Vascular Endothelial Growth Factor Antagonists: Promising Players in the Treatment of Neovascular Age-Related Macular Degeneration

Abstract: Neovascular age-related macular degeneration (nAMD) treatment has been revolutionized by the introduction of vascular endothelial growth factor antagonists (anti-VEGF), but the need for frequent intravitreal injections poses a heavy burden to patients and physicians. Evolving anti-VEGF therapies include longer duration agents, approaches that target multiple pathways, topical anti-VEGF agents, sustained-release, and genetic therapies. Abicipar pegol, a designed ankyrin repeat protein (DARPin), demonstrated the ability to maintain stable visual acuity with 12-week dosing, but was not approved by the FDA due to higher than usual rates of intraocular inflammation. Conbercept, a recombinant anti-VEGF fusion protein, has been approved in China, and is in Phase 3 trials globally. KSI-301 is an anti-VEGF antibody biopolymer conjugate that allowed 66% of nAMD patients to maintain at least a 6-month treatment-free interval in Phase 1b studies. OPT-302, an inhibitor of VEGF-C/D, will be tested in phase 3 studies that compare anti-VEGF-A monotherapy against combination therapy with OPT-302. Faricimab is a bispecific anti-VEGF/Ang-2 antibody that upregulates the Tie-2 signaling pathway and promotes vascular stability; it is undergoing phase 3 trials with potential for 12- or 16-week dosing. PAN-90806 is a topical anti-VEGF agent that showed the ability to reduce injection frequency by 79% compared to ranibizumab monotherapy in a phase 1/2a trial. Sustained-release anti-VEGF therapies include the ranibizumab Port Delivery System (in phase 3 studies), GB-102 (Phase 2b), OTX-TKI (phase 1), and Duraset (preclinical). Suprachoroidal delivery of the tyrosine kinase inhibitor, axitinib, is in preclinical studies. Genetic therapies in phase 1 studies include RGX-314 and ADVM-022, which introduce a viral vector that modifies the retina’s cellular apparatus to create an anti-VEGF biofactory, potentially serving as a one-time treatment. Further investigation is warranted for drugs and delivery systems that hope to advance visual outcomes and reduce treatment burden of nAMD.

Keywords: ADVM-022, age-related macular degeneration, abicipar pegol, CLS-AX, conbercept, faricimab, GB-102, KSI-301, PAN-90806, OTX-TKI, ranibizumab port delivery system, RGX-314, VEGF

Background

Age-related macular degeneration (AMD) is the most common cause of irreversible blindness in the elderly population of the industrialized world. The number of patients globally with AMD is estimated to be 196 million in 2020 and will reach 288 million in 2040. AMD is classified into one of two main subtypes: the non-neovascular (dry) and neovascular (wet) forms of the disease. Approximately 90% of all AMD case are characterized as dry AMD, in which accumulation of yellow...
lipofuscin-rich deposits known as drusen, along with atrophy of the macula, result in gradual vision loss over years.\textsuperscript{3} Neovascular AMD (nAMD), on the other hand, may result in acute and substantial loss of central vision; the growth of pathologic choroidal neovascularization (CNV) under the macula results in exudation of blood and/or serous fluid into the macula.\textsuperscript{4,5} The neovascular tissue and associated exudates can progress into submacular fibrosis that causes permanent loss of central vision.

Irregular neovascularization and exudation are caused by increased levels of the pro-angiogenic cytokine, vascular endothelial growth factor-A (VEGF).\textsuperscript{6} This cytokine was originally called vascular permeability factor because it results in increased exudation from blood vessels.\textsuperscript{7} Intravitreally administered therapies that neutralize VEGF have substantially reduced the degree of vision loss for patients with nAMD compared to the previous standard of care.\textsuperscript{8}

### Existing Treatments

Before the introduction of intravitreal anti-VEGF therapy, treatment options for nAMD included photodynamic therapy with verteporfin, focal laser photocoagulation, intravitreal corticosteroids, and surgical removal of CNV. These treatments have fallen out of favor since they did not significantly improve visual outcomes.\textsuperscript{9} In 2004, pegaptanib (Macugen, Eyetech) was approved by the United States Food and Drug Administration (US FDA) for the treatment of nAMD, marking the beginning of the anti-VEGF era; more recently it has also fallen out of favor compared to newer and more effective anti-VEGF agents such as ranibizumab, aflibercept, bevacizumab, and brolucizumab (summarized in Table 1). Most recently, in 2019, Brolucizumab (Novartis, Basel, Switzerland) was approved for 12-week dosing intervals in nAMD,\textsuperscript{10} it has recently come under scrutiny due to reports of occlusive retinal vasculitis associated with its use.\textsuperscript{11} All of these agents inhibit VEGF-A, which binds VEGF receptor (VEGFR)-2, that mediates angiogenesis and vascular leakage.

