



# Comparative Efficacy of Brolucizumab in the Treatment of Neovascular Age-Related Macular Degeneration: A Systematic Literature Review and Network Meta-Analysis

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## ABSTRACT

**Introduction:** A systematic literature review (SLR) and network meta-analysis (NMA) were conducted to evaluate the comparative efficacy of brolucizumab relative to other anti-vascular endothelial growth factor (VEGF) treatments for neovascular age-related macular degeneration (nAMD) at 1 and 2 years, and overall safety and injection frequency of each treatment.

**Methods:** An SLR identifying randomized controlled trials (RCTs) published before June 2021 according to a pre-specified protocol was followed by a Bayesian NMA to compare brolucizumab (6 mg q12w/q8w) against sham and all relevant anti-VEGF regimens. Pooled mean injection frequency, serious adverse ocular events, and discontinuation rates were estimated for each treatment regimen.

**Results:** Nineteen RCTs were included in NMA base-case analysis. Brolucizumab (6 mg q12w/

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q8w) with loading-phase (LP) demonstrated superior best-corrected visual acuity (BCVA) gains to sham both at year 1 (mean difference 16.8 [95%CrI 13.3, 20.4]) and year 2 (mean difference 21.2 [95%CrI 17.4, 25.0]) and was comparable to other anti-VEGFs. Brolucizumab (6 mg q12w/q8w) also showed superior retinal thickness reduction to most comparators including ranibizumab (0.5 mg q4w; year 1 mean difference - 50.1 [95%CrI - 70.3, - 29.8]; year 2 mean difference - 49.5 [95%CrI - 70.8, - 28.6]), aflibercept (2 mg q8w; year 1 mean difference - 39.7 [95%CrI - 52.9, - 26.4]; year 2 mean difference - 35.0 [95%CrI - 49.1, - 21.4]), and faricimab (6 mg q16w/q8w; year 1 mean difference - 27.6 [95%CrI - 42.3, - 12.8]). Brolucizumab (6 mg q12w/q8w) showed similar rates of treatment discontinuation and serious and overall adverse events (both years). At year 2, pooled annualized injection frequency was lowest for brolucizumab (6 mg q12w/q8w) and highest for ranibizumab (0.5 mg q4w) at 5.7 and 11.5 injections annually, respectively.

**Conclusion:** Among all licensed anti-VEGF treatments, brolucizumab showed superior reduction in retinal thickness and comparable BCVA gains and discontinuation rates, despite having the lowest injection frequency. The current study provides the most up-to-date, robust comparison of treatments for nAMD.

**Keywords:** Brolucizumab; Neovascular age-related macular degeneration; nAMD; Network meta-analysis; NMA; Anti-VEGF

## Key Summary Points

### *Why carry out this study?*

Current pharmacologic therapy for neovascular age-related macular degeneration (nAMD) involves intravitreal injections of anti-vascular endothelial growth factor (anti-VEGF) agents.

We conducted a network meta-analysis (NMA) comparing the efficacy and safety of brolucizumab to other anti-VEGF treatments in patients with nAMD.

By combining direct and indirect evidence, NMA makes it possible to simultaneously compare multiple treatments that were not compared in head-to-head trials.

### *What was learned from the study?*

Among all licensed anti-VEGF treatments, brolucizumab showed superior reduction in retinal thickness and comparable visual acuity gains and discontinuation rates, despite having the lowest injection frequency.

The current NMA provides the most up-to-date, robust comparison of treatments for nAMD.

## INTRODUCTION

Age-related macular degeneration (AMD) is the main cause of severe visual loss and blindness in all high-income countries [1]. AMD progresses slowly over on average 10 years from early to intermediate to late stage disease which can include dry, or non-exudative, late-stage AMD and wet, or neovascular, AMD as well as combinations thereof [2, 3]. Neovascular AMD

(nAMD) occurs in about two-thirds of late-stage AMD cases, develops rapidly, and commonly results in severe vision loss if left untreated.

Current pharmacologic therapy for nAMD involves intravitreal injections of anti-vascular endothelial growth factor (anti-VEGF) agents. Anti-VEGF monotherapy is the preferred treatment for nAMD with prompt intervention limiting visual loss in 95% of patients with nAMD [4, 5]. Patients with nAMD demonstrate high individual variability in the need for anti-VEGF injections. Delay in initiating treatment as well as undertreatment caused by poor adherence and/or persistence leads to worsening treatment outcomes over time [6–8]. Differing treatment regimens have been shown to impact treatment adherence and persistence, with treat and extend or comparable regimens fairing best. Thus, further research on the comparison between existing treatment regimens for nAMD is necessary.

Brolucizumab is a humanized single-chain antibody fragment that inhibits all isoforms of VEGF-A. Owing to its small size, brolucizumab demonstrates good tissue penetration in the eye, reaching high concentrations in the retina [9]. In two pivotal phase III trials, HAWK and HARRIER, brolucizumab demonstrated non-inferiority to aflibercept in the primary endpoint of change in best-corrected visual acuity (BCVA) in the treatment of nAMD [10]. Moreover, among key secondary endpoints of disease activity, brolucizumab was superior to aflibercept. These include reduction in retinal thickness (central subfield thickness) as well as retinal fluid markers (subretinal and intraretinal fluids), which are anatomical outcomes used to determine treatment efficacy and injection frequency in clinical practice [10]. Brolucizumab has been associated with uncommon events of retinal vasculitis and retinal vascular occlusion in patients with nAMD [11].

HAWK and HARRIER provided insight into the relative efficacy and safety of brolucizumab versus aflibercept. However, indirect comparisons of brolucizumab with other anti-VEGFs are needed.

The objective of the present study was to conduct a network meta-analysis (NMA) comparing the efficacy and safety of brolucizumab

to other approved and marketed anti-VEGF treatments in patients with nAMD. By combining direct and indirect evidence, NMA makes it possible to simultaneously compare multiple treatments that were not compared in head-to-head trials.

## METHODS

### Search Strategy

A systematic literature review of studies comparing anti-VEGFs in nAMD was conducted using Embase, Medline, Medline-in-Process, and Cochrane Library from database inception to June 3, 2021. The search strategy included English-language publications on clinical trials that evaluated treatments for patients with nAMD. The following treatments were included in the search strategy: brolucizumab, ranibizumab, aflibercept, faricimab, pegaptanib, bevacizumab, photodynamic therapy with verteporfin, laser photocoagulation therapy, and macular surgeries. The NMA only included licensed anti-VEGF therapies. While faricimab was not licensed when we conducted the study, it was included because Roche had filed an application in the European Union, and submitted to the National Institute for Health and Care Excellence (NICE) [12, 13]. Faricimab has been recently approved by the US Food and Drug Administration (FDA) [14]. The search strategy was not restricted to specific outcomes, as this was evaluated during the screening phase of the literature review. The detailed search strategy is available in the Supplementary Material.

In addition, relevant abstracts from 2015 to 2021 were searched from the following congresses: American Society of Retina Specialist, The American Macular Degeneration Foundation, European Society of Retina Specialists, The Retina International World Congress of Ophthalmology, The Association for Research and Vision in Ophthalmology, American Academy of Ophthalmology, The Royal Australian and New Zealand College of Ophthalmologists, Asia-Australia Controversies in Ophthalmology, and The Royal College of Ophthalmologists.

**Table 1** PICOS framework for the network meta-analysis

Topic	Description
Population	Patients $\geq$ 18 years old with nAMD (also known as wet AMD)
Intervention	Brolucizumab
Comparators	Ranibizumab (Lucentis <sup>®</sup> ) Aflibercept (Eylea <sup>®</sup> ) Faricimab (RO6867461)
Outcomes	Mean change in BCVA Mean change in retinal thickness Proportion of patients gaining at least 15 ETDRS letters Proportion of patients losing at least 15 ETDRS letters Discontinuation Injection frequency <sup>a</sup> Adverse events <sup>a</sup>
Study type	RCTs of 44 weeks or longer, crossover RCTs (if data presented at the time of crossover) Open-label extension studies of RCTs

nAMD neovascular age-related macular degeneration, AMD age-related macular degeneration, ETDRS: Early Treatment Diabetic Retinopathy Study, RCT randomized controlled trial

<sup>a</sup>Only the pooled absolute treatment effects were estimated for these outcomes

The US National Library of Medicine (ClinicalTrials.gov) and the EU Clinical Trials Register were also searched for any additional studies of interest. This article is based on previously conducted studies and does not contain any new studies with human participants or animals performed by any of the authors.

