Intravitreal aflibercept for exudative age-related macular degeneration with good visual acuity: 2-year results of a prospective study

Aflibercept is a recombinant fusion protein comprising a vascular endothelial growth factor (VEGF) receptor dimer fused to a human serum albumin tail. It has been shown to have robust efficacy for the treatment of exudative age-related macular degeneration (AMD) with good visual acuity. The purpose of this study was to report the 2-year outcomes of intravitreal aflibercept (IVA) for exudative AMD with good visual acuity (VA) and examine the baseline factors associated with good visual outcome.

Materials and methods: This multicenter, prospective study evaluated 39 eyes (39 AMD patients) enrolled from August 2013 to August 2014 at 12 and 24 months. Only patients with initial best-corrected VA (BCVA) >0.3 logarithm of the minimum angle of resolution (20/40 Snellen) were eligible. Three consecutive monthly IVA injections were followed by 2 monthly injections for 12 months. Thereafter, patients received injections on a treat-and-extend regimen for up to 24 months. Outcome measures included BCVA and central macular thickness (CMT) at 12 and 24 months. Post hoc analysis, BCVA, and CMT were evaluated by AMD types (typical AMD [tAMD], type 1, and type 2 polypoidal choroidal vasculopathy [PCV]). Baseline characteristics and BCVA associations were evaluated with linear regression analysis and Student’s t-test.

Results: Mean age was 69 years and 26 of 39 eyes were male. tAMD, type 1 and type 2 PCV occurred in 18, 12, and 9 eyes, respectively. Baseline mean BCVA was 0.097 logarithm of the minimum angle of resolution (20/25 Snellen) and showed significant improvement to 0.058 (20/22 Snellen, P=0.03) at 12 months and 0.066 (20/23) at 24 months. CMT improved significantly from 320 (99) µm (mean [SD]) to 250 (93) µm (P=0.002) at 12 months and 240 (93) µm (P=0.0005) at 24 months. BCVA and CMT were not significantly different among the three groups. Only subretinal hemorrhage (SRH) was significantly associated with improved BCVA. BCVA change from baseline was −0.12 with SRH and −0.011 without SRH (P=0.017) at 12 months.

Conclusion: IVA showed good efficacy for exudative AMD with good VA at 24 months. tAMD and type 1 and 2 PCV showed similar prognosis. Baseline SRH predicted favorable long-term vision in AMD with good VA.

Keywords: polypoidal choroidal vasculopathy, aflibercept, age-related macular degeneration, fixed regimen, good visual acuity, prospective study, treat-and-extend

Introduction

Age-related macular degeneration (AMD), the leading cause of blindness in developed countries, has seen several changes in treatment modalities over time. The clinical course of AMD is unfavourable, but treatment with anti-vascular endothelial growth factor (anti-VEGF) agents has improved visual acuity (VA), and better long-term prognosis for VA is expected. Aflibercept was approved in November 2012 for use in Japan and is reported to have long-term therapeutic effects. Aflibercept is a recombinant fusion protein comprising...
two main components: vascular endothelial growth factor (VEGF)-binding portions from the extracellular domains of human VEGF receptors 1 and 2, which are fused to the fragment crystallizable portion of human immunoglobulin G1. This soluble decoy receptor acts by binding to VEGF-A and placentals growth factor. Both factors are thought to be involved in the pathologic neovascularization and angiogenesis seen in ophthalmologic diseases such as AMD.

Aflibercept also binds VEGF-B, which may be involved in ophthalmologic disease. In the VIEW 1 and VIEW 2 studies, which assessed the efficacy and safety for treating AMD, intravitreal aflibercept (IVA) injections were administered once monthly or bimonthly after three administrations during the loading phase for 12 months and, as-needed regimen, but at least every 12 weeks dosing, from 12 to 24 months. Efficacy and safety were found to be similar to those of once-monthly intravitreal injections of ranibizumab.