### Unmet Needs and Current Research Goals

Despite the substantial progress in nAMD treatments, there are a number of shortcomings to anti-VEGF therapy, including the necessity of repeated injections and inadequate anatomic response in a subset of eyes. Even with monthly repeated injections of anti-VEGF therapy over 12 months, one fifth of patients have declining visual acuity (VA), about half do not achieve 20/40 VA (the requirement for an unrestricted driver’s license in parts of the USA and in most European Union countries), and over 2/3 fail to gain 3 or more lines of ETDRS letters.\textsuperscript{8,12}

Although favorable visual outcomes with treat and extend dosing of aflibercept have been reported,\textsuperscript{13} many “real world” anti-VEGF injection studies show visual outcomes inferior to those reported in randomized clinical trials (RCTs), likely as a result of non-compliance and under-treatment in the real world.\textsuperscript{14–29} Many reports have documented a positive correlation between the quantity of

### Table 1 Summary of Currently Available Intravitreal Anti-VEGF Therapies

| Anti-VEGF Agents | Description | Approval Date/Indication |
|------------------|-------------|--------------------------|
| Macugen (pegaptanib) | Pegylated synthetic RNA-based oligonucleotide targeting VEGF\textsubscript{165} | December 17, 2004/nAMD |
| Avastin (bevacizumab) | Recombinant humanized monoclonal IgG1 antibody against VEGF | Approved on February 24, 2004 for metastatic colorectal cancer; used off-label for nAMD |
| Lucentis (ranibizumab) | Humanized IgG1 antibody fragment against VEGF | June 30, 2006/nAMD |
| Eylea (aflibercept) | VEGF-trap, a recombinant protein created by fusing the second Ig domain of human VEGF receptor 1 with the third domain of human VEGF receptor 2, which in turn is fused to the constant region of human IgG1 | November 18, 2011/nAMD |
| Beovu (brolucizumab) | Humanized single-chain antibody fragment against VEGF | October 8, 2019/nAMD |

**Abbreviations:** RNA, ribonucleic acid; VEGF, vascular endothelial growth factor; nAMD, neovascular age-related macular degeneration; IgG1, immunoglobulin G1.
anti-VEGF injections over 1 year and better visual outcomes.\textsuperscript{15–17,21,24,30–32}

Compliance with frequent anti-VEGF injection can be limited by factors such as patients’ anxiety toward the needle, financial hardship, scheduling conflicts, and lack of transportation. More durable anti-VEGF formulations may help overcome some of these barriers. Although anti-VEGF drugs improve VA by limiting leakage from CNV, the pathophysiology of nAMD is indeed a complex process requiring attention to many more pathologic mechanisms than just VEGF inhibition. Unfortunately, a large RCT showed that nearly half of anti-VEGF treated eyes developed submacular fibrosis despite 2 years of injections, which limits further visual benefit despite treatment.\textsuperscript{33} As a result, new therapies and delivery systems are being developed to target other complementary mediators involved in angiogenesis.

In particular, inhibition of VEGF-A has been shown to upregulate other forms of VEGF, which may contribute to limited outcomes. For example, after bevacizumab (anti-VEGF-A) administration in nAMD patients, VEGF-C which also activates the VEGF2 receptor is upregulated.\textsuperscript{34} This upregulation of other VEGF ligands could contribute to treatment resistance, and a ceiling of efficacy. Tyrosine kinase inhibitors (TKIs), which achieve pan-VEGF receptor inhibition, could address this issue and, consequently, there has been interest in TKIs as investigational agents. TKIs assessed in clinical trials include oral vorolanib, topical sorafenib, regorafenib, and pazopanib, as well as intraocularly-injected sunitinib and axitinib. Although a few investigational anti-platelet-derived growth factor (anti-PDGF) drugs were developed with the objective of combining them with anti-VEGF injections, none have shown advantage over anti-VEGF monotherapy in a phase 3 clinical trial.\textsuperscript{35,36} The Tie-2 receptor kinase, which enhances vascular stability to minimize leakage, has been another target for investigational therapies.\textsuperscript{37} Sustained-release drug delivery systems, along with topical and oral anti-VEGF formulations, may also help to reduce injection frequency in the future. Viral vector-based gene therapy can produce continuous anti-VEGF expression after a single treatment.\textsuperscript{38}

**Drugs and Delivery Systems in Development**

See Table 2 for a summary of investigational anti-VEGF medications and delivery systems.

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**Abicipar Pegol**

Abicipar pegol is a novel anti-VEGF drug based on DARPin® (Designed Ankyrin Repeat Protein) technology. Two phase 3 clinical trials (SEQUOIA and CEDAR) sponsored by Allergan (Dublin, Ireland) and Molecular Partners (Zurich, Switzerland) compared abicipar against ranibizumab.\textsuperscript{39} Both studies were comprised of three treatment groups: abicipar every 8 weeks, abicipar every 12 weeks, and monthly ranibizumab. The 8-week group was treated with three loading doses and the 12-week abicipar group with two loading doses before switching to their specified treatment interval. The proportion of treated patients with stable vision (<15 ETDRS letters lost) at 1 year was the primary endpoint.

In SEQUOIA, 94.8% and 91.3% of those treated with abicipar every 8 and 12 weeks maintained stable vision. In the group that received monthly ranibizumab, 96% of patients maintained stability. In CEDAR, 91.7% and 91.2% of patients in the abicipar groups maintained stable vision when dosed every 8 and 12 weeks, compared with 95.5% of patients in the monthly ranibizumab group.\textsuperscript{39}

While the durability of abicipar was impressive, the rates of intraocular inflammation were significantly higher in the abicipar groups (15.1–15.7%) compared to the ranibizumab group (0–0.6%). A newer modified version was studied in the smaller MAPLE study. In this open-label study, 123 nAMD patients received three monthly 2 mg abicipar injections followed by 2 mg injections every 8 weeks through week 28. The incidence of intraocular inflammation was 8.9%, mostly mild-to-moderate in severity. The incidence of severe inflammation was 1.6%, with one case ofiritis and one case ofuveitis. There were no reported cases ofendophthalmitis or retinal vasculitis.\textsuperscript{39} Despite this, in June 2020 the FDA rejected abicipar as a nAMD drug candidate, citing an unfavorable benefit–risk ratio of intraocular inflammation compared to the current treatment options for nAMD. Allergan/AbbVie are to discuss further steps with the FDA.\textsuperscript{40}