### Study Population and Selection Criteria

The selection criteria for the NMA are presented in Table 1. In terms of participants, adults with

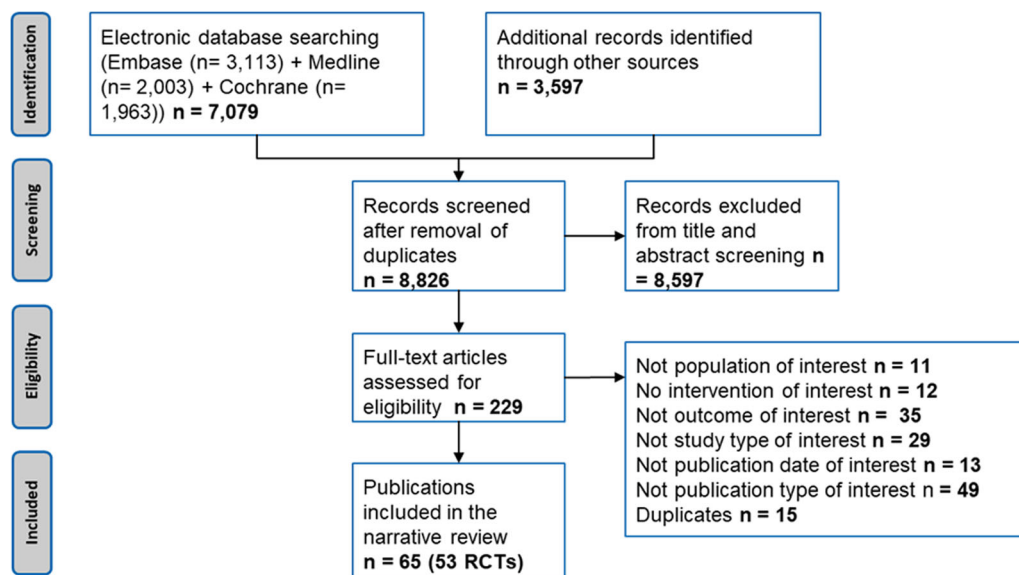
a primary diagnosis of nAMD were included. Studies where patients with polypoidal choroidal vasculopathy (PCV) made up more than 10% of the population were excluded, as patients with PCV present different demographic and clinical characteristics than the overall nAMD population. Randomized controlled trials (RCTs) of 44 weeks or longer, crossover RCTs, and open-label extension studies of RCTs were also considered. In terms of treatments, licensed doses of brolucizumab, ranibizumab, and aflibercept were included in the NMA. As faricimab was not yet licensed at the time of analysis, all doses and regimens in its clinical trials were considered. Acknowledging unlicensed bevacizumab is reimbursed in some countries under specific protocols, a sensitivity analysis including bevacizumab is reported in Supplementary Material. Each treatment was evaluated separately by its administration regimen, including if a loading phase was used. The efficacy outcomes of interest included mean change in BCVA, proportion of patients gaining at least 15 Early Treatment Diabetic Retinopathy Study (ETDRS) letters, proportion of patients losing at least 15 ETDRS letters, and mean change in retinal thickness. The safety outcomes included discontinuation, serious adverse events (SAE), including intraocular inflammation, endophthalmitis, retinal detachment, retinal tear, retinal pigment epithelial tear, cataract, and stroke. Additionally, the mean number of injections was evaluated.

### Study Screening

Titles and abstracts of all citations were screened by two independent reviewers (ND, AQ), and discrepancies resolved by a third independent reviewer (CM). The full texts of relevant publications were screened using the same process.

### Data Extraction and Quality Assessment

Two independent researchers (ND, AQ) extracted data from the identified publications into a prepared data extraction form. A third independent reviewer (CM) performed a quality



**Fig. 1** PRISMA diagram of the studies included in the systematic literature review

check of all extractions. The quality and risk of bias of each included study were assessed using the Cochrane Collaboration’s quality tool [15].

**Statistical Analysis**

Heterogeneity was assessed for each direct comparison using pairwise meta-analyses when at least two studies reported the same treatment comparison. The Cochran’s Q test was conducted and the  $I^2$  statistic was calculated; if the  $p$  value of the Cochran’s Q test was less than 0.10 or the  $I^2$  statistic was greater than 50%, then heterogeneity was suspected [16]. An inconsistency assessment using the Bucher approach was conducted for each closed loop in the networks with at least two trials. For each outcome, the indirect comparison based on the Bucher approach was compared to a direct pairwise meta-analysis [17]. The difference of the point estimates obtained from indirect versus direct evidence was calculated and a z-score was obtained. If the  $p$  value was lower than 0.05, then inconsistency was considered to be present.

The NMAs were conducted using a Bayesian framework in accordance with NICE Decision Support Unit (DSU) guidelines, and also accepted by most health technology appraisal bodies

[18, 19]. The estimates were calculated using Monte Carlo Markov chains (MCMCs). Two MCMCs were used with 20,000 iterations and a burn-in of 20,000 for fixed-effects models; for random-effects models, 100,000 iterations were used with a burn-in of 100,000. The model with the lowest deviance information criterion (DIC) was selected. If the DIC of the random-effects model was less than 3 units lower than that of the fixed-effects model, then the fixed-effects model was chosen.

For the continuous outcomes, mean change in BCVA and mean change in retinal thickness, the posterior mean change from baseline was calculated along with 95% credible intervals (95%CrI). For the categorical outcomes, patients losing or gaining at least 15 letters and discontinuation, the odds ratio was reported along with the 95%CrI. Finally, for serious adverse events and injection frequency, the pooled absolute treatment effects were calculated using inverse variance weighting and random-effect models [20]. Serious adverse events were pooled by treatment molecule while injection frequency was pooled by treatment regimen.

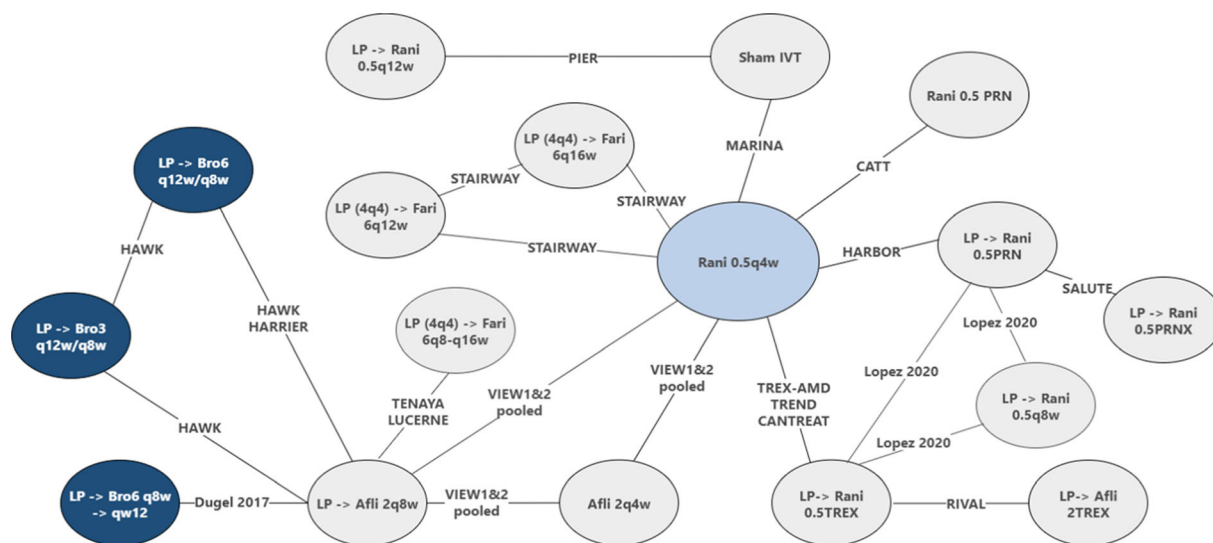
**Table 2** Details of trials included in the network meta-analysis