Polypoidal choroidal vasculopathy (PCV) accounts for about half of all cases of AMD in East Asia. Photodynamic therapy (PDT) is effective in PCV, but not in typical AMD (tAMD; AMD without PCV and retinal angiomatosus proliferation). Aflibercept is almost equally effective in tAMD and PCV. Kawamura et al defined type 1 PCV as multiple polypoidal lesions around a network of vessels and type 2 PCV as a single polypoidal lesion without a network of vessels. Studies from other Japanese groups proved useful for predicting the treatment outcome of PDT, showing that PDT is not effective in type 1 PCV but effective in type 2 PCV. Another report compares the short-term treatment outcomes between type 1 and 2 PCV in patients receiving anti-VEGF therapy.

There are, however, two important unanswered questions. First, there is the lack of evidence for the use of aflibercept in AMD patients with good VA. Notably, in real-world practice, a substantial proportion of AMD patients present with better VA than those included in the aforementioned clinical trials. Many AMD studies have focused on patients with better VA than those included in the aforementioned clinical trials. Many AMD studies have focused on patients with better VA than those included in the aforementioned clinical trials. Many AMD studies have focused on patients with better VA than those included in the aforementioned clinical trials.

Primary endpoints were mean change of BCVA from baseline to 12 months and ratio of patients with improved BCVA at 12 months. Secondary endpoints were mean change of BCVA from baseline to 24 months and ratio of patients with improved BCVA at 24 months, mean change of retinal thickness in the central subfield on OCT from baseline to 12 and 24 months, ratio of patients with subretinal fluid at 12 and 24 months, and mean change of greatest linear diameter (GLD) of disease area. Further secondary endpoint of PCV was ratio of patients with regression of polypoidal lesions. Safety outcomes were adverse events and side effects of the procedure.

Patients

Inclusion criteria were adults aged \( \geq 50 \) years; AMD with exudative signs on indocyanine green (ICG) angiography, fluorescein angiography, or optical coherence tomography (OCT); treatment-naive status; and initial BCVA over 0.3 logMAR (20/40 Snellen equivalent).

Exclusion criteria were intraocular surgery in the target eye within 3 months; history of vitreous surgery in the target eye; inflammation or infection in the intraocular, outer ocular area, or around either eye; history of uveitis in either eye; hypersensitivity or allergy to fluorescein or ICG; clinically significant drug allergy or known hypersensitivity to therapeutic or diagnostic protein products, or to any of the study drugs or their components; pregnancy, possibility of pregnancy, or lactation; and any history that might affect the interpretation of the results of the study or put the patient at an increased risk of treatment complications.

In a post hoc analysis, we also investigated the baseline factors associated with good visual outcomes at 12 months.

Materials and methods

Study design

This multicenter, prospective, non-randomized, interventional exploratory study was performed in an institutional setting.

Institutional review board approval

Institutional review board approval was obtained from Jichi Medical University (A13-23, B14-54) and the Japan Community Health Care Organization (JCHO) Tokyo Shinjuku Medical Center (H25-3, H.26-3).

Patients

Inclusion criteria were adults aged \( \geq 50 \) years; AMD with exudative signs on indocyanine green (ICG) angiography, fluorescein angiography, or optical coherence tomography (OCT); treatment-naive status; and initial BCVA over 0.3 logMAR (20/40 Snellen equivalent).

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Subjects were 47 AMD patients (47 eyes) with BCVA >0.3 logMAR (Table 1). All subjects received their first IVA at either Jichi Medical University or JCHO Tokyo Shinjuku Medical Center (H25-3, H.26-3).
Table 1 Enrolled patient characteristics

<table>
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<tr>
<th>Characteristics</th>
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<td>Age, years</td>
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<td>Sex</td>
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<tr>
<td>Male</td>
<td>30 eyes</td>
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<tr>
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</tr>
<tr>
<td>Type 1 PCV</td>
<td>15 eyes</td>
</tr>
<tr>
<td>Type 2 PCV</td>
<td>10 eyes</td>
</tr>
<tr>
<td>BCVA</td>
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</tr>
<tr>
<td>CMT (µm)</td>
<td>320 (100, 200 to 700)</td>
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</tbody>
</table>

Abbreviations: AMD, age-related macular degeneration; BCVA, best-corrected visual acuity; CMT, central macular thickness; PCV, polypoidal choroidal vasculopathy.

