0%             | 18%                  | \textit{p}<0.001|
| All eyes | 3%             | 76%                 | \textit{p}<0.001|
| Highest-risk NPDR | 2%             |                     |                 |

DRSS, diabetic retinopathy severity scale; NPDR, nonproliferative diabetic retinopathy.
Post hoc analysis was performed to determine rate of diabetic retinopathy regression in the three treatment groups. In NPDR eyes, 31.2% of aflibercept-treated eyes, 37.7% of ranibizumab-treated eyes, and 22.1% of bevacizumab-treated eyes demonstrated improvement in severity of diabetic retinopathy. Aflibercept (p=0.004) and ranibizumab (p=0.01) were more strongly associated with diabetic retinopathy regression. At 2 years, there was no significant difference among the three groups (aflibercept, 24.8%; ranibizumab 31%; bevacizumab 22.1%). More improvement (51.9 and 64.6%) was seen with more severe forms of NPDR (severity levels 47 and 53).

In eyes with PDR, a significantly higher percentage of eyes demonstrated improvement in diabetic retinopathy after aflibercept treatment (75.9%) when compared to ranibizumab (55.2%, p=0.02) or bevacizumab (31.4%, p<0.001). The difference between ranibizumab and bevacizumab was nonsignificant (p=0.09). Seventy percent (70%) of eyes showed sustained improvement into year 2. Among eyes that did not improve at year 1, 21.4% did experience improvement by year 2 (aflibercept: 66.7%, bevacizumab: 13%, and ranibizumab: 15.4%), suggesting continued treatment despite initial nonresponse may be beneficial.

The proportion of eyes with worsening diabetic retinopathy was not significantly different among the three groups in NPDR or PDR eyes. At year 2, worsening NPDR was seen in 10.2% of eyes in the aflibercept group, 10.2% in the bevacizumab group, and 7.1% in the ranibizumab group. Two-point-four percent of eyes (2.4%) progressed to PDR. In eyes with PDR, worsening retinopathy was seen in 17.2% of aflibercept-treated eyes, 26.4% in the bevacizumab group, and 17.6% in the ranibizumab group. Vitreous hemorrhage was the most common indication of NPDR and PDR progression.

DME studies show that anti-VEGF treatment can promote regression of diabetic retinopathy. Anti-VEGF therapy has a greater effect in severe NPDR or PDR than mild or moderate NPDR. Aflibercept may be more effective at promoting regression, but the difference diminishes with time. Despite active treatment, diabetic retinopathy continues to worsen in a subset of eyes, especially in those with PDR.

Clinical trials of anti-VEGF medications in PDR

Anti-VEGF is frequently used as an adjunct to PRP in PDR. In eyes with persistent, active neovascularization, despite repeated PRP treatments, anti-VEGF medications were shown to significantly reduce the area of leakage and improve VA. In treatment-naïve PDR, eyes receiving a combination of PRP with anti-VEGF treatment had a more rapid and larger reduction in area of active neovascularization than PRP alone. Figuiera et al. studied 87 patients with high-risk PDR and found similar results. However, these studies did not address whether anti-VEGF monotherapy is sufficient for PDR management.

DRCR Protocol S: intravitreal ranibizumab compared with PRP for PDR

DRCR Protocol S was a randomized clinical trial designed to determine whether ranibizumab, 0.5 mg, was noninferior to PRP for PDR. Three hundred and ninety-four (394) treatment-naïve eyes with PDR were randomized 1:1 to treatment with intravitreal ranibizumab or PRP. Primary outcome was mean VA improvement at 2 years. Ranibizumab was injected every 4 weeks through week 12, after which further injections were administered based on a prespecified algorithm. Injection at weeks 16 and 20 were required unless all neovascularization had regressed. The PRP group received full laser treatment over 1–3 sessions and was allowed to receive anti-VEGF medication if DME was present.

The average change in VA at 2 years was +2.8 letters in the ranibizumab group, compared to +0.2 letters in the PRP group (p=0.001 for noninferiority). There was no difference in percentage of eyes with quiescent PDR (PRP: 30%, ranibizumab: 35%). Although 98% of eyes in the PRP group received the per-protocol initial PRP treatment, 45% of eyes needed supplemental PRP for PDR progression. Only 6% of eyes in the ranibizumab group needed rescue PRP treatment. Of 155 eyes without baseline DME and received PRP, 62 received ranibizumab over 2 years of follow-up. The cumulative 2-year probability of developing visually significant central DME was 9% in the ranibizumab group compared with 28% in the PRP group.