**OPT-302**

OPT-302 (Opthea; Victoria, Australia) inhibits both VEGF-C and VEGF-D, which are thought to serve a supplemental role to VEGF-A in nAMD pathophysiology. As discussed above, broad VEGF inhibition has the potential to address secondary upregulation of other forms of VEGF that follows VEGF-A blockade. The phase 1 trial (NCT02543229) demonstrated that monthly intravitreal...
injections of combination OPT-302 2 mg and ranibizumab 0.5 mg (to inhibit VEGF-A) had a favorable safety profile. After 12 weeks of follow-up, this regimen showed significant extra visual gains and reduction in CST compared to ranibizumab monotherapy. Of 51 patients enrolled, 25 were treatment-naive patients, and 26 had been previously treated with intravitreal anti-VEGF-A inhibitors.41 Seven of 13 (54%) patients receiving OPT-302 monotherapy did not need rescue anti-VEGF-A injections and the mean change in BCVA from baseline to week 12 was +5.6 letters. Mean BCVA gains from baseline to week 12 following combination OPT-302 with ranibizumab were +10.8 letters in treatment-naive patients and +4.9 letters in patients with prior anti-VEGF treatment. At week 12, change in mean CST in both groups were −119 μm and −54 μm, respectively; half of treatment-naive patients did not show any detectable CNV at that point. The proportion gaining ≥5, ≥10, or ≥15 ETDRS letters at week 12 was 67%, 44%, and 33%, respectively, in treatment-naive subjects (baseline mean BCVA of 56.4 letters); for those patients with a history of prior anti-VEGF therapy it was 53%, 16%, and 0% (baseline mean BCVA was higher at 64.5 letters).41

Phase 2b trials compared ranibizumab monotherapy with two doses of combination OPT-302/ranibizumab in 366 treatment-naive patients. Patients receiving 2 mg OPT-302 combination therapy gained a mean of 14.2 letters by week 24, compared to 10.8 letters in ranibizumab group (difference of 3.4 letters, p=0.0107). The 0.5 mg OPT-302 low dose group had a similar outcome to the ranibizumab group (+9.4 letters). High dose OPT-302 (2 mg) combination treatment showed improvements across multiple secondary endpoints, including a higher proportion of patients with stable vision (defined as ≤15 letter loss from baseline), and those gaining ≥10 and ≥15 letters of visual acuity.42

Opthea has two upcoming phase 3 trials, ShORe and COAST, which will enroll treatment-naive patients and

### Table 2 Summary of Emerging Drugs to Treat Neovascular Age-Related Macular Degeneration (nAMD)

| Compound        | Company                  | Stage of Development                           | Structure/Mechanism of Action                                      |
|-----------------|--------------------------|-----------------------------------------------|-------------------------------------------------------------------|
| Abicipar pegol  | Allergan                 | Phase 3 completed – FDA did not approve        | DARPin antagonist of VEGF-A                                        |
| Faricimab       | Roche                    | Phase 3 ongoing                               | Bispecific Ang-2/VEGF-A antibody                                  |
| Ranibizumab PDS| Genentech                | Phase 3 ongoing                               | Refillable port of VEGF-A mAb                                     |
| Conbercept      | Chengdu Kanghong Biotech Co., Ltd. | Phase 3 terminated in April 2021 | Anti-VEGFR1/2                                                    |
| OPT-302         | Ophthea                  | Phase 3 starting 2021                         | Anti-VEGF-C/VEGF-D                                               |
| GB-102          | GrayBug Vision           | Phase 2b ongoing                              | Bioerodible nanoparticles encapsulate TKI of VEGFR/PDGFR        |
| ADVM-022/ADVM-032 | Adverum Biotechnologies | Phase 2 enrolling in late 2021               | AAV encoding anti-VEGF-A similar to aflibercept                 |
| RGX-314         | Regenxbio                | Phase 2b/3 expected to begin 2021; phase 2 enrolling for suprachoroidal injection | AAV8 encoding anti-VEGF-A similar to ranibizumab                |
| KSI-301         | Kodiak Sciences          | Phase 1b ongoing; phase 2b recruitment completed | Anti-VEGF antibody biopolymer conjugate                            |
| PAN-90806       | PanOptica                | Phase 1/2 completed                           | TKI of VEGF-A/PDGF                                               |
| OTX-TKI         | Ocular Therapeutix       | Phase 1 ongoing                               | Sustained release anti-VEGF-A                                    |
| Durasept        | pSivida                  | Preclinical                                   | TKI of VEGF-A/PDGF                                               |
| ARP-1536        | Aerpio                   | Preclinical                                   | Tie-2 receptor activation (via VE-PTP inhibition)                 |
| CLS-AX (axitinib)| Clearside Biomedical     | Preclinical                                   | Suprachoroidal injection of TKI                                   |

**Abbreviations:** MW, molecular weight; Fab, fragment antibody; VEGF, vascular endothelial growth factor; DARPin, Designed Ankyrin Repeat Protein; PDGF, platelet-derived growth factor; TKI, tyrosine kinase inhibitor; Ang-2, Angiopoietin-2; mAb, monoclonal antibody; VE-PTP, vascular endothelial-protein tyrosine phosphatase; VEGFR, vascular endothelial growth factor receptor; AAV, adeno-associated virus; cDNA, complementary deoxyribonucleic acid.