Medical Center between August 2013 and August 2014. Written informed consent was obtained from all consented patients. The study protocol adhered to the tenets of the Declaration of Helsinki and was registered in the University Hospital Medical Information Network Clinical Trials registry prior to study commencement (UMIN000011337 for the first year and UMIN000016598 for the second year). The protocol was based on the VIEW 1 and VIEW 2 studies. First, IVA injections were administered once monthly for three consecutive months and bimonthly thereafter. Treatment continued for 12 months. Second, IVA injections were administered on treat-and-extend regimens from 13 to 24 months. On signs of recurrent choroidal neovascularization (CNV) activity (increased exudative change at any visit after 12 months), IVA was administered and the interval to the next visit was shortened by 1 month versus the last interval. Additionally, on signs of new subretinal hemorrhage (SRH) at any visit after 12 months, IVA was administered and the interval to the next visit was set to 1 month. If there was a minor exudative lesion (residual exudation), IVA was administered and the interval to the next visit remain unchanged. With no sign of recurrent CNV activity, IVA was administered and the interval to the next visit was extended by 1 month versus the last interval. Minimum dosing interval was 1 month, and maximum dosing interval was not regulated.

Among the 47 subjects, three dropped out before starting treatment (one withdrew consent and two experienced decreased VA and no longer satisfied the inclusion criteria). Also, five subjects dropped out after the third administration of the loading phase (one demonstrated no effects during the loading phase, withdrew consent, and received PDT; one showed insufficient effects after the first course of maintenance therapy, withdrew consent, and received PDT; one developed uveitis; and two withdrew consent and stopped IVA due to the high treatment costs). At 12 months, 39 subjects completed the study. Of these, four dropped out during the second year (two withdrew consent and stopped IVA due to high treatment costs; one withdrew only study consent and continued treat and extend; and one experienced worsening of exudative lesions with bimonthly injections, withdrew consent, and received PDT). At 24 months, 35 subjects completed the study. A flow diagram of the study is shown in Figure 1.

Measurements

The full analysis set was 39 subjects; four subjects dropped out during the second year. Their missing values up to 24 months were imputed using the last observation carried forward analysis.

On study commencement, we performed comprehensive ophthalmologic examination on all 39 subjects, including measuring decimal BCVA using Landolt rings and performing color fundus photography. Fluorescein angiography and ICG angiography were performed using a confocal scanning laser ophthalmoscope (HRA2; Heidelberg Engineering GmbH, Heidelberg, Germany) at Jichi Medical University and a fundus camera (TRC-50; TOPCON Corporation, Tokyo, Japan) at JCHO Tokyo Shinjuku Medical Center. Swept source OCT (DRI OCT-1 Atlantis, TOPCON) or spectral domain OCT (RS-3000 Lite; NIDEK Co, Ltd, Aichi, Japan) was performed at Jichi Medical University, and spectral domain OCT (Cirrus HD-OCT Model 4000; Carl Zeiss Meditec AG, Oberkochen, Germany) was performed at JCHO Tokyo Shinjuku Medical Center. Decimal BCVA assessment and dilated fundus examination were performed at every visit. Axial length was examined using A-mode ultrasonography (UD-6000; TOMEY Corp., Aichi, Japan).

Diagnosis of AMD was made by experienced retinal specialists using color fundus photography, fluorescein angiography, and ICG angiography. Fluorescein angiography and ICG angiography were done at baseline, 12 months, and 24 months after treatment. PCV was diagnosed based on the presence of characteristic polyp-like aneurysmal dilatations in the choroidal vessels in ICG angiography findings and the presence of orange-red protruding lesions on the ocular fundus in ophthalmoscopy findings. GLD was measured using fluorescein angiography.