Trial #	Author, year	Trial name	Time of assessment (months)	Previous anti-VEGF therapy	Intervention	Comparator
1	Martin 2011/Martin 2012	CATT	12/24	Treatment-naive	Rani 0.5q4	Rani 0.5PRN
2	Dugel 2017	NR	12	Treatment-naive	LP → Bro 6q8 → q12	LP → Afi 2q8
3	Dugel 2018/2019	HARRIER	48 weeks/ 96 weeks	Treatment-naive	LP → Bro 6q12/q8	LP → Afi 2q8
4	Dugel 2018/2019	HAWK	48 weeks/ 96 weeks	Treatment-naive	LP → Bro 3q12/q8 LP → Bro 6q12/q8	LP → Afi 2q8
5	Eldem 2015	SALUTE	12	Treatment-naive	LP → Rani 0.5PRNX	LP → Rani 0.5PRN
6	Feltgen 2017	RABIMO	12	Treatment-naive	LP → Rani 0.5q8	LP → Rani 0.5PRN
7	Hunyor 2018/Gillies 2019 (1)/Gillies 2019 (2)	RIVAL	12/24	Treatment-naive	LP → Rani 0.5TREX	LP → Afi 2TREX
8–9	Schmidt-Erfurth 2014	VIEW 1&2 Pooled	12/96 weeks	Treatment-naive	Afi 2q4 → PRN LP → Afi 2q8 → PRN	Rani 0.5q4 → PRN
10	Ho 2014	HARBOR	12/24	Treatment-naive	Rani 0.5q4	LP → Rani 0.5PRN
11	Kertes 2019/Kertes 2020	CAN-TREAT	12	Treatment-naive	LP → Rani 0.5TREX	Rani 0.5q4
12	Lopez 2020	NR	12	Treatment-naive	LP → Rani 0.5TREX LP → Rani 0.5PRN	LP → Rani 0.5q8
13	Regillo 2008	PIER	12	Not reported	LP → Rani 0.5q12	Sham injection
14	Rosenfeld 2006	MARINA	12/24	Not reported	Rani 0.5q4	Sham injection
15	Silva 2017	TREND	12	Treatment-naive	LP → Rani 0.5TREX	Rani 0.5q4

**Table 2** continued

Trial #	Author, year	Trial name	Time of assessment (months)	Previous anti-VEGF therapy	Intervention	Comparator
16	Wykoff 2015/Wykoff 2017	TREX-AMD	12/24	Treatment-naive	LP → Rani 0.5TREX	Rani 0.5q4
17	Heier 2021	TENAYA	48 weeks	Treatment-naive	LP (4q4) → Fari 6q8-q16	LP → Afi 2q8
18	Heier 2021	LUCERNE	48 weeks	Treatment-naive	LP (4q4) → Fari 6q8-q16	LP → Afi 2q8
19	Khanani 2020	STAIRWAY 12	12	Treatment-naive	LP (4q4) → Fari 6q12 LP (4q4) → Fari 6q16	Rani 0.5q4

Rani ranibizumab; Bro brolocizumab; Afi aflibercept; Fari faricimab; LP loading phase of 3 monthly injections, LP (4q4) loading phase of 4 monthly injections; PRN pro re nata or as needed; PRNX as needed with the potential to extend the treatment intervals; TREX treat and extend; q4, q8, q12, q16 injections administered every 4, 8, 12, and 16 weeks, respectively



**Fig. 2** Network for mean change in BCVA at 1 year. Rani ranibizumab; Bro brolocizumab; Afi aflibercept; Fari faricimab; LP loading phase of 3 monthly injections, LP (4q4) loading phase of 4 monthly injections; PRN pro re nata or as needed; PRNX as needed with the potential to

extend the treatment intervals; TREX treat and extend; q4, q8, q12, q16 injections administered every 4, 8, 12, and 16 weeks, respectively

## RESULTS

### Systematic Literature Review

A total of 10,676 citations were captured from the electronic database searches. After removal of duplicates, 8826 citations remained. The screening of these titles and abstracts led to the review of 229 publications to assess their eligibility for inclusion. On the basis of the selection criteria, 65 publications were included, and their data was extracted in the narrative synthesis, as shown in the PRISMA diagram (Fig. 1). These publications encompassed 53 clinical trials. After applying the PICOS (Population, Intervention, Comparators, Outcomes Study design) criteria for inclusion in the NMA, a total of 19 trials were considered relevant for the NMA (Table 1). The details of the included trials are reported in Table 2, and the network for the primary outcome of interest, the mean change in BCVA, is represented in Fig. 2. For all other outcomes, the networks are reported in the Supplementary Material.

The evidence provided by these 19 trials were overall of high quality. The greatest risk of bias was due to insufficient information identified in the publication during the quality assessment. Most studies did not report unexpected imbalances in dropouts between groups.

An additional ten trials were included in the sensitivity analyses with unlicensed bevacizumab. The details of the included trials are reported in Supplementary Material.

### Characteristics and Comparability of Included Studies

Similar eligibility criteria were observed amongst the included trials. Nearly all trials included treatment-naive patients only, with the exception of MARINA and PIER. While MARINA and PIER did not report any inclusion or exclusion criteria related to prior anti-VEGF therapy, these trials were conducted when pegaptanib was the only authorized anti-VEGF agent. Given the lack of other authorized anti-VEGF treatments at the time, patients enrolled

**Fig. 3** Forest plot of results obtained through the network meta-analysis for **a** mean change in BCVA, **b** mean change in retinal thickness, **c** patients losing at least 15 letters, **d** patients gaining at least 15 letters, **e** overall discontinuation at 1 year

in these trials were likely to be predominantly treatment naive.

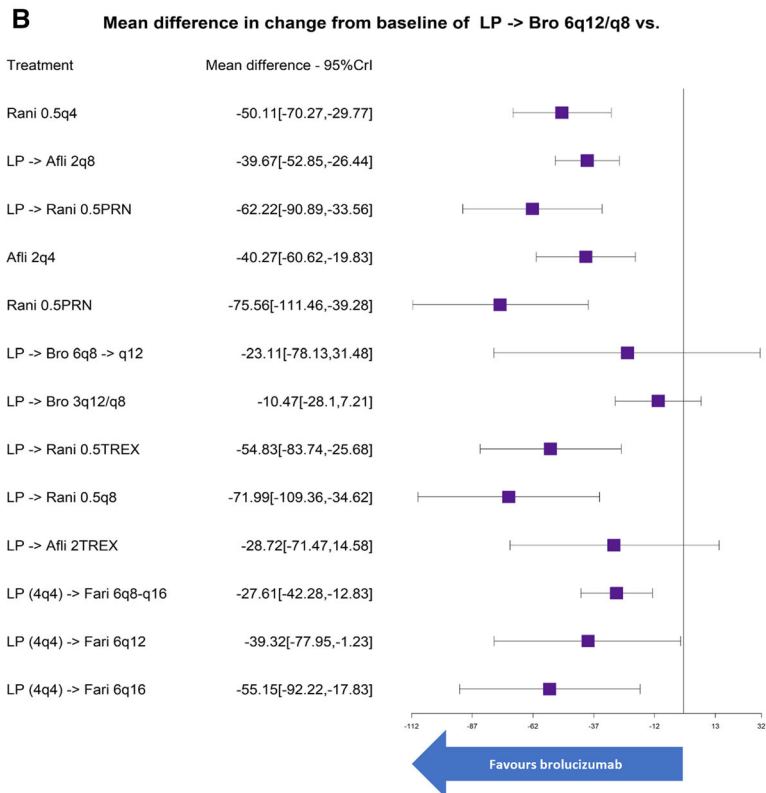
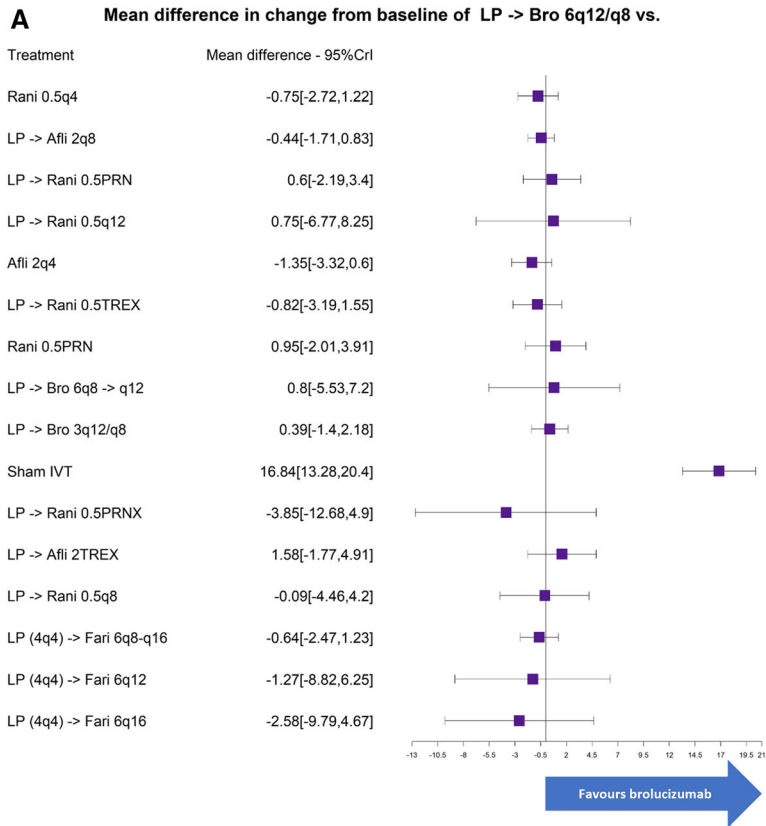
The average age at baseline varied between 65 years (TRESX-AMD: LP → Rani 0.5TRESX) and 81 years (RABIMO: LP → Rani 0.5PRN), with most trials reporting a mean age between 75 and 78 years. The proportion of male patients ranged from 30% (RABIMO: LP → Rani 0.5q8) to 56% (SALUTE: LP → Rani 0.5PRN). The mean BCVA at baseline was generally higher amongst the more recent studies, but this did not vary considerably. The lowest mean BCVA at baseline was reported for the sham injection arm of MARINA, at 53.6 ETDRS letters. The highest mean BCVA at baseline was reported for the LP → Rani 0.5TRESX arm of RIVAL, at 65.3 letters.

