ORIGINAL RESEARCH

Short-Term Outcomes of 3 Monthly intravitreal Faricimab On Different Subtypes of Neovascular Age-Related Macular Degeneration

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Purpose: To evaluate the efficacy and safety of faricimab injections for treatment-naïve neovascular age-related macular degeneration (nvAMD) patients, including subtypes and pachychoroid phenotypes, and identify predictive factors for visual outcomes.

Methods: nvAMD patients were prospectively recruited, receiving three monthly faricimab (6 mg) injections. Best-corrected visual acuity (BCVA) two months after the last injection (month 4) was compared between subtypes, and between pachychoroid neovasculopathy (PNV) and non-PNV eyes. Regression analysis determined factors influencing month 4 BCVA.

Results: The study involved 23 patients (12 typical AMD [tAMD], 10 polypoidal choroidal vasculopathy [PCV], 1 retinal angiomatous proliferation [RAP]). Eleven exhibited PNV phenotype. Significant BCVA ($P = 4.9 \times 10^{-4}$) and central retinal thickness (CRT) ($P = 1.3 \times 10^{-5}$) improvements were observed post-faricimab treatment. The therapy demonstrated favourable results for both tAMD and PCV eyes, and non-PNV and PNV eyes. Faricimab achieved dry macula in 77.3% of eyes, with subretinal fluid resolution in most cases, although intraretinal fluid (IRF) often persisted. Multivariable analysis identified external limiting membrane (ELM) presence and IRF as BCVA contributors at month 4.

Conclusion: Faricimab demonstrated significant effectiveness and safety in treatment-naïve nvAMD patients, particularly for PCV and PNV eyes. ELM presence and IRF is predictive of visual outcomes.

Keywords: anti-VEGF, anti-vascular endothelial growth factor, CRT, central retinal thickness, ELM, external limiting membrane, faricimab, IRF, intraretinal fluid, MNV, macular neovascularization, nvAMD, neovascular age-related macular degeneration, PCV, polypoidal choroidal vasculopathy, PNV, pachychoroid neovasculopathy, RAP, retinal angiomatous proliferation

Introduction

Age-related macular degeneration (AMD) is a leading cause of blindness worldwide, affecting individuals in North America, Europe, Australia, and Asia.¹ Late AMD is classified into two types: geographic atrophy and neovascular AMD (nvAMD). Pathological neovascularization of the choroidal blood vessels underneath the macula or intraretinal neovascularization (macular neovascularization, MNV) is observed in patients with nvAMD, which rapidly compromises their central visual field. Over the past decade, standards of care for nvAMD have primarily focused on targeting vascular endothelial growth factor-A (VEGF-A). Intravitreal injection of anti-VEGF drugs, such as ranibizumab and aflibercept, is considered the first-line therapy for nvAMD.^{2,3} However, over 10% of patients show a suboptimal response to anti-VEGF therapy.^{4,5} Additionally, frequent injections of anti-VEGF therapy are required to maintain the vision gain achieved during the loading phase, resulting in treatment burden and possibly macular atrophy.^{6–8} Brolucizumab, which was recently approved as an anti-VEGF agent based on the HAWK and HARRIER, has demonstrated better fluid control and longer dosing intervals.⁹ However, there are significant concerns regarding intraocular inflammation,

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including retinal occlusive vasculitis, following intravitreal brolucizumab treatment, which can result in permanent vision loss.¹⁰ Therefore, new therapeutic approaches, including novel agents that target other pathways involved in nvAMD that exhibit acceptable safety profiles while potentially being more effective, must be explored.

Faricimab, an innovative bispecific monoclonal antibody, has been specifically designed for intraocular use through intravitreal injection, to target two distinct pathways by binding and neutralizing both VEGF-A and angiopoietin-2 (Ang-2) simultaneously.^{11,12} Anti–Ang-2 stabilizes vessels, reduces vascular leakage, and alleviates inflammation, whereas anti–VEGF-A inhibits neovascularization in addition to reducing vascular leakage.¹¹ The efficacy of faricimab in nvAMD has been demonstrated in the pivotal Phase 3 TENAYA and LUCERNE trials.^{13,14} Data from the phase 3 global trials indicate that dosing of faricimab up to every 16 weeks (Q16W) was non-inferior to aflibercept Q8W in terms of changes in best-corrected visual acuity (BCVA) from baseline. Additionally, rates of ocular adverse events were comparable between faricimab Q16W and aflibercept Q8W.¹³ These results suggest that faricimab may overcome some of the limitations of existing anti-VEGF drugs in treating nvAMD. Therefore, real-world evidence on the use of faricimab is highly desirable. In addition, although the efficacy of anti-VEGF therapy differs among the subtypes of nvAMD¹⁵ and eyes with pachychoroid phenotypes,¹⁶ the current literature lacks data on the efficacy of faricimab in these populations.