Risk factors for progression of PDR were identified through a post hoc analysis. A composite outcome was used to determine worsening PDR, including vitreous hemorrhage, neovascularization of the iris or angle, neovascular glaucoma, retinal detachment, as well as need for further PRP or vitrectomy. Overall, the PRP group had a higher proportion (42 versus 34%, p=0.063) of outcomes associated with PDR progression. When the analysis was adjusted for severity of PDR, there was a significantly higher proportion of outcomes associated with PDR progression in the PRP group, when compared to the ranibizumab group (HR 1.45, p=0.024). This became more apparent when assessing eyes with center-involving DME that were randomized to PRP. These eyes did not receive adjunct ranibizumab and had a higher hazard ratio for adverse outcomes (HR 1.73, p=0.004) compared to ranibizumab alone.

The most significant risk factor for development of poor outcomes in PRP and ranibizumab treatment groups was severity of PDR based on ETDRS severity scale. Additional risk factors for worsening PDR included: presence of epiretinal membrane, neovascularization of the disk with neovascularization elsewhere, and presence of vitreous hemorrhage. In the PRP group, treatment with pattern scan PRP was associated with worse outcomes compared to single-spot PRP (60 versus 39%, p<0.008). Vitreous hemorrhage was the most common adverse event.

In Protocol S, ranibizumab-treated eyes had better VA outcomes (+2.8 versus 0.2 letters), less development of DME (PRP: 28%, ranibizumab: 9%, p<0.001), and less need for vitrectomy (PRP: 15%, ranibizumab: 4%, p<0.001). They
were less likely to experience events associated with PDR progression than eyes treated with PRP, especially in PDR with high-risk characteristics. The authors emphasized close follow-up of both PRP- and ranibizumab-treated eyes, because 42% of PRP-treated eyes and 34% of ranibizumab-treated eyes demonstrated progression of PDR as defined by the post hoc composite outcome. The most common adverse outcome was vitreous hemorrhage, which was also the most common reason for vitrectomy. PRP-treated eyes required more vitrectomy and additional PRP sessions.

**CLARITY: intravitreal aflibercept compared with PRP for PDR**

CLARITY was a phase 2b, single-blind, noninferiority trial comparing intravitreal aflibercept to PRP in newly diagnosed or previously laser-treated active PDR. Patients were treated with three-monthly injections of aflibercept (2 mg/0.05 mL) and reassessed every 4 weeks thereafter for further injections as needed, or treated with standard PRP and reassessed every 8 weeks for further PRP. Primary outcome was change in BCVA at 12 weeks. A total of 392 patients (191 untreated and 201 previously treated) underwent randomization. Aflibercept was found to be noninferior to PRP with a difference of +3.9 ETDRS letters (95% CI: 2.3–5.6) favoring aflibercept at 2 years. Regression of retinal neovascularization at 2 years was seen in 81% of aflibercept-treated eyes and 78% of PRP-treated eyes. Sixty-four (64%) of aflibercept-treated eyes had total regression of neovascularization, compared to 34% in PRP eyes. Accordingly, the aflibercept group had greater improvement in retinopathy as graded by ETDRS retinopathy severity score (p=0.016).

Eyes receiving PRP were more likely to develop new or increasing vitreous hemorrhage (18 versus 9%, p=0.034) and more likely to need vitrectomy, though this was not statistically significant (6 versus 1%, p=0.066). Aflibercept-treated eyes appeared to have more inflammation (8 versus 3%, p=0.075) and corneal-related problems such as abrasion, epithelial erosions, and conjunctival lacerations (4 versus 0%, p=0.060). Patients with baseline macular edema were excluded. At 2 years, proportion of macular edema was 11% in the aflibercept group and 29% in the PRP group.