In the included studies there were 11 different anti-VEGF regimens considered: injections administered on a fixed schedule every 4, 8, 12, or 16 weeks (q4, q8, q12, and q16), treatment administered as needed (PRN), PRN with the potential to extend the assessment interval (PRNX), treat with the potential to extend the treatment interval when no signs of exudation are present (TRESX), injections every 8 weeks until week 40 then every 12 weeks until week 56 (q8 → q12), injections every 12 weeks unless there were signs of disease progression in which case the patient switched to bimonthly injections (q12/q8), and injections every 8 or 16 weeks on the basis of disease activity and assessment at weeks 20 and 24 (q8–q16). Some of the regimens included a loading phase of three initial monthly injections (LP). The faricimab trials, however, included a loading phase of four initial monthly injections (LP (4q4)).

### Network Meta-Analysis

On the basis of convergence and DIC values, the fixed-effects model was chosen for each outcome. Among the closed loops, no statistical





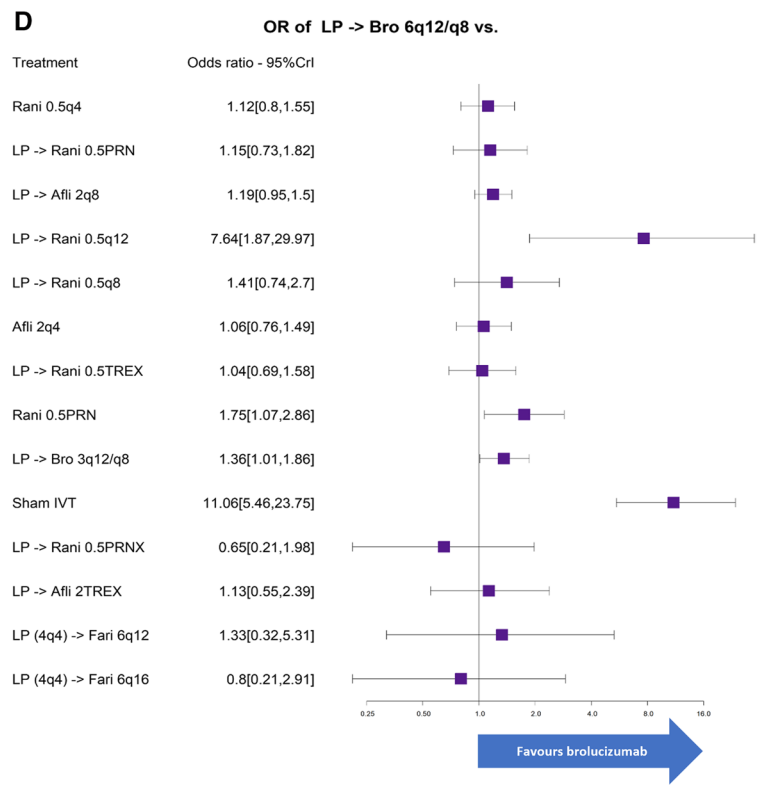
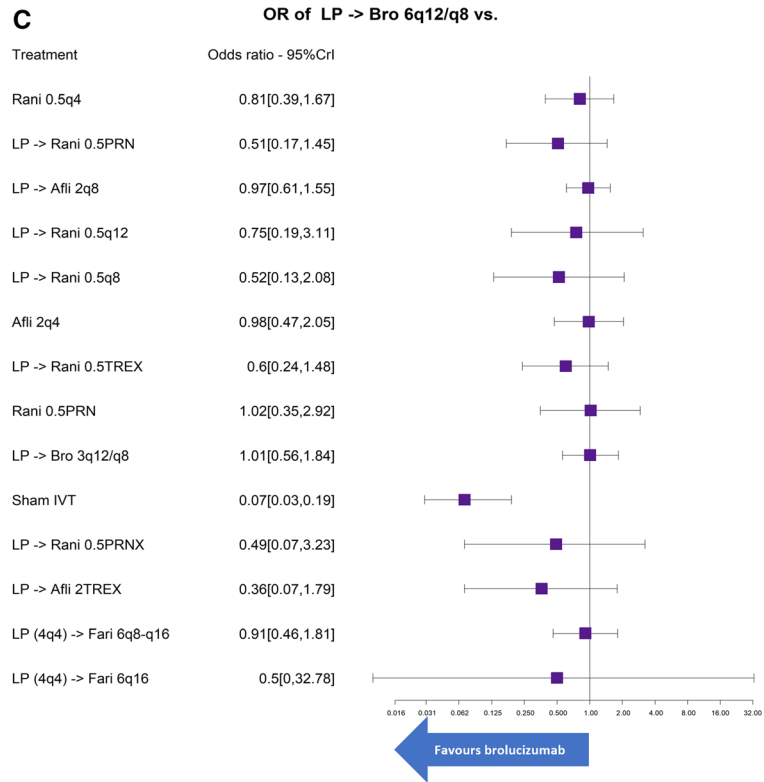