In this study, we investigated the effectiveness and safety of three monthly intravitreal injections of faricimab in treatment-naïve patients with nvAMD. We also evaluated the differences in treatment effects by subtypes and with and without pachychoroid phenotypes.

Methods

Study Design

This prospective, non-randomised, observational study was conducted in an institutional setting. The study design was approved by the Institutional Review Board (IRB) of Kyoto University Graduate School of Medicine (R0532). All study conduct adhered to the tenets of the Declaration of Helsinki. Prior to enrolment, each patient provided written informed consent to participate in this study.

Study Population

Participants were recruited at Kyoto University Hospital between June 2022 and October 2022. The inclusion criteria were age >50 years, axial length <26.5 mm, presence of nvAMD, and willingness to participate in the study. Only one eye from each patient was included in the study and the other eye was not considered if it developed AMD. The exclusion criteria comprised previous treatment for MNV, and the presence of other retinal diseases, such as retinal vein or artery occlusion, diabetic retinopathy, angioid streaks, and vitelliform macular dystrophy. Patients with a chronic course of AMD, as indicated by disease history and/or massive fibrotic lesions, were also excluded. Those who dropped out of the study were not included in the analysis.

Intervention and Observation Procedure

All participants received 3 monthly faricimab injections (6.0 mg). We collected clinical data at baseline and 2 months after the third faricimab injection (month 4; Figure 1). If patients did not show subretinal fluid (SRF) or intraretinal fluid (IRF) at month 4, their clinical data were collected at month 5.





The patients underwent comprehensive examinations, including BCVA measurement with Landolt C charts (Takagi Seiko, Nakano, Japan),¹⁷ fundus photography, measurement of axial length (IOL Master, Carl Zeiss Meditec, Dublin, California, USA), fluorescein angiography (FA), indocyanine green angiography (IGA), fundus autofluorescence imaging (HRA2; Heidelberg Engineering), and spectral-domain optical coherence tomography (SD-OCT; Spectralis; Heidelberg Engineering, Heidelberg, Germany) at baseline. We performed fundus photography, angiography, fundus autofluorescence imaging, and SD-OCT under pupillary dilation. Visual acuity measurements and SD-OCT were also performed in month 4, and the results at baseline and month 4 were analysed. The SD-OCT images were obtained using Spectralis and Heidelberg Eye Explorer (version 1.8.6.0; Heidelberg Engineering). Thirty-degree horizontal and vertical scans through the fovea were recorded using normal and enhanced depth imaging modes, with an average of 100 scans.¹⁸

The measurements of central retinal thickness (CRT), defined as the distance between the vitreoretinal surface and the inner surface of the retinal pigment epithelium (RPE), and subfoveal choroidal thickness (SFCT), defined as the length between the outer border of the Bruch's membrane and the chorioscleral interface, were done in horizontal and vertical scans and subsequently averaged. SFCT was measured using enhanced depth imaging scans. Height of pigment epithelium detachment (PED) was defined as the distance between the outer border of the retinal pigment epithelium and the inner border of the Bruch's membrane. The subretinal hyperreflective material (SHRM) was identified in the subretinal space¹⁹ and the SHRM height was defined as the distance between the inner surface of the hyperreflective material and the inner surface of the RPE. Graders reviewed all the B scans to determine and measure the maximum PED and SHRM heights in the raster scans. Furthermore, the presence of the foveal external limiting membrane (ELM) and ellipsoid zone (EZ) were evaluated using vertical and horizontal scans. The presence of vitreoretinal adhesions was assessed using raster scanning. The presence of fluid accumulation was determined using SD-OCT at month 4. A dry macula was defined as the complete resolution of intraretinal (IRF) and/or subretinal fluid (SRF) detected on SD-OCT raster scans. The presistence of only PED was considered indicative of a dry macula.

We evaluated the presence of polypoidal lesions using indocyanine green angiography at baseline. Polypoidal choroidal vasculopathy (PCV) was diagnosed based on the finding of characteristic polypoidal lesions at the border of the branching choroidal vascular networks. Patients with signs of retinochoroidal anastomosis were diagnosed with retinal angiomatous proliferation (RAP), whereas others were diagnosed with typical AMD (tAMD).