**Anti-VEGF therapy for vitreous hemorrhage associated with PDR**

In eyes with vitreous hemorrhage due to PDR, PRP is not possible until the blood clears or vitrectomy is performed. Anti-VEGF therapy can instead be used to treat the underlying PDR, but it may not reduce the need for vitrectomy to remove nonclearing vitreous hemorrhage. The DRCR network Protocol N studied rate of vitrectomy after intravitreal anti-VEGF for vitreous hemorrhage. Two-hundred sixty-one (261) eyes with PDR and vitreous hemorrhage that prevented PRP treatment were randomized to intravitreal ranibizumab, 0.5 mg, compared with intravitreal saline at baseline, week 4, and week 8. No difference was found in rate of vitrectomy by week 16 (ranibizumab: 12%; saline: 17%, p=0.37). Due to low overall rate of vitrectomies, the study was underpowered to detect a difference in vitrectomy rates.

The ranibizumab group was more likely to complete PRP without need for vitrectomy (44 versus 31%, p=0.05), and VA outcomes also favored the ranibizumab group (22±23 letters) compared to the saline group (16±31 letters, p=0.04). However, there was no difference in percentage of eyes with poor (20/200 or worse) or very poor (20/800 or worse) VA between the groups. Twenty percent (20%) of ranibizumab-treated eyes had poor VA compared to 27% of saline-treated eyes, and 11% of ranibizumab-treated eyes had very poor VA compared to 16% of saline-treated eyes.

**Side effects of anti-VEGF medications**

There are limitations and risks to anti-VEGF use. Common side effects include transient increase in intraocular pressure and floaters. Operator complications including traumatic intraocular injuries, such as to the lens, have also been reported. Endophthalmitis is uncommon, with prevalence estimated at 1 per 1000 injections. Rare side effects include uveitis, which has been reported with aflibercept and bevacizumab. A serious vision-threatening complication of anti-VEGF use in PDR is the development of tractional retinal detachment in the setting of pre-existing membranes. It is hypothesized that VEGF-associated neovascularization and connective tissue growth factor (CTGF)-associated fibrosis balance each other in the PDR eye. When anti-VEGF medications are used, the balance favors CTGF and an ‘angio-fibrotic switch’ is flipped. CTGF promotes cell apoptosis, accelerates fibrosis, and produces traction on the underlying retina. This occurs more frequently in the first month of anti-VEGF treatment and in eyes with long-standing (>15 years), poorly controlled diabetic retinopathy. Other risk factors identified include vitreous hemorrhage, PDR resistant to PRP, and higher dose of intravitreal anti-VEGF.

Systemic side effects of anti-VEGF medications are also a concern. In cancer trials, high doses of anti-VEGF medications were associated with increase in bowel perforation, arterial thromboembolism, myocardial infarction (MI), stroke, and hypertension. These side effects have not been confirmed with the lower dose and intravitreal route of ophthalmic anti-VEGF use, though studies were not powered to detect systemic adverse effects. The RISE/RIDE studies found an increase in incidence of stroke in ranibizumab, 0.5 mg, compared with 0.3 mg dose (4.8 versus 2.0%), while the incidence of MI at 36 months was 7.2% in the 0.3 mg cohort and 3.6% in the 0.5 mg cohort. However, the study was not powered to detect a difference in MI incidence between the cohorts. In VIVID/VISTA, the overall nonocular side effects were similar between the three groups and related to underlying comorbidities. No difference was found among three anti-VEGF medications.

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in Protocol T. Protocol S also did not find a difference in serious adverse events, including hospitalization, death, Antiplatelet Trialists’ Collaboration arteriothromboembolic events, or MedDRA organ system class events between PRP or ranibizumab. Multiple meta-analyses of the data have both demonstrated and refuted increases in risk of death or cerebrovascular events.

**Discussion**

In summary, anti-VEGF medications are an important addition to the arsenal of PDR treatments. Studies in DME have demonstrated that anti-VEGF medications can promote regression of diabetic retinopathy in both NPDR and PDR, keeping in mind that PDR can still develop despite monthly treatments. The severity of the retinopathy is associated with a more robust response. Protocol T demonstrated that all three medications – aflibercept, ranibizumab, and bevacizumab – have a favorable effect in diabetic retinopathy. In NPDR, aflibercept had better outcomes in year 1, but no visually significant difference among the three groups was found by year 2. VIVID/VISTA results up to 148 weeks of treatment showed that with continued treatment, the effect on retinopathy severity is durable.