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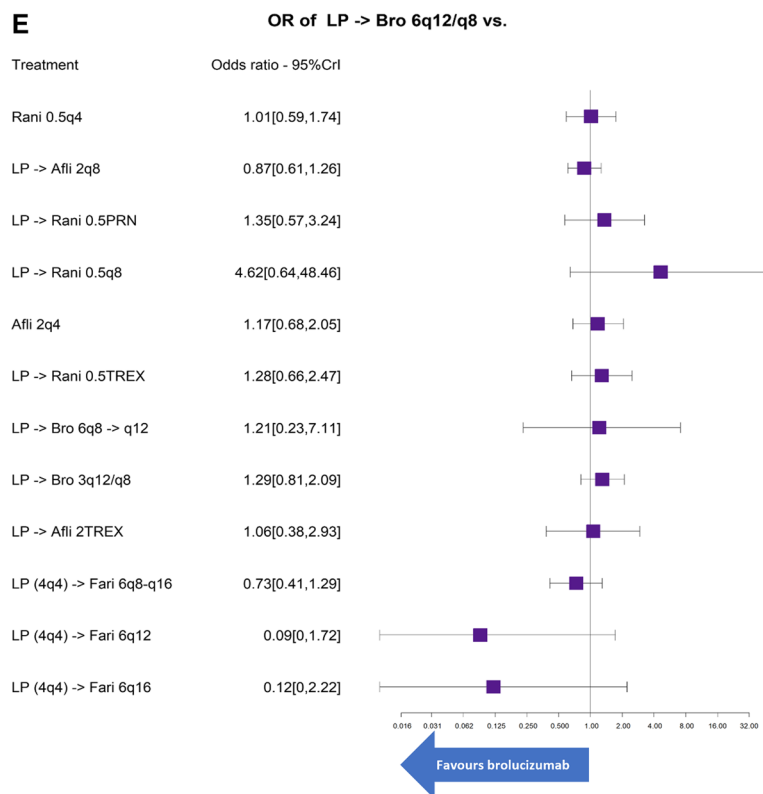


Fig. 3 continued

evidence of inconsistency was identified (Supplementary Material). The treatment comparisons for which significant heterogeneity was identified for mean change in BCVA at 1 year were LP → Rani 0.5TRES versus Rani 0.5q4, Rani 0.5q4 versus Afli 2q4, and LP → Afli 2q8 vs. Afli 2q4. In the first comparison, baseline characteristics were assessed and were similar across the three trials. This heterogeneity is therefore likely due to the inherent variability of the follow-up treatment intervals in the TRES regimen and cannot be controlled for. The second two comparisons were between similarly designed trials, VIEW 1 and VIEW 2, which used the same inclusion and exclusion criteria. The main difference between these studies was the geographical area considered, suggesting that the heterogeneity in results is due to random variability. These trials were therefore pooled in the NMA and considered one trial. This is the same approach that NICE used in their NMA in nAMD [21]. Detailed

information from the heterogeneity assessment can be found in the Supplementary Material.

The relative efficacy and safety results for the brolocizumab 6 mg arm evaluated in HAWK and HARRIER (LP → Bro 6q12/q8) for each of the NMAs conducted at 1 and 2 years are presented in Figs. 3 and 4, respectively. The surface under the cumulative ranking curve (SUCRA) and probability for LP → Bro 6q12/q8 to perform better than each of its comparators are reported in Table 3. Compared to the sham injection, LP → Bro 6q12/q8 had a higher mean change in BCVA at both 1 and 2 years. None of the other relative treatments had significant effects based on the 95%CrI. LP → Bro 6q12/q8 had greater than a 50% probability of performing better than seven out of 16 of its comparators in terms of increasing BCVA at 1 year.

Brolocizumab 6 mg also had a significantly greater odds of gaining at least 15 letters than sham injections and a high probability of performing better than 12 out of 14 and eight out

of nine of its comparators at 1 and 2 years, respectively. Additionally, LP → Bro 6q12/q8 had the third highest SUCRA for patients gaining at least 15 letters at 1 year, with a value of 73.7%. Similar results were obtained for the odds of losing at least 15 letters, with LP → Bro 6q12/q8 having a greater than 50% probability of performing better than 12 out of 15 of its competitors at 1 year.

A significantly greater decrease in mean retinal thickness was observed for brolocizumab 6 mg when compared to most of the competitors at both 1 and 2 years (10 out of 13 at 1 year, and 5 out of 8 at 2 years), with LP → Bro 6q12/q8 having the highest mean rank in SUCRA value at 1 year with 96.6%. Each treatment had similar odds of discontinuation, with no significant differences in relative treatment effects based on the 95%CrI.

### Adverse Events and Injection Frequency

This NMA includes SAEs as reported in the original pivotal trials. Information for faricimab is limited at this stage, and isolated trial results do not reflect the latest brolocizumab safety information available. Results from the molecule-based pooled absolute treatment are briefly reported below, and fully included in the Supplementary Material for completion.

Endophthalmitis rates ranged from 0.31% for brolocizumab to 0.50% for ranibizumab at 1 year and from 0.54% for aflibercept to 0.94% for ranibizumab at 2 years. All anti-VEGF treatments reported similar rates of retinal detachment at 1 year, 0.30%, and ranging between 0.30% and 0.49% at 2 years. Rates of retinal pigment epithelial tear ranged between 0.30% and 1.10% at 1 year and between 0.20% and 0.40% at 2 years. Retinal tear was reported among 0.30% of patients receiving ranibizumab and none receiving aflibercept at 1 year. At 2 years, these rates ranged between 0.30% and 0.50%. Finally, at 1 year, stroke was reported by between 0.42% and 0.63% of patients; at 2 years, these rates ranged between 0.63% and 1.04%.

The results for the injection frequency at 1 year indicated that Afli 2q4 had the highest

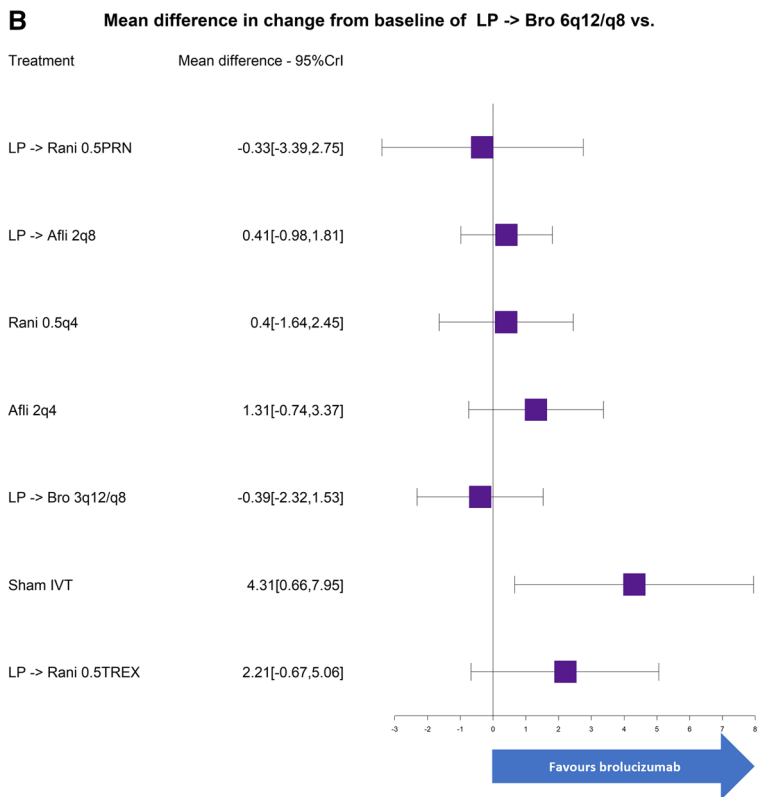
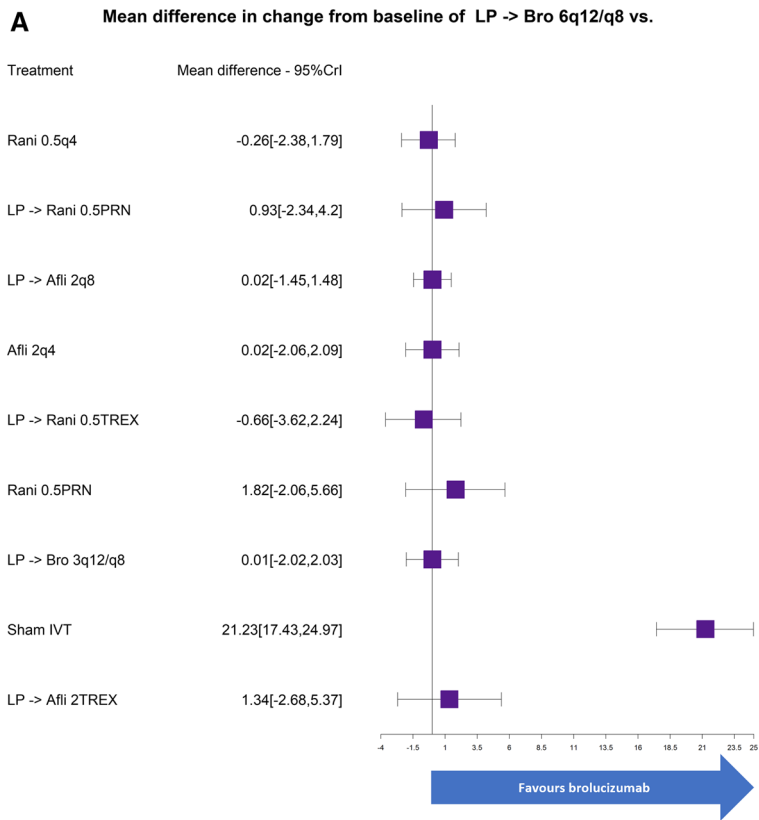
**Fig. 4** Forest plot of results obtained through the network meta-analysis for **a** mean change in BCVA at 2 years, **b** mean change in BCVA from 1 to 2 years, **c** mean change in retinal thickness at 2 years, **d** patients losing at least 15 letters at 2 years, **e** patients gaining at least 15 letters at 2 years, **f** overall discontinuation at 2 years