We measured the central macular thickness (CMT) manually during OCT between the inner limiting membrane and Bruch’s membrane using an OCT measurement function. Bruch’s membrane was identified by the presence of a highly reflective line photographed under the retinal pigment.
epithelium in cases with subfoveal pigment epithelial detachment, and using the lower end of the highly reflective line in the retinal pigment epithelium in cases without pigment epithelial detachment.

Subanalysis
Eyes with multiple polypoidal lesions around a network of vessels on ICG angiography were set to type 1 PCV. Eyes with a single polypoidal lesion without a network of vessels were set to type 2 PCV. Similarly, we assessed polyp regression on ICG angiography at 12 and 24 months. In this study, only two eyes had type 2 CNV. We could not perform statistical analysis on types 1 and 2 CNV.

Statistical analysis
Statistical analysis was performed using JMP Pro software version 11.2.0 (SAS Institute, Cary, NC, USA). Decimal BCVA was converted to logMAR. BCVA and CMT were compared with the first visit by single-sided paired t-test. Improvement of BCVA and CMT were compared between type 1 and type 2 PCV by double-sided t-test. Power analysis was performed, and the least significant number, which was needed for analysis of significance, was calculated. A P-value of <0.05 was considered statistically significant. The equality test of BCVA prognosis was calculated with logMAR 0.15 as clinically equivalent. Maximum value of an upper P-value and a lower P-value was used. Analysis of variance was used to evaluate the baseline characteristics of AMD and type 1 and type 2 PCV, except sex and GLD, which were analyzed by Fisher’s exact test and Kruskal–Wallis test, respectively. Since this research is an explanatory study, Bonferroni correction was not done.

Associations between the baseline factors and good visual prognosis were examined. Influence of baseline characteristics (age, axial length, logMAR BCVA, GLD, CMT and numbers of injections) for logMAR BCVA was evaluated by linear regression analysis. We also examined the baseline characteristics (sex, presence of posterior vitreous detachment, types of CNV, intraretinal fluid, subretinal fluid, SRH, pigment endothelial detachment, and hemorrhagic pigment endothelial detachment) by Student’s t-test. Correlations were evaluated between statistically significant baseline characteristics and awareness ratio of VA drop at the first visit to local ophthalmologists and SRH recurrences in 1 year.

Figure 1 Flow diagram of the study.
Abbreviation: PDT, photodynamic therapy.
Results

A total of 47 eyes in 47 subjects were analyzed (Table 1). At 12 months of treatment, we were able to analyze the clinical course of 39 eyes of 39 subjects (25 men and 14 women) of mean (SD) age 67 (9) years with an age range of 52–88 years. Mean CMT was 250 (93) µm. There were 18 eyes with tAMD (16 type 1 CNV and 2 type 2 CNV) and 21 eyes with PCV (12 type 1 PCV and 9 type 2 PCV). There were no significant differences between tAMD, type 1 PCV, and type 2 PCV in age, sex, BCVA, GLD, and CMT (Table 2). Representative PCV cases are shown in Figure 2. BCVA at the start of treatment was 0.10 (0.11) logMAR. At the end of 2 years of treatment, the mean CMT was 240 (93) µm.

At 12 months, the BCVA was 0.06 (0.15) logMAR, indicating improvement (P=0.03; Figure 3). There were 5/39 patients (12.8%) in whom the BCVA improved by 0.2 logMAR or more, 34 patients (87.2%) in whom there was no change, and no cases in which there was worsening of 0.2 logMAR or more. At 24 months, the BCVA was 0.07 (0.14) logMAR, indicating no significant improvement (P=0.09).