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assess the efficacy and safety of 2 mg OPT-302 in combination with anti-VEGF-A therapy compared to anti-VEGF-A monotherapy. Patients will be dosed every 4 or 8 weeks (treated with ranibizumab for ShORe and aflibercept for COAST). The primary endpoint of both trials will be the mean change in VA from baseline to week 52, though patients will continue to be dosed until week 96 to evaluate long-term safety. Opthea plans to commence both trials in the first half of 2021.43

Conbercept
Conbercept (Chengdu Kanghong Biotech Co. Ltd.; Chengdu, Sichuan, China) is a recombinant anti-VEGF fusion protein that is composed of portions of VEGFR1 and VEGFR2. The phase 3 PHOENIX trial conducted in China led to its approval there. One treatment arm received conbercept 0.5 mg monthly for 3 months followed by 0.5 mg every 3 months. The delayed treatment group received three monthly sham injections, followed by three monthly conbercept 0.5 mg injections, and finally quarterly conbercept 0.5 mg injections until month 12. The group with the loading doses of conbercept demonstrated an improvement of 9.9 letters, while the delayed treatment arm had 8.8 letter improvement, both of which were statistically significant.44

PANDA-1 (NCT03577899) and PANDA-2 (NCT03630952) are both international phase 3 studies that evaluated safety and efficacy in comparing conbercept 0.5 mg and 1.0 mg to aflibercept. Patients in each of the trials’ three study arms received three monthly loading doses, after which the treatment groups formed were: conbercept 0.5 mg at 8-week intervals, conbercept 1 mg at 12-week intervals, and aflibercept 2 mg at 8-week intervals. The primary outcome was mean change in BCVA at 36 weeks. The PANDA-1 and PANDA-2 trials with a combined enrollment target of 1,140 patients were originally targeted for completion in 2022.45,46 Due to disruptions related to the global pandemic of 2020–2021, in which many patients did not complete adequate follow-up to obtain results with registrational value, the trials were terminated early in April 2021.47

KSI-301
KSI-301 (Kodiak Sciences; Palo Alto, CA) has developed a novel anti-VEGF antibody biopolymer conjugate (ABC platform48) for the treatment of nAMD and other retinal vascular diseases. The ABC platform comprises a humanized IgG1 antibody with inert immune effector function and a biopolymer that is an optically clear high molecular weight phosphorylcholine polymer, which was designed to increase intraocular durability. The phase 1b study for nAMD compares 8-week dosing of aflibercept with 12, 16, and 20-week dosing of KSI-301. Patients were randomized to receive a 2.5 mg/50 µL or 5 mg/100 µL dose with three loading doses at monthly intervals and a mandatory re-treatment at 6 months after the last treatment.48

New KSI-301 data were presented at the Angiogenesis, Exudation, and Degeneration 2021 virtual meeting in February 2021. This data demonstrated promise in 1-year durability, efficacy, and safety in the Phase 1b study. Of nAMD patients, 66% achieved a 6-month or longer treatment-free interval at the 1-year mark and 78% were on a 4-month or longer interval. An average of only 2.0 retreatments were given in the 10 months after the initial three loading doses. In nAMD patients a 5.7 letter improvement was noted after 1 year.49

There has also been recent completion of recruitment in the USA for Kodiak’s DAZZLE Phase 2b/3 study (NCT04049266) in treatment naive nAMD patients. In this study, after three monthly loading doses patients are randomized to receive KSI-301 on a dosing regimen no more than every 3 months and as infrequently as every 5 months compared to aflibercept every 8 weeks. BCVA at 1 year will mark primary study outcome, with continued following of patients for 2 more years.50

Anti-VEGF-A Plus Tie-2 Receptor Modulator Combination Therapy
The Tie-2 tyrosine kinase receptor, which is expressed on vascular endothelial cells, has also become a target of interest in treating exudative retinal disease. Activation of this receptor reinforces junctional proteins and stabilizes vasculature to limit permeability.51 Angiopoietin-1 (Ang-1) is a ligand that activates the Tie-2 receptor, thus reducing vascular leakage and promoting homeostasis.52 Angiopoietin-2 (Ang-2) is upregulated in pathologic states and is a competitive antagonist to Ang-1; elevated Ang-2 levels promote vascular leakage.53 Several new drugs have been designed to inhibit Ang-2, thus allowing the constitutively secreted Ang-1 to activate the Tie-2 receptor. Another approach targets the enzyme, vascular endothelial protein tyrosine phosphatase (VE-PTP), which inactivates the Tie-2 receptor and more directly affects the phosphorylation and activation of Tie-2.54 Enhancing Tie-2 receptor
function in combination with anti-VEGF therapy may provide enhanced efficacy over anti-VEGF monotherapy.

**Faricimab**

Faricimab (Roche/Genentech; Basel, Switzerland) is a unique bispecific antibody targeting both VEGF-A and Ang-2. The phase 2 AVENUE trial included 273 patients, which compared treatment with ranibizumab 0.5 mg every 4 weeks, faricimab 1.5 mg every 4 weeks, faricimab 6 mg every 4 weeks, faricimab 6 mg every 8 weeks, and faricimab 6 mg/ranibizumab 0.5 mg combination therapy (three ranibizumab injections every 4 weeks, followed by faricimab every 4 weeks) in treatment-naïve nAMD. The primary outcome was mean change in BCVA from baseline at 36 weeks. The 1.5 mg faricimab arm given every 4 weeks demonstrated the best gains of +9.1 letters at 36 weeks. All arms showed significant decreases in central subfield thickness (CST), with the combination therapy arm demonstrating the largest reduction of −185 µm. 55

Another phase 2 trial, STAIRWAY, compared faricimab 6 mg given either every 12 or 16 weeks to ranibizumab 0.5 mg every 4 weeks in 76 treatment-naïve nAMD patients. In the two faricimab groups, patients initially received four monthly faricimab injections followed by injections either every 12 or 16 weeks. The ranibizumab group received monthly treatment. At week 24, 65% of all patients treated with faricimab had no protocol-defined activity 12 weeks after their last injection. At 52 weeks, patients in the 12-week group gained a mean of 10.1 letters, those in the 16-week group gained 11.4 letters, and the ranibizumab group gained 9.6 letters. 56