injection frequency and LP → Rani 0.5PRNX had the lowest. At 2 years, Rani 0.5q4 presented the highest injection frequency and the two brolocizumab regimens presented the lowest annualised injection frequency (Table 4). Fixed monthly injections are expected to have the highest mean number of injections based on their protocol for treatment administration.

## DISCUSSION AND CONCLUSION

This study is the first of its kind in indirectly comparing the efficacy of brolocizumab and faricimab to all comparators for the treatment of nAMD. Across the board, the quality of evidence included in the NMA was high with mostly low bias risk. Among all anti-VEGF treatments, the visual acuity outcomes were similar, which is consistent with previous meta-analyses conducted in nAMD [21–23]. Many of the studies included ranibizumab and aflibercept as active comparators. However, ranibizumab is often administered on a monthly schedule, whereas newer treatments have less frequent or more flexible dosing regimens based on disease activity. In particular, when compared to the faricimab arm evaluated in TENAYA and LUCERNE at 1 year, brolocizumab had a lower number of injections, at 6.7 compared to 7.0. This is likely in part due to the loading phase consisting of intravitreal injections every 4 weeks for the first four doses required in the faricimab trials.

Brolocizumab also showed greater reductions in retinal thickness than its comparators. Retinal thickness is a common anatomical measure of disease activity in nAMD, and studies have shown that greater thickness may be associated with worse visual acuity outcomes [24, 25]. These measurements also play an important role in determining dosing intervals for variable