Anti-VEGF can be used as an adjunct to PRP and as primary treatment in proliferative retinopathy; Protocol S demonstrated that ranibizumab alone was noninferior to PRP. Outcomes favored ranibizumab, with improved VA gains (2.8 versus 0.2 letters), improvement of pre-existing DME, lower rates of DME development, and less need for rescue laser or vitrectomy. Additionally, in subgroup analysis, comparing those eyes with DME and PDR, the visual outcomes favored treatment with ranibizumab (median +10, range 0 to +16 letters) compared to PRP (median +5, range –4 to +13 letters). Post hoc analysis also favored ranibizumab treatment. Eyes receiving PRP were more likely to have signs of PDR progression, especially vitreous hemorrhage, than intravitreal ranibizumab treatment alone. Again, severity of disease was the strongest predictor of adverse outcomes. It is important to recognize that a large percentage of eyes in both groups had complications of active PDR despite continuous treatment. These results suggest that anti-VEGF can be started early in severe NPDR or PDR disease course, and aggressive treatment may be necessary in severe PDR.

In eyes with proliferative retinopathy and vitreous hemorrhage that preclude PRP, anti-VEGF treatment can be used to treat the underlying retinopathy and may delay but may not reduce need for surgery. DRCR Protocol H, though underpowered, did not show a significant difference in rates of vitrectomy at 16 weeks between saline injection or ranibizumab-injected eyes. Eyes that received ranibizumab did have better VA outcomes (+22±23 versus +16±31 ETDRS letters).

There are limitations to anti-VEGF therapies for diabetic retinopathy. PRP is cost-effective, can be completed in a few office visits, and treatment has long-lasting effects. In contrast, there is no known treatment period after which it is safe to discontinue anti-VEGF therapy. Studies have shown that once the anti-VEGF medication is stopped, neovascularization will reoccur or continue to progress. Additionally, while PRP targets ischemic retina and reduces the drive for VEGF, effects of anti-VEGF on retinal ischemia are unclear. Further analysis of DRCR Protocol T and S also remind us that despite treatment, approximately 30% of eyes with retinopathy will continue to worsen. In young patients with PDR, the need for close monitoring and the uncertainty of long-term outcomes with anti-VEGF treatment are especially of concern.

Major ocular adverse effects of anti-VEGF intravitreal injections, such as intraocular pressure rise, are transient but may be relevant in patients with glaucoma. Serious adverse effects such as endophthalmitis are rare but accumulate with continuous injections. In proliferative disease, there is also the risk of tractional retinal detachments after anti-VEGF use in eyes with pre-existing vitreoretinal membranes, which can lead to vision loss. The cost and time of anti-VEGF monotherapy is also a burden upon patients and the healthcare system. Many patients with diabetic retinopathy are of working age, suffer from poorly controlled diabetes with systemic manifestations, and face socioeconomic strains. For healthcare systems, it was estimated that the cost utility for PRP was 85% lower than intravitreal anti-VEGF therapy in facility settings. For each line of vision saved, the estimated cost of anti-VEGF therapy ($16,849) was more than double that of PRP ($725). The efficacy of the two treatments is similar. Protocol S did not show a significant difference between PRP and ranibizumab in overall percentage of eyes achieving quiescent PDR.

In light of these studies, the American Academy of Ophthalmology (AAO) Preferred Practice Pattern committee now states that there is sufficient evidence for the treatment of diabetic retinopathy with anti-VEGF treatment. The Preferred Practice Pattern states, “It would be reasonable to consider use of ranibizumab in severe NPDR patients in settings where laser surgery would be considered,” and in high-risk PDR, an anti-VEGF alternative to PRP could be considered for patients who can follow-up regularly, especially if there is macular edema. Anti-VEGF medications monotherapy can be considered as first-line therapy with PDR assuming judicious oversight by the clinician for disease progression. It also has a role in situations where PRP is not possible, due to media opacity, dense cataract, or nonclearing vitreous hemorrhage, where the patient is unable to go to the operating room.

The role of anti-VEGF treatment in NPDR is also evolving. The AAO Preferred Practice Pattern recommends considering anti-VEGF treatment in severe NPDR where PRP would otherwise be considered. Anti-VEGF treatment has also been shown to reduce diabetic retinopathy severity and is associated with lower rates of PDR development. Currently, the management of NPDR consists of risk factor modification. Use of anti-VEGF medications in addition to risk factor modification may prevent development of PDR. Ongoing studies are addressing the...
REVIEW – Anti-VEGF in the management of proliferative diabetic retinopathy

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