Definition of Pachychoroid Neovasculopathy

In this study, pachychoroid neovasculopathy (PNV) was diagnosed according to the following criteria^{20,21}: (1) MNV in either eye and (2) clinical features of the pachychoroid phenotype²² such as reduced fundus tessellation in fundus photographs, choroidal vascular hyperpermeability (CVH) in IA images, and dilated choroidal vessels in SD-OCT and IA images. Dilated choroidal vessels were defined as dilated outer choroidal vessels with attenuation and thinning of the choriocapillaris on OCT.²² In IA, dilated choroidal vessels extended from one or more vortex veins;²³ (3) there was none or only non-extensive hard drusen in both eyes (Age-Related Eye Disease Study [AREDS] level 1, no AMD²⁴). Diagnoses were made by two ophthalmologists, and in case of discrepancy, a senior retinal specialist determined the final diagnosis.

Outcome Measures

Main outcome measure was a change in visual acuity. Secondary outcome measures included subgroup comparisons between patients with and without PNV and those with AMD subtypes. Additionally, we performed a multivariable regression analysis to identify baseline characteristics predicting visual outcomes.

Statistical Analysis

Statistical analysis was performed using SPSS version 21 (IBM Japan, Tokyo, Japan) with a significant *P*-value set at <0.05. Visual acuity measured with Landolt C charts was converted to the logarithm of the minimal angle of resolution (logMAR) for statistical analysis. To compare baseline characteristics, visual acuity, CRT, SFCT, PED height, and SHRM height between PNV and non-PNV, PCV and tAMD, and eyes with and without fluid at month 4, unpaired *t*-test or Fisher's exact test was applied. A paired *t*-test was used to compare baseline and month 4 values. Bivariate relationships were examined using the Pearson correlation coefficient for continuous variables and the Spearman correlation coefficient test for binary variables. Stepwise multivariable regression analysis was performed with age, presence of polypoidal

lesions, presence of IRF and/or SRH, CRT, maximum SHRM, and presence of ELM as independent factors, and visual acuity at month 4 as a dependent factor.

Results

This study included 24 patients who met the eligibility criteria with one patient dropping out after the second injection for unknown reasons. Thus, the final study sample size was 23 patients, consisting of 12 patients with tAMD, 10 with PCV, and 1 with RAP. One patient developed a retinal pigment epithelium tear as an adverse event but no other ocular complications, such as intraocular inflammation or systemic extraocular complications, were observed.

The baseline characteristics of the study participants are presented in Table 1. The mean age of patients was 75.4 ± 7.6 years and baseline BCVA was $0.29 \pm 0.30 \log$ MAR (20/39). BCVA significantly improved to $0.18 \pm 0.32 \log$ MAR (20/30) two months after the third injection (month 4) ($P = 4.9 \times 10^{-4}$, Figure 2A). Regardless of the subtype (PCV or tAMD), BCVA significantly improved (Figure 2B). In addition, significant improvements were observed in OCT parameters, where CRT and maximum SHRM height significantly decreased from $324.5 \pm 193.2 \mu$ m to $164.3 \pm 90.1 \mu$ m ($P = 1.3 \times 10^{-5}$, Figure 2C), and 159.3 ± 127.0 to 44.6 ± 132.3 ($P = 5.0 \times 10^{-6}$, Figure 2D) respectively. For the effects of treatments on the structures under the RPE, SFCT, and PED height significantly reduced from 287.8 ± 103.1 to $259.7 \pm 103.2 \mu$ m ($P = 1.1 \times 10^{-2}$, Figure 2E), and 244.1 ± 182.9 to $145.8 \pm 89.6 \mu$ m ($P = 7.2 \times 10^{-3}$, Figure 2F) respectively.

Among the 23 eyes, 11 showed pachychoroid phenotypes (PNV) and 12 did not (non-PNV). Comparing baseline characteristics between eyes with PNV and non-PNV, PNV eyes had a higher frequency of CVH (81.2% vs 8.3%, $P = 6.0 \times 10^{-4}$) and significantly greater SFCT as expected (333.1 ± 100.8 vs 246.3 ± 86.3 µm, $P = 4.6 \times 10^{-2}$) (Table 2). After 3 monthly faricimab injections, BCVA showed favourable results in both non-PNV eyes and PNV eyes at month 4 ($P = 4.1 \times 10^{-2}$ and 1.1×10^{-3} , respectively, Figure 3A). Faricimab significantly decreased SFCT in PNV (12.5% reduction in SFCT, $P = 5.0 \times 10^{-3}$; Figure 3B) and decreased CRT, PED height, and SHRM height in both non-PNV ($P = 3.0 \times 10^{-3}$, 4.1×10^{-2} , and 3.0×10^{-3} , respectively), and PNV eyes ($P = 2.0 \times 10^{-3}$, 1.8×10^{-2} , and 1.0×10^{-3} , respectively) (Figure 3C–E).