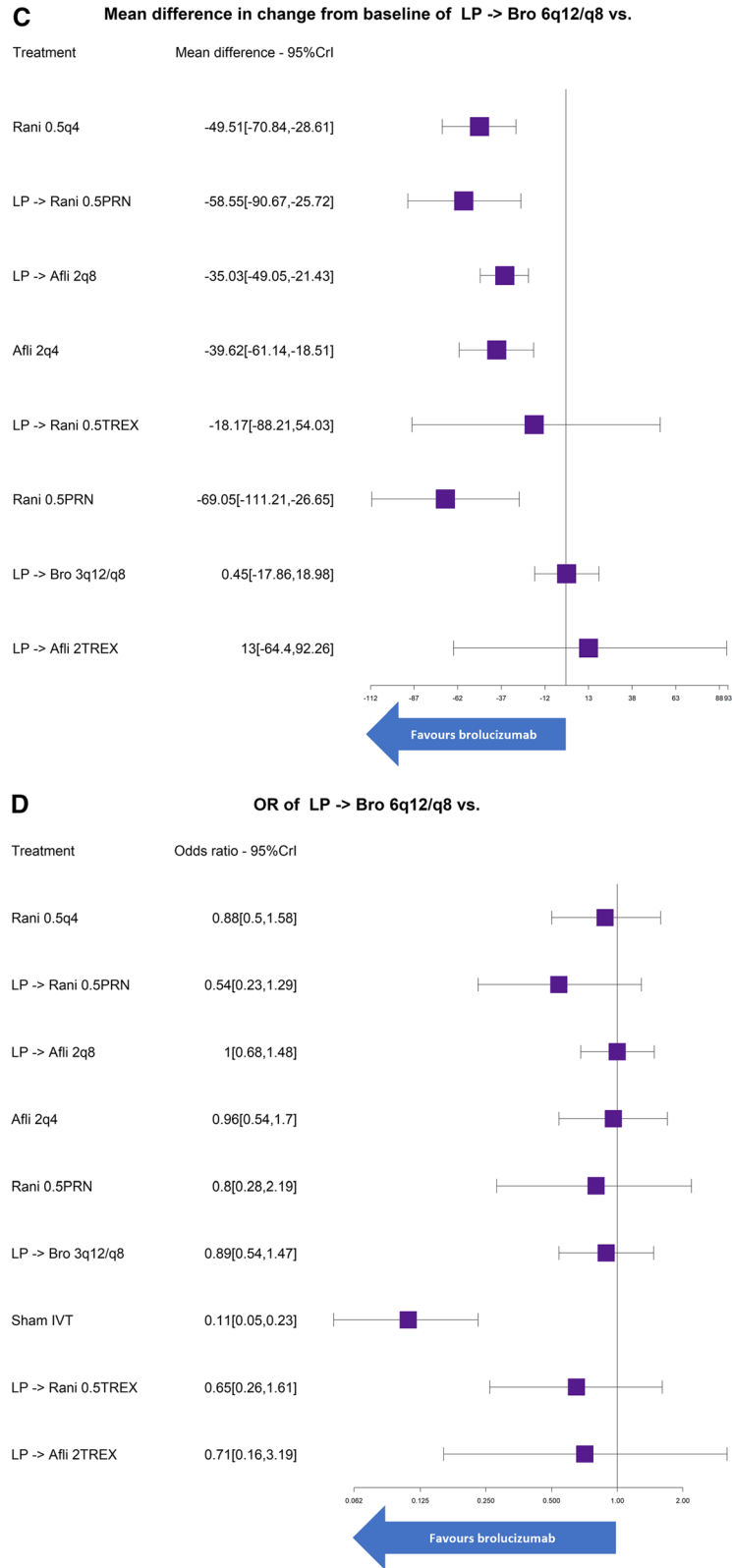


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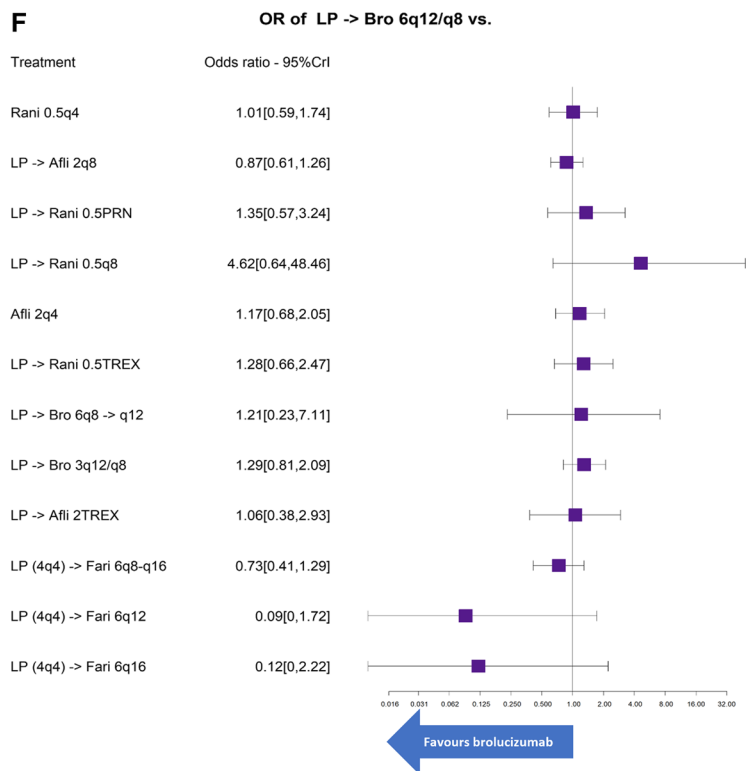
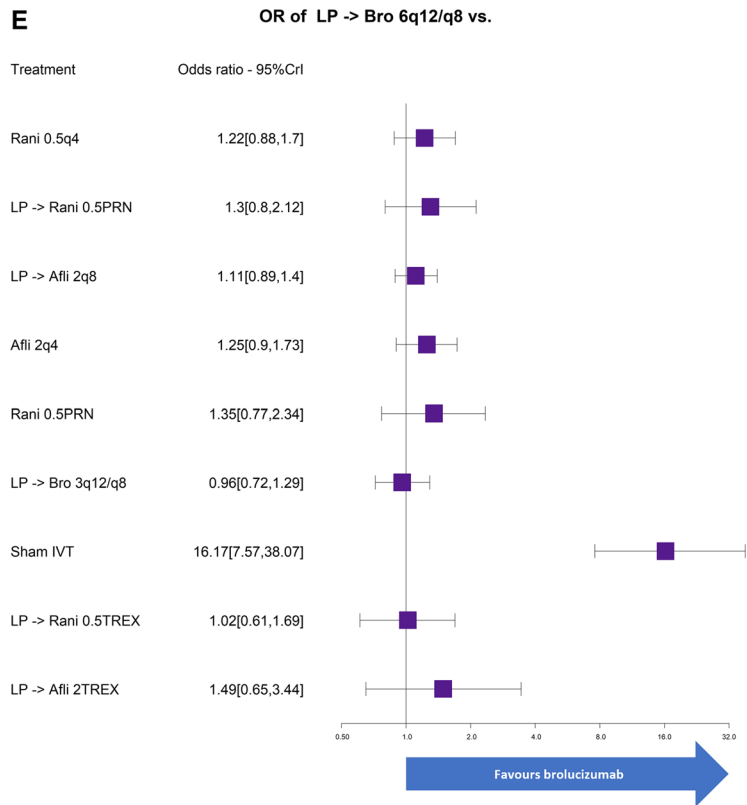


Fig. 4 continued

**Table 3** Cumulative surface under the cumulative ranking curves (SUCRA) and probabilities for LP → Bro 6q12/q8 to perform better than each comparator for all outcomes at 1 and 2 years

Treatment	SUCRA	Probability for LP → Bro 6q12/q8 to perform better than each comparator	SUCRA	Probability for LP → Bro 6q12/q8 to perform better than each comparator
Mean change in BCVA	1 year		2 years	
Rani 0.5q4	63.9%	22.7%	70.8%	40.5%
LP → Aflī 2q8	56.6%	24.9%	60.6%	50.8%
LP → Rani 0.5PRN	36.3%	66.5%	41.1%	71.6%
LP → Rani 0.5q12	43.5%	57.6%	–	–
Aflī 2q4	76.0%	8.7%	60.7%	50.9%
LP → Rani 0.5TREX	65.0%	24.8%	78.2%	32.7%
Rani 0.5PRN	31.7%	73.4%	29.5%	82.1%
LP → Bro 6q8 → q12	41.8%	59.7%	–	–
LP → Bro 6q12/q8	46.4%	–	61.4%	–
LP → Bro 3q12/q8	39.3%	66.6%	61.0%	50.2%
Sham injection	0.0%	100%	0.0%	100%
LP → Rani 0.5PRNX	79.2%	19.2%	–	–
LP → Aflī 2TREX	24.5%	82.5%	36.6%	74.2%
LP → Rani 0.5q8	50.4%	48.5%	–	–
LP (4q4) → Fari 6q8-q16	60.9%	25.0%	–	–
LP (4q4) → Fari 6q12	61.2%	37.0%	–	–
LP (4q4) → Fari 6q16	73.3%	24.6%	–	–
Mean change in retinal thickness	1 year		2 years	
Rani 0.5q4	37.0%	100%	25.2%	100%
LP → Aflī 2q8	54.5%	100%	49.8%	100%
LP → Rani 0.5PRN	20.3%	100%	15.8%	100%
Aflī 2q4	54.2%	100%	42.0%	100%
Rani 0.5PRN	9.2%	100%	9.0%	99.9%
LP → Bro 6q8 → q12	70.2%	79.7%	–	–
LP → Bro 6q12/q8	96.6%	–	81.8%	–
LP → Bro 3q12/q8	88.1%	87.6%	82.3%	48.1%
LP → Rani 0.5TREX	30.7%	100%	58.3%	68.5%
LP → Rani 0.5q8	11.5%	100%	–	–
LP → Aflī 2TREX	68.1%	90.4%	85.9%	37.6%
LP (4q4) → Fari 6q8-q16	73.5%	100%	–	–



**Table 3** continued

Treatment	SUCRA	Probability for LP → Bro 6q12/q8 to perform better than each comparator	SUCRA	Probability for LP → Bro 6q12/q8 to perform better than each comparator
LP (4q4) → Fari 6q12	55.2%	97.9%	–	–
LP (4q4) → Fari 6q16	30.9%	99.8%	–	–
Patients losing at least 15 letters	1 year		2 years	
Rani 0.5q4	56.6%	72.4%	60.1%	66.4%
LP → Rani 0.5PRN	33.3%	90.2%	26.5%	91.7%
LP → Afli 2q8	69.3%	54.7%	73.2%	50.7%
LP → Rani 0.5q12	54%	66.2%	–	–
LP → Rani 0.5q8	38.2%	82.7%	–	–
Afli 2q4	69.6%	52.4%	68.4%	55.7%
LP → Rani 0.5TREX	39.3%	87.4%	37.8%	82.3%
Rani 0.5PRN	70.2%	49.0%	53.6%	66.5%
LP → Bro 6q12/q8	70.2%	–	72.1%	–
LP → Bro 3q12/q8	70.1%	48.9%	59.5%	67.8%
Sham injection	3.2%	100%	0.1%	100%
LP → Rani 0.5PRNX	39.5%	77.9%	–	–
LP → Afli 2TREX	27.1%	89.9%	48.8%	67.4%
LP (4q4) → Fari 6q8-q16	63.5%	61%	–	–
LP (4q4) → Fari 6q12	61.2%	46.8%	–	–
LP (4q4) → Fari 6q16	34.8%	73.6%	–	–
Patients gaining at least 15 letters	1 year		2 years	
Rani 0.5q4	59.8%	74.2%	46.8%	88.3%
LP → Rani 0.5PRN	56.2%	72.6%	40.6%	85.4%
LP → Afli 2q8	50.5%	93.7%	62.0%	82.7%
LP → Rani 0.5q12	6.1%	99.7%	–	–
LP → Rani 0.5q8	39.1%	85.0%	–	–
Afli 2q4	66.7%	64.4%	42.9%	90.5%
LP → Rani 0.5TREX	68.9%	57.8%	74.2%	52.9%
Rani 0.5PRN	22.7%	98.7%	38.2%	85.3%
LP → Bro 6q12/q8	73.7%	–	78.7%	–
LP → Bro 3q12/q8	38.1%	97.9%	82.2%	39.9%
Sham injection	1.9%	100%	0.0%	100%

**Table 3** continued

Treatment	SUCRA	Probability for LP → Bro 6q12/q8 to perform better than each comparator	SUCRA	Probability for LP → Bro 6q12/q8 to perform better than each comparator
LP → Rani 0.5PRNX	85.0%	22.8%	–	–
LP → Aflī 2TRES	57.7%	63.1%	34.4%	83.0%
LP (4q4) → Fari 6q12	48.5%	65.8%	–	–
LP (4q4) → Fari 6q16	75.0%	36.4%	–	–
Discontinuation	1 year		2 years	
Rani 0.5q4	32.5%	49.1%	35.9%	62.3%
LP → Aflī 2q8	17.7%	76.7%	11.9%	93.2%
LP → Rani 0.5PRN	61.0%	24.6%	74.4%	24.4%
LP → Rani 0.5q8	91.4%	6.6%	–	–
Aflī 2q4	52.1%	28.6%	44.7%	54.3%
LP → Rani 0.5TRES	60.0%	23.4%	79.5%	20.5%
LP → Bro 6q8 → q12	49.2%	41.2%	–	–
LP → Bro 6q12/q8	34.5%	–	48.4%	–
LP → Bro 3q12/q8	60.8%	14.3%	61.4%	31.9%
LP → Aflī 2TRES	41.0%	45.7%	43.7%	52.6%
LP (4q4) → Fari 6q8-q16	25.6%	86.2%	–	–
LP (4q4) → Fari 6q12	8.0%	94.0%	–	–
LP (4q4) → Fari 6q16	12.6%	91.0%	–	–

dosing regimens. The results of the present study demonstrate that flexible dosing regimens of ranibizumab and aflibercept have similar efficacy on visual acuity as fixed dosing regimens while decreasing the disease burden for patients, caregivers, and physicians [10, 26, 27]. In addition, even with fewer injections, brolucizumab had significantly better morphological outcomes in terms of retinal thickness than its comparators (with comparable incidence of geographic atrophy and fibrosis across both treatment arms). However, further research is necessary to better understand the intricacies of the relationship between these anatomical outcomes and visual acuity.