Two multi-center Phase 3 trials, LUCERNE (NCT03823300) and TENAYA (NCT03823287), have been launched with 1,280 participants randomized into treatment arms of faricimab every 16 weeks (with the ability to decrease the interval to 12 or 8 weeks) or aflibercept every 8 weeks with the average change of BCVA from baseline to week 48 serving as the primary outcome measure. The expected completion of the Phase 3 trials is Fall 2022. Interim results released in February 2021 revealed that both studies met their primary endpoint, with faricimab demonstrating non-inferior VA gains to aflibercept. In TENAYA and LUCERNE, the mean vision gains from baseline in the faricimab arms were +5.8 and +6.6 letters, respectively, compared to +5.1 and +6.6 letters in the aflibercept arms. Of note, 45.7% (144/315) of patients in TENAYA and 44.9% (142/316) in LUCERNE were able to be treated every 4 months in the first year. An additional 34% (n=107/315) of patients in TENAYA and 32.9% (n=104/316) in LUCERNE were able to be treated every 3 months. Combined, approximately 80% of faricimab-treated patients were able to go 3 months or longer between treatments during the first year. In both studies, faricimab given at intervals of up to 4 months offered reductions in CST similar to aflibercept given every 2 months. 57

**ARP-1536**

ARP-1536 (Aerpio Therapeutics; Blue Ash, OH) is a monoclonal antibody that activates the Tie-2 receptor by inhibiting VE-PTP, which is the most downstream negative regulator of Tie-2. ARP-1536 is administered by intravitreal injection and binds to the extracellular domain of VE-PTP, hindering its function, and thus causing increased activation of the Tie-2 receptor. Pre-clinical studies have established its biologic activity. AKB-9778, Aerpio’s other VE-PTP inhibitor, is administered by subcutaneous injection. 37 At the time of this writing, Aerpio has not made any public updates on plans for a phase 1 clinical trial for ARP-1536.

**Topical Treatments**

There have been several topical therapies developed to inhibit VEGF and PDGF, including pazopanib (a TKI, GlaxoSmithKline; Brentford, UK), squalamine lactate (Ohr Pharmaceutical; New York, NY), regorafenib (Bayer Healthcare; Leverkusen, Germany), and LHA510 (a TKI, Alcon; Fort Worth, TX). None of these drugs were shown to reduce the need for intravitreal anti-VEGF injections, and their clinical development has been discontinued. 58

**PAN-90806**

PAN-90806 (PanOptica; Mount Arlington, NJ), a TKI eyedrop, has been shown to produce anti-VEGF-A biological signaling with topical once daily dosing, according to a phase 1/2 study (NCT02022540). A newer suspension was designed to reduce the incidence of punctate keratopathy as a side-effect, which occurred due to off target inhibition of corneal epithelial growth factor receptor. A new phase 1/2 clinical trial of the updated suspension for patients with nAMD (NCT03479372) has shown significantly improved safety and tolerability compared to the prior (solution) formulation. It demonstrated anti-VEGF biological response at all doses (2 mg/mL, 6 mg/mL, 10 mg/mL) in 51 treatment naïve wet AMD patients as
once daily monotherapy for 12 weeks. Twenty-six of 51 (51%) patients completed the study on PAN-90806 eye-drops alone (without rescue ranibizumab injections) through the 1-month post-treatment (week 16) visit; 23 of 26 (88%) non-rescued patients showed clinical improvement or stability based on independent masked review. Those patients requiring rescue treatment at week 4 or earlier had greater CST (444 μm) and baseline BCVA (20/100) compared to those that were never rescued (331 μm, 20/63) or rescued after 4 weeks (365 μm, 20/80). The mean number of rescue ranibizumab injections per patient was 0.84, which represents a reduction of 79.4% compared to monthly ranibizumab treatment. 59

**Sustained Delivery Treatments**

**Ranibizumab Port Delivery System**

The need for frequent intravitreal injections has spurred several companies to develop sustained-release anti-VEGF-A devices. Genentech (South San Francisco, CA) developed the ranibizumab Port Delivery System (PDS), which is a nonbiodegradable port surgically fixed to the sclera that has a 0.05 mL reservoir that can be refilled in the office.

The phase 2 LADDER trial randomized 179 patients to implantation of the PDS with doses of ranibizumab of either 10 mg/mL, 40 mg/mL, or 100 mg/mL. Among the PDS group with the 100 mg/mL ranibizumab dose, approximately 80% of patients were maintained for 6 months or longer without requiring a medication refill. In the 10 mg/mL and 40 mg/mL groups, 63.5% and 71.3%, respectively, went 6 months or more without a refill. At 9 months, BCVA change from baseline was –3.2, –0.5, +5.0, and +3.9 letters in the PDS 10 mg/mL, PDS 40 mg/mL, PDS 100 mg/mL, and monthly intravitreal ranibizumab 0.5-mg arms, respectively. Initially in the trial, rate of vitreous hemorrhage related to the surgical procedure was as high as 50%. Modifications to the surgical approach were made, including cauterization of the choroid with laser photocoagulation, which then resulted in a reduced postoperative vitreous hemorrhage rate of 4.5%. Other notable complications included endophthalmitis (1.7%), retinal detachment (2.2%), hyphema (4.5%), and cataract (7.3%). 60