Mean CMT after 12 and 24 months was 250 (93) µm (P=0.002) and 240 (93) µm (P=0.0005), indicating a significant reduction, respectively (Figure 4). The mean number of aflibercept injections during the 12 months was 6.7±0.72 (39 cases) and from 13 to 24 months was 4.8±1.38 (35 cases). There was no significant difference in BCVA or CMT between the AMD cases with and without PCV (Figure 5).

At 12 months, the polypoidal lesions had completely regressed in 12/21 PCV patients (57%), partially regressed in 2 (10%), and enlarged in 2 (10%). In type 1 PCV, the polypoidal lesions had completely regressed in 6/12 PCV patients (50%), partially regressed in 2 (17%), and enlarged in 2 (17%). In type 2 PCV, the polypoidal lesions had completely regressed in 6/9 PCV patients (67%). There was no statistical difference in the ratio of complete regression between type 1 and 2 PCV (P=0.66, Fisher’s exact test).

At 24 months, the polypoidal lesions had regressed completely in two additional type 1 PCV cases and were unchanged in type 2 CNV from 12 months. Ratio of complete regression between type 1 and 2 PCV (P=0.66, Fisher’s exact test) did not differ significantly. Mean 1- and 2-year prognosis of BCVA and CMT was −0.035 and −0.027 logMAR, 87 and 103 µm in type 1 PCV, 0.0086 and 0.046 logMAR, and 57 and 77 µm in type 2 PCV, respectively. There were no significant differences between type 1 and type 2 PCV in BCVA prognosis (P=0.29 and 0.79, least significant number: 72 at 1 year and 1,100 at 2 years, respectively) or CMT prognosis (P=0.56 and 0.68, respectively). It was rejected that there were over 0.15 logMAR differences between type 1 and type 2 PCV in terms of BCVA prognosis (P=0.024 and 0.043, respectively; Figure 6).

Baseline SRH was a significantly good factor for BCVA prognosis at 12 months. Change of logMAR BCVA at 12 months was −0.12 with SRH and −0.011 without SRH (P=0.017; Table 3; Figure 7). Baseline logMAR BCVA correlated with the prognosis of logMAR BCVA at 12 months (R−=−0.44, P=0.005). Other factors were not significant. Baseline SRH was correlated with the awareness ratio of VA drop at the first visit to local ophthalmologists (11/12 eyes with SRH, 16/35 eyes without SRH, P=0.006, Pearson’s chi-square test) and SRH recurrences in 1 year (mean: 1.6 versus 0.12 times, P<0.0001). AMD was pointed out in most patients without self-awareness by local ophthalmologists during the visits for common eye diseases, such as cataract or dry eye.

Among the 44 subjects who started this study, three withdrew consent due to inadequate effects of monthly or bimonthly IVA and requested PDT combination therapy in anticipation of reducing the frequency of IVA. One patient had type 1 PCV and the others had type 1 CNV. All three patients did not receive IVA after PDT up to month 24, and only one CNV patient experienced VA loss over 0.15 logMAR.

The other four patients withdrew consent and stopped injections due to high costs. One patient had type 1 PCV, one had type 2 PCV, and two had type 1 CNV. Three of the four patients continued to visit the hospital up to month 24 and received IVA once in the second year, and one of the three patients experienced VA loss of over 0.15 logMAR.

One patient developed endophthalmitis 2 days after IVA at 24 months. However, treatment was successful and did not affect visual prognosis. No other serious adverse events.