The summary from our work on adverse events amongst anti-VEGF treatments in nAMD is in line with previous NMAs that evaluated ranibizumab and aflibercept [22, 28–32].

After the approval from the FDA, brolucizumab was associated with a safety signal of retinal vasculitis and/or retinal vascular occlusion (RV/RO), typically in the presence of intraocular inflammation [33]. These events were not reported with this wording, i.e., RV/RO, in the HAWK and HARRIER trials. The latest brolucizumab safety information can be found elsewhere, including in the brolucizumab.info website [11, 34–39].

**Table 4** Pooled mean number of injections received by each treatment regimen at 1 year, 2 years, and between 1 and 2 years

Intervention	Number of trials	Mean number of injections	Standard error	Cochran <i>Q</i> statistic	<i>p</i> value of Cochran <i>Q</i>	Between-trial variance
1 year						
Afli 2q4	1	11.90	0.13			
LP → Afli 2TREX	1	9.70	0.22			
LP → Afli 2q8	3	7.14	0.15	26.72	0.00	0.06
LP → Bro 3q12/q8	1	6.60	0.05			
LP → Bro 6q12/q8	2	6.66	0.05	1.64	0.20	0.00
LP (4q4) → Fari 6q12	1	6.70	0.43			
LP (4q4) → Fari 6q16	1	6.20	0.37			
LP (4q4) → Fari 6q8-q16 <sup>a</sup>	2	6.95	0.04	0.35	0.56	0.00
LP → Rani 0.5PRN	3	7.20	0.39	14.27	0.00	0.40
LP → Rani 0.5PRNX	1	5.50	0.31			
LP → Rani 0.5TREX	5	9.48	0.10	5.63	0.23	0.01
LP → Rani 0.5q8	1	7.60	0.05			
Rani 0.5PRN	1	6.90	0.18			
Rani 0.5q4	7	11.90	0.13	26.54	0	0.08
2 years						
Afli 2q4 → PRN	1	8.70	0.07			
LP → Afli 2TREX	1	8.50	0.27			
LP → Afli 2q8	2	6.35	0.25	19.23	0.00	0.12
LP → Afli 2q8 → PRN	1	6.10	0.06			
LP → Bro 3q12/q8	1	5.70	0.07			
LP → Bro 6q12/q8	2	5.70	0.20	14.16	0.00	0.07

**Table 4** continued

Intervention	Number of trials	Mean number of injections	Standard error	Cochran <i>Q</i> statistic	<i>p</i> value of Cochran <i>Q</i>	Between-trial variance
LP → Rani 0.5PRN	1	6.70	0.12			
LP → Rani 0.5TREX	3	8.88	0.11	2.16	0.34	0.00
Rani 0.5PRN	1	6.30	0.20			
Rani 0.5q4	4	11.53	0.35	50.86	0.00	0.43
Rani 0.5q4 → PRN	1	8.90	0.08			
1–2 years						
Afi 2q4 → PRN	1	4.80	0.08			
LP → Afi 2TREX	1	7.30	0.54			
LP → Afi 2q8	2	5.47	0.25	6.10	0.01	0.10
LP → Afi 2q8 → PRN	1	5.00	0.07			
LP → Bro 3q12/ q8	1	4.80	0.12			
LP → Bro 6q12/ q8	2	4.76	0.35	15.46	0.00	0.23
LP → Rani 0.5PRN	1	5.60	0.23			
LP → Rani 0.5TREX	3	8.21	0.21	0.38	0.83	0.00
Rani 0.5q4	3	11.29	0.70	26.65	0.00	1.23
Rani 0.5q4 → PRN	1	5.60	0.10			

<sup>a</sup>The average number of injections was calculated on the basis of the proportion of patients receiving faricimab doses every 8, 12, and 16 weeks after the initial loading phase of 4 monthly injections

RCTs, which are often considered the best available published evidence on comparative treatment efficacy and safety, have been shown to have low external validity and thus are limited when trying to decide which of multiple treatments is the most effective when used in clinical practice. This is in part due to the

inclusion of only a highly select group of patients into RCTs, the lack of data on direct clinical comparisons, and the lack of clinically pertinent endpoints. In addition, a conventional pairwise meta-analysis cannot fully inform the comparisons between each treatment due in part to different populations being

studied [40]. The application of an NMA helped overcome some of these limitations as all available evidence from the literature was pooled to indirectly compare treatments that had not been previously directly compared in a clinical trial. In particular, since only HAWK and HARRIER were head-to-head trials assessing the comparison between brolocizumab and aflibercept, conducting an NMA helped to estimate the relative efficacy of brolocizumab compared to other treatments not assessed in these two studies.

The process for conducting the present systematic literature review and NMA was based on reference standard guidelines, e.g., NICE [18, 41]. The results from this study were generally similar to the results of the NMA in nAMD conducted by NICE [21], and in the few differing cases the source of the discrepancy was found to be due to the decisions made on which studies to include. Patient characteristics among the included trials were similar at baseline, and little heterogeneity was identified in the direct comparisons, allowing for more robust results.

While the results from this NMA provide pertinent details comparing brolocizumab to its licensed comparators, several limitations were identified. First, it was not possible to conduct a meta-regression adjusted on relevant covariates such as baseline BCVA, age, and year of RCT start. The addition of these covariates yielded an over-parametrized model that could not converge. In addition, pooling indirect clinical evidence in an NMA does not necessarily reflect actual clinical practice, as the study population is selective in terms of inclusion and exclusion criteria, many included trials are historic, and patients must adhere to the study protocol. Further real-world studies evaluating these outcomes are therefore warranted. Moreover, patients with nAMD were evaluated in this study, and no specific subgroup analyses were conducted. Further research on the impact of anti-VEGFs on relevant subgroups, such as patients with PCV and types 1 (subpigment epithelial), 2 (subretinal), and 3 (retinal angiomatous proliferation) would provide a greater understanding of how these patient populations respond to the different treatments. Additionally, both patients naive to

other anti-VEGF and mixed populations were included in the analyses. Only two trials, PIER and MARINA, did not report prior use of other anti-VEGF as an inclusion/exclusion criterion. However, patients included in these trials were likely to be treatment naive as they were pivotal ranibizumab trials that took place when pegaptanib was the only licensed anti-VEGF treatment. In order to include all available evidence for the treatments of interest, time equivalence was assumed between 48 and 52 weeks for 1-year outcomes and between 96 and 104 weeks for 2-year outcomes. Finally, an NMA for the mean number of injections was not conducted since the number of injections was directly related to the type of regimen specified in the study protocol. As such, the pooled absolute number of injections was estimated using robust methods. Similarly, while an NMA is a superior method to study relative effects for serious adverse events, it was not feasible because of serious adverse events being rare events and a general lack of reported data on individual adverse events.

On the basis of the results of the present study, brolocizumab has comparable BCVA gains and discontinuation rates compared to other licensed anti-VEGF treatments and is superior in terms of reducing retinal thickness to most treatments. Fewer injections were administered for brolocizumab than its other comparators to achieve these results during 2 years of treatment, while injection frequency was following the respective study protocol specifications.

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**Compliance with Ethics Guidelines.** This article is based on previously conducted studies and does not contain any new studies with human participants or animals performed by any of the authors.

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