The Phase 3 ARCHWAY (NCT03677934) trial is currently underway. Primary analysis of results were presented at the American Society of Retina Specialists (ASRS) annual meeting in July 2020, which demonstrated achievement of primary endpoint visual outcomes in PDS patients and reduced treatment burden compared with monthly injections of ranibizumab. Of note, 98% of patients in the PDS cohort did not require a supplemental ranibizumab injection at 6 months. There were no significant differences in safety between the two treatment arms. Expected completion of the ARCHWAY trial is in 2022. 61

**GB-102**

GB-102 (GrayBug Vision; Redwood City, CA) is sunitinib maleate, a TKI with activity against both VEGF-A and PDGF. It is an injectable depot designed for twice per year formulation. The drug is encapsulated within bioerodible polymer nanoparticles that degrade slowly over time. The patented GrayBug formulation is designed to avoid the inflammatory response seen with other nanoparticles, and the nanoparticles remain at the injection site to avoid clouding the visual axis. Phase 1/2a study (ADAGIO) results were announced in January 2019. The study met its primary endpoint of safety and tolerability, with no dose limiting toxicities or serious ocular adverse events. There was a long-lasting pharmacodynamic effect, with 3- and 6-month dosing duration achieved in 88% and 68% of patients, respectively. However, there was microparticle-related anterior chamber migration and mild ocular hypertension, which was self-limited. 62

The phase 2b ALTISSIMO study in wet AMD tested a new GB-102 manufacturing process optimized to eliminate particle dispersion. Two doses of GB-102 (1 mg and 2 mg) injected every 6 months were compared against aflibercept injected every 2 months. Following an interim safety analysis, the 2 mg dose arm was discontinued and those patients received the 1 mg dose for their second treatment. The median time to first supportive therapy was 5 months for the GB-102 1 mg group. Forty-eight percent of the patients did not require supportive therapy for at least 6 months, and 62% did not require supportive therapy for at least 4 months more. In all, the GB-102 1 mg dose was well-tolerated by patients. The majority of drug-related adverse events were mild-to-moderate, and medication was detected in the anterior chamber in less than 10% of the 1 mg injections. No vision-threatening inflammation was observed and there were no significant rises in intraocular pressure. CST in the GB-102 1 mg arm was similar to that of the aflibercept control arm, though mean change in BCVA for all 20 completers was approximately 9 ETDRS letters lower compared to aflibercept across all time points. A 6-month extension has been
added to the trial, after which full analysis of the trial results will determine future directions.

**OTX-TKI and OTX-AFS**

Ocular Therapeutix (Bedford, MA) developed a sustained release formulation of axitinib, a TKI, that is delivered through intravitreal injection. OTX-TKI is a bioresorbable hydrogel that contains TKI particles in an injectable fiber, designed to deliver drug to target tissues for up to 12 months. The hydrogel creates a meshwork that encloses the drug and gradually dissolves, allowing the medication to be released over time. Phase 1 studies of axitinib intravitreal implant (OTX-TKI) are in process; the first patient was dosed in the second quarter of 2018. OTX-AFS is an aflibercept suprachoroidal injection that is being developed in collaboration with Regeneron, which is still in the preclinical phase of investigation.

**CLS-AX**

Clearside Biomedical (Alpharetta, GA) is currently sponsoring a Phase 1/2a trial of CLS-AX, a suprachoroidally administered formulation of axitinib. Suprachoroidal delivery can target therapy to affected chorioretinal tissues for potential efficacy benefits, compartmentalizing it away from unaffected vitreous and anterior segment tissues for potential safety benefits, and achieve durability through the use of insoluble suspension formulations. This approach was assessed with proof of concept therapy, CLS-TA or Xipere, a proprietary suspension of triamcinolone, for uveitic macular edema, culminating in a successful phase 3 clinical trial; durability was demonstrated in an extension study, with a new drug application planned for 2021. In preclinical studies of axitinib, compared to intravitreal administration, suprachoroidal administration resulted in much greater targeted levels of axitinib in retinal and choroidal tissues, with lower levels in the vitreous and anterior segment, which could minimize the risk of off-target effects.

**Durasert**

The Durasert™ Bioerodible TKI (EyePoint Pharmaceuticals; Watertown, MA, USA) is a sustained-release implant that delivers vorolanib, a TKI, which was previously assessed in an oral formulation by another sponsor. IND-enabling studies with the Durasert implant are in progress.

**Gene Therapy**

The eye has become a target for investigational gene therapy due to the monogenic nature of many inherited retinal diseases (IRD), its accessibility, tight blood–ocular barrier, the ability to non-invasively monitor for functional and anatomic outcomes, as well as its relative immune privileged state. Gene therapy offers potential for long-term continuous expression of anti-VEGF-A protein with a single administration. Viral vectors are conduits for transferring desired genetic material to host cells. Vectors currently used in ocular gene therapy clinical trials include adeno-associated virus (AAV), small single-stranded DNA viruses of the parvovirus family, and lentivirus, RNA viruses of the retrovirus family. After transduction, the target cells transcribe and translate the viral genetic information into therapeutic protein, which then curbs the pathogenesis of the targeted disease process. Previous gene therapy candidates such as rAAV.sFLT-1 (Avalanche Biotechnologies, now known as Adverum Biotechnologies; Redwood City, CA), AAV2-sFLT01 (Sanofi Genzyme; Cambridge, MA), and Retinostat (Oxford Biomedica; Oxford, UK) did not advance into later studies but have paved the way for current candidates, ADVM-022 and RGX-314.