	Total Eyes (n = 23)
Age (y)	75.4±7.6
Sex (male/female)	13/10
Axial length (mm)	23.46±0.90
Subtype (tAMD/PCV/RAP)	12/10/1
LogMAR visual acuity	0.29±0.30 (20/39)
Greatest linear dimension (µm)	3439.1±1901.2
Fluid	
SRF (±)	23/0
IRF (±)	5/18
SRH (±)	9/14
CMT (µm)	324.5±193.2
Subfoveal CT (µm)	287.8±103.1
Maximum PED height (µm)	244.1±182.9
Maximum SHRM height (µm)	159.3±127.0
Vitreoretinal adhesion (±)	6/17
Intact foveal EZ (±)	11/12
Intact foveal ELM (±)	20/3

 Table I Baseline Demographic and Clinical Characteristics of

 Patients with Neovascular Age-Related Macular Degeneration

Abbreviations: tAMD, typical age-related macular degeneration; PCV, polypoidal choroidal vasculopathy; RAP, retinal angiomatous proliferation; SRF, subretinal fluid; IRF, intraretinal fluid; SRH, subretinal hemorrhage; PED, pigment epithelial detachment; SHRM, subretinal hyper-reflective material; EZ, ellipsoid zone; ELM, external limiting membrane; logMAR, logarithm of minimum angle resolution; CMT, central macular thickness; CT, choroidal thickness.



Figure 2 Visual and anatomic outcome of age-related macular degeneration (AMD) eyes treated with faricimab. (A and B) Changes in the logarithm of minimum angle resolution (logMAR) visual acuity (VA) in total eyes (A), and typical age-related macular degeneration (tAMD) and polypoidal choroidal vasculopathy (PCV) eyes (B). (C–F) Changes in central retinal thickness (CRT) (C), subfoveal choroidal thickness (SFCT) (D), maximum pigment epithelial detachments (PED) height (E), and maximum subretinal hyperreflective material (SHRM) height (F) in total eyes. *P < 0.05, ***P < 0.005, ***P < 0.001 compared between two groups (A and C–F), and between baseline and month 4 (B).

We explored predictive factors for visual outcomes. The univariate analysis showed that age, CRT, maximum SHRM height, and presence of foveal ELM, IRF, and SRH were associated with visual acuity at month 4 (Table 3). A multiple regression model analysis was performed, yielding significant overall regression (R2 = 0.788, $P < 1.0 \times 10^{-3}$), indicating that

	Pachychoroid Phenotypes (+) N = 11	Pachychoroid Phenotypes (-) N = 12	P value
Age (y)	72.9±8.2	77.8±6.2	0.140
Sex (male/female)	7/4	6/6	0.680
Axial length (mm)	23.66±0.93	23.28±0.83	0.334
Subtype (tAMD/PCV/RAP)	5/6/0	7/4/1	1.000
LogMAR visual acuity	0.23±0.18 (20/34)	0.34±0.37 (20/44)	0.423
Greatest linear dimension (µm)	2916.6±1371.2	3918.2±2173.6	0.225
CVH (%)	9 (81.2)	I (8.3)	0.0006

 Table 2 Comparison of Baseline Characteristics Between Neovascular AMD Patients with and without Pachychoroid Phenotypse

(Continued)

	Pachychoroid Phenotypes (+) N = 11	Pachychoroid Phenotypes (-) N = 12	P value
Fluid			
SRF (±)	11/0	12/0	1.000
IRF (±)	2/9	3/9	1.000
SRH (±)	4/7	5/7	1.000
CMT (μm)	271.2±97.5	373.3±240.5	0.223
Subfoveal CT (µm)	333.1±100.8	246.3±86.3	0.046
Maximum PED height (μm)	186.4±127.8	297.1±208.1	0.161
Maximum SHRM height (μm)	141.6±68.9	175.6±161.3	0.543
Vitreoretinal adhesion (±)	2/9	4/8	0.640
Intact foveal EZ (±)	7/4	4/8	0.220
Intact foveal ELM (±)	11/0	9/3	0.217

Table 2 (Continued).