Table 2 Baseline characteristics of tAMD, type 1 PCV, and type 2 PCV

<table>
<thead>
<tr>
<th></th>
<th>tAMD (n=18)</th>
<th>Type 1 PCV (n=12)</th>
<th>Type 2 PCV (n=9)</th>
<th>P-value</th>
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<tr>
<td>Age (years)</td>
<td>69 (11)</td>
<td>65 (9)</td>
<td>67 (9)</td>
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<tr>
<td>Sex (male:female)</td>
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<td>7:5</td>
<td>4:5</td>
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<td>BCVA (logMAR)</td>
<td>0.059 (0.14)</td>
<td>0.045 (0.10)</td>
<td>0.041 (0.15)</td>
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<td>GLD (µm)</td>
<td>2,500 (2,100)</td>
<td>3,400 (1,300)</td>
<td>2,400 (1,100)</td>
<td>0.09</td>
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<tr>
<td>CMT (µm)</td>
<td>260 (120)</td>
<td>230 (51)</td>
<td>250 (62)</td>
<td>0.7</td>
</tr>
</tbody>
</table>

Notes: Data expressed as mean (SD). P-value using ANOVA, except sex (Fisher’s exact test) and GLD (Kruskal–Wallis test).

Abbreviations: ANOVA, analysis of variance; BCVA, best-corrected visual acuity; CMT, central macular thickness; GLD, greatest linear diameter; logMAR, logarithm of minimum angle of resolution; PCV, polypoidal choroidal vasculopathy; tAMD, typical age-related macular degeneration.
Figure 2 Indocyanine green angiogram of representative cases of type 1 and 2 PCV.

Notes: (A) Type 1 PCV at initial injection. (B) Type 1 PCV at 12 months. Polypoidal lesions are enlarged and number of polyps is increased. (C) Type 1 PCV at 24 months. Some polyps are enlarged and the lesions remain unchanged. (D) Type 2 PCV at initial injection. (E) Type 2 PCV at 12 months. Polypoidal lesions have disappeared. (F) Type 2 PCV at 24 months. No reappearance of polypoidal lesions.

Abbreviation: PCV, polypoidal choroidal vasculopathy.

Figure 3 BCVA over 2 years of treatment (mean, SD).

Notes: Thirty-nine eyes with wet age-related macular degeneration and BCVA of ≥0.3 logMAR received three monthly intravitreal injections of aflibercept (IVA) followed by IVA every other month for up to 12 months and then with a modified treat-and-extend regimen from 12 to 24 months. Mean BCVA of 0.058 (20/22 Snellen equivalent) at 12 months significantly improved from initial mean BCVA of 0.097 (20/25 Snellen equivalent; paired t-test, P=0.03). Mean BCVA of 0.066 (20/23 Snellen equivalent) at 24 months was not significantly different from initial mean BCVA (paired t-test, P=0.09). *P<0.05.

Abbreviations: BCVA, best-corrected visual acuity; IVA, intravitreal aflibercept; logMAR, logarithm of minimum angle of resolution.
Aflibercept for AMD with good VA

were observed due to physical adverse drug reactions, such as cardiovascular events.

**Discussion**

In Japan, the treatment guidelines for AMD recommend anti-VEGF monotherapy for AMD patients with good BCVA.\(^1\) However, there is no direct prospective evidence supporting this. Hence, we planned this study and found that aflibercept monotherapy could maintain good BCVA. Here, we observed maintenance of BCVA and improvement in CMT at 12 and 24 months after the start of the planned aflibercept regimen, even in favorable cases with BCVA >0.3 logMAR. Because untreated AMD is a progressive disease, treatment should be started early, even in patients with good BCVA.\(^2\)

In the VIEW 1 and VIEW 2 studies, AMD subjects received their first treatment with BCVA ranging from 0.3 to 1.2 logMAR\(^4\) (Early Treatment Diabetic Retinopathy Study acuity: 25–73 letters). Improvement was observed in both studies. In the group that received aflibercept 2 mg bimonthly after three monthly administrations during the loading phase, mean BCVA was 53.6 before treatment, with an 8.4-letter improvement at 12 months. Between 12 and 24 months, the same group received aflibercept 2 mg as needed with mandatory dosing at least every 12 weeks, and mean BCVA showed a 7.6-letter improvement at 24 months.