**ADVM-022**

Adverum Biotechnologies’ gene therapy product, ADVM-022, utilizes an AAV vector (AAV.7m8) that has been optimized for intravitreal injection of vectors carrying anti-VEGF cDNA, leading to the continuous expression of aflibercept. The Phase 1 OPTIC trial (NCT03748784) is a multi-center, open-label, dose-escalation trial that enrolled 30 patients with nAMD who were good responders to prior anti-VEGF therapy. Interim results were presented at the Angiogenesis, Exudation and Degeneration 2021 virtual meeting. There were four cohorts studied. Cohort 1 (n=6) received a high dose of $6 \times 10^{11}$ vector genomes (vg)/eye and a 13-day course of oral steroids, cohort 2 (n=6) received a low dose ($2 \times 10^{11}$ vg/eye) and a 13-day course of oral steroids, cohort 3 (n=9) received a low dose and a 6-week course of topical steroids, and cohort 4 (n=9) received a high dose and a 6-week course of topical steroids.

ADVM-022 had a favorable safety profile in all groups, with mild inflammation responsive to steroid drops. Patients were eligible for aflibercept rescue injection if they had an increase in CST >75 μm from baseline, loss of ≥10 letters from baseline attributed to intraretinal
or subretinal fluid, or the presence of vision threatening hemorrhage due to nAMD. In cohort 1, which had follow-up of median 86 weeks (range=64–92 weeks), a single intravitreal injection of ADVM-022 resulted in no need for supplemental injections in all six patients. Cohort 2 had a median follow-up of 64 weeks (range=64–68 weeks), and 3/6 patients did not need supplemental injections. Cohort 3 had a median follow-up of 48 weeks (range=32–48 weeks), and 7/9 patients did not need supplemental injections. Cohort 4 had a follow-up range of 12–24 weeks, and 8/9 patients did not need supplemental injections. Mean BCVA was maintained and CST was maintained or improved in all groups. The phase 2 Pivotal-a and Pivotal-b trials are planned to initiate trials in late 2021, and will compare ADVM-022 against aflibercept given every 8 weeks in treatment-naïve nAMD patients.\textsuperscript{70}

**RGX-314**

Regenxbio (Rockville, MD) has developed a novel AAV8 vector, RGX-314, which expresses a soluble anti-VEGF monoclonal antibody fragment, similar to ranibizumab, in transduced retinal cells. The company advocates that their proprietary gene delivery platform (NAV\textsuperscript{®} Technology Platform) may yield higher levels of anti-VEGF expression than earlier generation AAV vectors.\textsuperscript{71}

In the phase 1/2a trial (NCT03066258), RGX-314 was administered via subretinal injection during vitrectomy. Forty-two patients with severe wet AMD requiring frequent anti-vascular endothelial growth factor (anti-VEGF) injections were treated across five dose cohorts, with doses ranging from $3 \times 10^9$ GC/eye to $2.5 \times 10^{11}$ GC/eye. New data were reported at the Angiogenesis, Exudation, and Degeneration 2021 conference from Cohort 4 and 5 of RGX-314 Phase I/IIa trial for nAMD and Cohort 3 of its Long-Term Follow-Up study. The data demonstrates potential continued treatment effect out to 3 years. As of January 22, 2021, there was a meaningful reduction in anti-VEGF treatment burden in both Cohorts 4 and 5 as well as stable VA and decreased CST (mean change of −46 µm and −93 µm, respectively). When compared to the mean annualized injection rate during the 12 months prior to RGX-314 administration, patients in Cohort 4 received a mean of 4.4 injections over 1.5 years after RGX-314 administration, and those in Cohort 5 received a mean of 1.7 injections over that time span (a decrease in anti-VEGF treatment burden of 58.3% and 81.2%, respectively). A number of study patients continued into the Long-Term Follow-Up study (NCT03999801) to assess safety up to 5 years after RGX-314 administration, and as of late January 2021 these patients have largely shown no new drug-related ocular adverse events. Over a 3-year period, all six patients from Cohort 3 of the Phase 1/2a trial enrolled in the LTFU study demonstrated a mean BCVA improvement of +12 letters from baseline, and retinal anatomy remained stable at 3 years compared to at 2 years. These patients have also shown a long-term reduction in anti-VEGF treatment burden over the 3-year period, with four out of the six patients receiving no anti-VEGF injections from 9 months to 3 years after RGX-314 administration.\textsuperscript{72}

The long-term data from dose-escalation in Phase 1/2a trials have allowed for further advancement in clinical trials for this potential gene therapy for the treatment of wet AMD. Regenxbio has announced the first of two planned phase 2b/3 trials, ATMOSPHERE (NCT04704921), to evaluate safety and efficacy of RGX-314. The goal is enrollment of approximately 700 patients with nAMD. In one trial the patients will be enrolled across two RGX-314 dose arms versus ranibizumab with primary endpoint being non-inferiority to ranibizumab based on BCVA at 1 year. The second trial, in a similar fashion, will be compared against aflibercept. Both trials are expected to begin dosing patients in 2021. The phase 2 AAVIATE trial (NCT 0454653) is currently enrolling nAMD patients for the evaluation of two dose arms of suprachoroidal injection of RGX-314, which will be compared against monthly ranibizumab injections.\textsuperscript{73}

**Discussion**

The development of anti-VEGF therapy has significantly changed the landscape for nAMD management. Landmark trials such as the ANCHOR study demonstrated superiority of anti-VEGF injections over photodynamic therapy for preserving and recovering visual acuity.\textsuperscript{74} The currently available anti-VEGF-A medications in North America include bevacizumab, ranibizumab, aflibercept and brolucizumab. While these medications have drastically improved visual outcomes for nAMD patients globally, there remains a need for new drugs and delivery systems to address the shortcomings of anti-VEGF treatment.