Abbreviations: tAMD, typical age-related macular degeneration; PCV, polypoidal choroidal vasculopathy; RAP, retinal angiomatous proliferation; CVH, choroidal vascular hyperpermeability; SRF, subretinal fluid; IRF, intraretinal fluid; SRH, subretinal hemorrhage; PED, pigment epithelial detachment; SHRM, subretinal hyper-reflective material; EZ, ellipsoid zone; ELM, external limiting membrane; logMAR, logarithm of minimum angle resolution; CRT, central retinal thickness; CT, choroidal thickness.

78.8% of the visual outcomes could be explained by the selected factors. Among the possible predictive factors, the presence of ELM and the absence of IRF were identified as contributing factors to better visual acuity at month 4 (Table 4).

Dry macula was achieved in 77.3% of eyes at month 4 (P < 0.0001); although faricimab completely resolved SRF in most cases (from 100% to 9.1%, $P < 1.0 \times 10^{-4}$), IRF often persisted (from 21.7% to 13.6%, $P = 7.0 \times 10^{-1}$) (Figure 4A). Among eyes with dry macula at month 4, 58.8% remained dry one month later (month 5). Upon comparing eyes with and



Figure 3 Comparison of visual and anatomic outcome between age-related macular degeneration (AMD) with and without pachychoroid phenotypes. Changes in the logarithm of minimum angle resolution (logMAR) visual acuity (VA) (**A**), subfoveal choroidal thickness (SFCT) (**B**), central retinal thickness (CRT) (**C**), maximum pigment epithelial detachments (PED) height (**D**), and maximum subretinal hyperreflective material (SHRM) height (**E**) in age-related macular degeneration (AMD) with and without pachychoroid phenotypes. *P < 0.05, ***P < 0.001 compared between baseline and month 4 (**A**–**E**), and [†]P < 0.05 compared between eyes with and without pachychoroid phenotypes (**C**).

	Pearson Correlation Coefficient	P value
CRT	0.443	0.039
Maximum SHRM height	0.512	0.015
	Spearman Rank Correlation Coefficient	P value
IRF +	0.551	0.008
SRH +	0.471	0.027
Intact foveal ELM	-0.587	0.004

Table 3 Predictive Factors of Visual Acuity at Month 4 in Patients with Faricimab

Abbreviations: IRF, intraretinal fluid; SRH, subretinal hemorrhage; SHRM, subretinal hyper-reflective material; ELM, external limiting membrane; CRT, central macular thickness.

Table 4 Results of Multiple Linear Regression After Step-Wise Selection

 in Patients Treated with Faricimab

Variable	В	95% CI	β	P value
(Constant)	0.710	0.468 to 0.952	NA	7.0x10 ⁻⁶
ELM	-0.670	-0.912 to -0.429	-0.708	1.4x10 ⁻⁵
IRF	0.242	0.027 to 0.457	0.288	0.029

Abbreviations: IRF, intraretinal fluid; ELM, external limiting membrane.

without fluid at month 4, although CRT significantly decreased in both groups (Figure 4B), VA improved in eyes without fluid ($P = 5.5 \times 10^{-4}$) only and not in eyes with fluid ($P = 6.6 \times 10^{-1}$) (Figure 4C).

Discussion

In the present study, we demonstrated the effectiveness and safety of three monthly intravitreal injections of faricimab in treatment-naïve patients with nvAMD, including those with and without pachychoroid phenotypes. Our results indicated that faricimab injections improved anatomical outcomes both above and below the RPE (CRT and SHRM, SFCT and PED, respectively). Notably, faricimab was effective not only in patients with tAMD but also in PCV as well as PNV. Dry macula was achieved in 77.3% of the eyes two months after the third injection, with 58.8% of these eyes remaining dry one month later, suggesting a prolonged treatment effect after the last dose in approximately half of the eyes. Furthermore, multivariable analysis identified the presence of ELM and IRF as factors contributing to visual outcomes.



Figure 4 Comparison of visual and anatomic outcome between age-related macular degeneration (AMD) with and without dry macula at month 4. (**A**) Prevalence of dry macula, SRF, and IRF at baseline and month 4. (**B** and **C**) Changes in central retinal thickness (CRT) (**B**), and logarithm of minimum angle resolution (logMAR) visual acuity (VA) (**C**) in age-related macular degeneration (AMD) with and without dry macula at month 4. **P < 0.005, ***P < 0.001 compared between baseline and month 4 (**B** and **C**).