Our subjects had good BCVA before treatment, so we observed an improvement of only 0.04 and 0.03 logMAR at 12 and 24 months, respectively, due to ceiling effect. Few cases showed improvement, but there was only one case that worsened. Three of the patients who dropped out developed residual serous retinal detachment and were treated with PDT, so worsening can occur even in patients with good VA.

Previous reports have indicated that treating AMD with anti-VEGF agents may not always be effective,\(^20–23\) and investigation of treatments for such cases is necessary. Consideration should be given as to whether to continue monthly administration, switch to another anti-VEGF agent, or use an anti-VEGF agent in combination with PDT. In this study, baseline SRH and baseline good BCVA were independently associated with improved BCVA at 12 months. Notably, even though eyes with initial SRH tend to have recurrent SRH, eyes with initial SRH had good prognosis. Thus, eyes with baseline SRH might likely receive IVA earlier than eyes without initial SRH, because most eyes with SRH had visual symptoms themselves, whereas most without SRH were found to have exudative AMD lesions incidentally.

To our knowledge, only three papers have described treatment outcomes of anti-VEGF therapy in AMD with good VA. Compared with previous reports,\(^16–18\) BCVA was favorable after 24 months, although more injections were administered.

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**Figure 4** CMT over 2 years of treatment (mean, SD).

**Notes:** Thirty-nine eyes with wet age-related macular degeneration. Mean CMT significantly improved at 12 months (250 µm) and 24 months (240 µm) compared with baseline (320 µm) (paired t-test, \(P=0.002\) and 0.0005, respectively). \(^*P<0.05\).

**Abbreviation:** CMT, central macular thickness.
owing to a fixed regimen and modified treat-and-extend regimen (Figure 3). Axer-Siegel et al reported that mean BCVA in 150 cases of AMD did not change from the start of treatment (0.22 logMAR) after administration of a mean 11.3 bevacizumab injections over a mean 20.2 months.16 Furthermore, a report on ranibizumab injections by Lee et al indicated worsening of mean BCVA. In 754 eyes with AMD, mean BCVA was 0.23 logMAR before treatment and 0.31 logMAR after 5.7 injections over 12 months.18 According to Kato et al, mean BCVA in 36 eyes with AMD showed no significant change from 0.11 logMAR at treatment initiation to 0.12 logMAR after 3.3 injections over 12 months.17 In this study, we treated 39 eyes with AMD. BCVA improved, ranging from −0.08 to 0.26 logMAR (mean 0.097), and from −0.08 to 0.40 logMAR at 12 months (mean 0.058, P=0.03), and was maintained at −0.08 to 0.52 logMAR at 24 months (mean 0.066, P=0.09), after a mean of 6.7±0.72 injections of aflibercept over 12 months and 4.8±1.8 injections from 13 to 24 months.

This study included 21 cases of PCV (54%). Good short-term outcomes have been reported when PCV is treated with aflibercept.24 Here, we found no significant differences in BCVA or CMT between AMD cases with and without PCV. We also found no significant differences in BCVA, CMT prognosis, or polypoidal regression ratio between type 1 and 2 PCV. In some cases of type 1 PCV, polypoidal lesions may enlarge with bimonthly aflibercept. We believe IVA should be administered during the early stages of treatment in cases with good VA, irrespective of the presence of type 1 or 2 PCV. Also, the ratio of disappearance of polypoidal lesions was reported as 69%16 and 55%24 in treatment-naïve PCV patients. Here, we found the ratio was almost the same – 57% at 12 months and 67% at 24 months, respectively – in treatment-naïve PCV patients with good BCVA.

We recognize some limitations of this study. It was an exploratory study and the sample size was small for the ad hoc analysis. Further large-scale studies are needed to identify prognostic imaging biomarkers for the treatment.
Conclusion

We observed improved BCVA and CMT at 12 months and no worsening of BCVA and improved CMT after 24 months of planned administration of aflibercept in patients with AMD with a mean baseline BCVA of $0.3$ logMAR. Our patients received monthly administrations over 3 months, followed by bimonthly administration up to 12 months and administrations with a treat-and-extend regimen from 13 to 24 months. There was no significant difference in BCVA or CMT prognosis between AMD with and without PCV or between type 1 and type 2 PCV. There was no significant difference in the regression ratio of polypoidal lesions between type 1 and type 2 PCV. Baseline SRH was a good indicator of change of logMAR BCVA after 1 year, but was not a significant factor after 2 years.