Drugs in the pipeline aim to achieve broad VEGF inhibition, often through TKIs, or to modulate synergistic targets such as Ang-2, PDGF, VEGF-C, and VEGF-D with hopes of outperforming anti-VEGF-A monotherapy. Other
drugs utilize sustained release therapy and nanoparticle technology in order to extend the duration of the anti-VEGF effect. Topical anti-VEGF therapies have been developed with hopes to reduce the burden of frequent injections. Gene therapy could serve as a one-time treatment by using the retina’s own cellular apparatus to continuously express anti-VEGF molecules.

The management of nAMD can be burdensome for physicians, patients, families, and the healthcare system. The current paradigm requires an indefinite length of treatment in many cases, and patients need to be counseled that maintaining stable vision (as opposed to achieving visual gains) is still considered a success given the degenerative nature of the disease. In general, real-world outcomes of anti-VEGF treatment do not live up to those obtained in clinical trials due to inadequate numbers of injections to optimize visual gains. The “treat-and-extend” regimen, in which injections are gradually spaced out further apart until a recurrence of fluid occurs, has gained popularity in attempt to reduce the treatment burden of frequent injections. This regimen appears to perform relatively well in practice based on results of the LUCAS, TREX-AMD, and TREND trials.

Given the unparalleled economic value of bevacizumab compared to other anti-VEGF treatments, it seems likely that it would continue to be a popular first line treatment for most VEGF-mediated diseases globally, with the newly-approved products of the future being utilized more often in refractory cases or in patients in need of longer duration therapy. As biosimilars for aflibercept and ranibizumab become approved and more widely available, they may rival bevacizumab as the most cost-effective first line treatment option.

Anti-VEGF/anti-PDGF combination therapy initially appeared like a promising strategy to address the treatment ceiling of anti-VEGF therapy, however results of the trials of intravitreal rinucumab/aflibercept and pegpleranib/ranibizumab failed to show any advantage over anti-VEGF monotherapy. Similarly, topical anti-VEGF/anti-PDGF therapies for nAMD so far have been unsuccessful in reducing the need for rescue intravitreal injections, as evidenced by the trials of squalamine lactate, pazopanib, and regorafenib. Based on the early promising results of the phase 1/2 trial, topical PAN90806 could have a possible role for nAMD patients with mild fluid as monotherapy, as an adjuvant to intravitreal anti-VEGF therapy for refractory cases, or as chronic maintenance after loading injections.

In 2019, brolucizumab became the first anti-VEGF drug approved for 12-week dosing. Similarly, abicipar showed promise as a possible 12-week treatment option. However, the issue of increased intraocular inflammation with both of these drugs compared to other anti-VEGF options has hampered their widespread use. In the case of brolucizumab, rare but severe cases of occlusive retinal vasculitis were reported after it became commercially available. The Research and Safety in Therapeutics (ReST) committee of the American Society of Retina Specialists (ASRS) has been working with Novartis in tracking the data, which as of June 2020 demonstrate incidence of retinal vasculitis to be 8.65 per 10,000 injections. Data reported at the American Academy of Ophthalmology meeting in November 2020 cited 26 eyes of 25 patients with retinal vasculitis in which vision loss was more severe, with 35% of eyes losing six lines of vision or more without recovery. Currently the FDA has not advised against use of brolucizumab, while they work with the ASRS to closely monitor these reports of intraocular inflammation.

Clinical research has continued to broaden the physiological scope of drug development to include medications that upregulate Tie-2 receptor functioning, including nesvacumab, faricimab, and ARP-1536. Of these, nesvacumab did not show sufficiently promising results in Phase 2 trials for further advancement. Faricimab is the furthest under development and is differentiated by its unique bispecific antibody design that targets VEGF and Ang-2. In the phase 2 STAIRWAY trial, it showed promising results with visual gains comparable to monthly dosing of ranibizumab with 12- and 16-week dosing of faricimab. Phase 3 trials (LUCERNE and TENAYA) comparing a flexible dosing regimen of faricimab (up to 16-week dosing) against aflibercept every 8 weeks are expected to have results in 2022.

The ranibizumab Port Delivery System is the furthest sustained-release anti-VEGF therapy in development. The phase 3 Archway Study is expected to conclude in 2022; interim results indicate that 98% of patients were able to go 6 months without additional treatment while maintaining visual acuity comparable to the monthly ranibizumab group. The need for surgery to implant the refillable port, with higher risk of complications compared to intravitreal injection, may be a deterring factor for some patients. While GB-102 showed an impressive durability and safety profile in its phase 2b trial, it is unclear why patients randomized to GB-102 gained 9 less letters compared to those receiving aflibercept, and this will need to be explored further in a larger scale trial.

Gene therapy with viral vectors could represent the ultimate form of sustained anti-VEGF treatment, as the
retina develops its own anti-VEGF biofactory that would diminish the need for subsequent intravitreal injections. While there have been several gene therapy candidates in the past, currently ADVM-022 and RGX-314 are the two that are primed for phase 2 trials in the near future.

In summary, the pathogenesis of nAMD is multifaceted and includes an array of factors that could be targeted in future treatments. Emerging therapies that target alternative pathways, along with novel anti-VEGF therapies that provide extended durability, will provide the retina community with better options for refractory nAMD and reduce the need for monthly injections in many patients.

Disclosure
Dr Audina M. Berrocal reports personal fees, DORC, ALCON, ProQR, AGTC, Novartis, during the conduct of the study. Dr. Rehan Hussain has served on an advisory board for Alimera Sciences. Dr. Jayanth Sridhar is a consultant for Alcon, DORC, Genentech, Regeneron, and Oxurion. The authors report no other conflicts of interest in this work.

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