Our findings are consistent with the results of the pivotal phase 3 TENAYA and LUCERNE trials, which demonstrated the non-inferiority of faricimab dosed up to Q16W compared to aflibercept Q8W, with similar ocular adverse event rates, and those of the retrospective clinical studies.^{13,14,25,26} Our study extends these previous findings by examining the efficacy of faricimab in different subtypes of nvAMD, including tAMD and PCV, and PNV and non-PNV eyes. Although several studies have reported that nvAMD subtypes can affect disease management^{15,27} we found that faricimab treatment resulted in significant improvements in BCVA in both the PCV and tAMD subtypes, suggesting that faricimab may be effective in managing a range of nvAMD presentations. However, further studies with longer follow-up periods are necessary to confirm the long-term safety and efficacy of faricimab. Overall, our study provides promising evidence for the real-world application of faricimab in patients with nvAMD.

PNV is a novel clinical entity of MNV driven by pachychoroid phenotypes, which indicate choroidal manifestations, including thickened choroid and dilated outer choroidal vessels.^{16,28} The significant decrease in SFCT observed in our study, especially in PNV eyes, suggests that faricimab may effectively target the underlying choroidal abnormalities associated with pachychoroid phenotypes. This finding is essential as previous reports have shown the suboptimal efficacy of anti-VEGF agents (ranibizumab, aflibercept, and bevacizumab) for PNV treatment.²⁹ We and others have previously revealed that the molecular mechanisms involved in angiogenesis differ between PNV and non-PNV eyes.²¹ Notably, the upregulation of Ang-2 was found in the aqueous humour of treatment-naïve patients with PNV.³⁰ The bispecific action of faricimab, which targets both VEGF-A and Ang-2, may therefore be crucial in addressing the complex pathophysiology of PNV. Further, though targeting both VEGF and Ang-2 by Faricimab may offer enhanced efficacy for PNV compared to other anti-VEGF agents, it is important to note that our current study did not include a direct comparison between Faricimab and other anti-VEGF treatments. Because our study's scope was limited to its efficacy within the context of PNV without comparative analysis to other treatments, future studies specifically designed to compare Faricimab with other anti-VEGF agents would be invaluable to substantiate this hypothesis and provide more definitive conclusions.

Our multivariable analysis identified the presence of ELM and IRF at baseline emerged as factors contributing to visual acuity at month 4. These findings align with those of previous studies that have reported the presence of ELM as a predictor of better visual outcomes following anti-VEGF therapy.³¹ The association between IRF presence before treatments and poorer visual outcomes underscores the importance of addressing fluid accumulation in the management of patients with nvAMD. The reasons for the potential negative impact of IRF on BCVA are not well understood, and it has been suggested that IRF may be indicative of a more aggressive lesion type or a late sign of the chronic stage of MNV.³² Despite a significant reduction in SRF following faricimab treatment as demonstrated in our study, IRF often persisted, highlighting the need for further investigation into the management of intraretinal fluid in patients with nvAMD.

A key strength of our study lies in its evaluation of faricimab treatment in a real-world setting, which complements the findings from the TENAYA and LUCERNE trials. Additionally, the inclusion of nvAMD subtypes and investigation of treatment effects in eyes with and without PNV provide valuable insights into the potential benefits of faricimab across different patient populations. However, our study has several limitations, including a small sample size and a relatively short follow-up period. We understand that the small sample size may impact the generalizability of our findings, but our results can be a valuable preliminary indicator and a basis for future studies. We believe that our findings contribute to the field, offering initial insights that can be further explored in subsequent research. Future studies with larger sample sizes and longer follow-up periods are needed to provide further evidence of the long-term efficacy and safety of faricimab in patients with nvAMD.

In conclusion, our findings provide compelling evidence supporting the effectiveness and safety of faricimab in treatment-naïve patients with nvAMD. The favourable outcomes observed in both PNV and non-PNV eyes suggest their potential as promising therapeutic options for a wide range of patients with nvAMD. Half of the treated eyes retained dry macula 3 months after the last dose, indicating the possibility of extending the dosing interval compared to conventional anti-VEGF agents, such as ranibizumab and aflibercept. However, further comparative studies are needed to confirm the long-term benefits of faricimab and optimise treatment regimens tailored to individual patients.

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