Data sharing statement

All dataset files are available from the Figshare database (https://doi.org/10.6084/m9.figshare.5734857).

Table 3  Analysis of baseline characteristics and mean change of logMAR BCVA

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<th>Mean change of logMAR BCVA</th>
<th>P-value</th>
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<td>SRF−</td>
<td>0.048</td>
<td></td>
<td>−0.063</td>
<td></td>
</tr>
<tr>
<td>PED+</td>
<td>−0.047</td>
<td>0.5</td>
<td>−0.046</td>
<td>0.3</td>
</tr>
<tr>
<td>PED−</td>
<td>−0.016</td>
<td></td>
<td>−0.013</td>
<td></td>
</tr>
<tr>
<td>hPED+</td>
<td>−0.04</td>
<td>1</td>
<td>0.016</td>
<td>0.5</td>
</tr>
<tr>
<td>hPED−</td>
<td>−0.039</td>
<td></td>
<td>−0.037</td>
<td></td>
</tr>
<tr>
<td>Typical AMD</td>
<td>−0.057</td>
<td>0.4</td>
<td>−0.026</td>
<td>0.9</td>
</tr>
<tr>
<td>PCV</td>
<td>−0.024</td>
<td></td>
<td>−0.035</td>
<td></td>
</tr>
<tr>
<td>SRH+</td>
<td>−0.12</td>
<td>0.02*</td>
<td>−0.010</td>
<td>0.06</td>
</tr>
<tr>
<td>SRH−</td>
<td>−0.011</td>
<td></td>
<td>−0.0056</td>
<td></td>
</tr>
</tbody>
</table>

Notes: P-value using linear regression analysis (Fisher’s exact test). *P<0.05.

Abbreviations: AMD, age-related macular degeneration; BCVA, best-corrected visual acuity; hPED, hemorrhagic PED; IRF, intraretinal fluid; logMAR, logarithm of minimum angle of resolution; PCV, polypoidal choroidal vasculopathy; PED, pigment epithelial detachment; PVD, posterior vitreous detachment; SRF, subretinal fluid; SRH, subretinal hemorrhage.
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Author contributions

SS: analysis, writing, and editing. HT: data collection, analysis, writing, and editing. YA, SI, NK, YF, TT, HK, and YY: data collection, editing, and review of manuscript. YI: data collection, analysis, editing, and review of manuscript. All authors contributed toward data analysis, drafting and revising the paper and agree to be accountable for all aspects of the work.

Disclosure

SS, YA, SI, and NK: none. YY: lecturer’s fees from Novartis Pharmaceuticals, Bayer Healthcare, and Santen Pharmaceuticals outside this work. HT: lecturer’s fees from Kowa Pharmaceuticals, Novartis Pharmaceuticals, Bayer Yakuhin, and Santen Pharmaceuticals; grants from Novartis Pharma outside this work. YI: lecturer’s fees from Kowa Pharmaceuticals, Novartis Pharmaceuticals, Bayer Yakuhin, and Santen Pharmaceuticals outside this work. YF: lecturer’s fees from ALCON Japan and Otsuka Pharmaceuticals outside this work. TT: lecturer’s fees from ALCON Japan outside this work. HK: lecturer’s fees from Kowa Pharmaceuticals, Novartis Pharmaceuticals, and Santen Pharmaceuticals outside this work. The authors report no other conflicts of interest in this work.

References

1. Bressler NM. Age-related macular degeneration is the leading cause of blindness. JAMA. 2004;291:1900–1901.


