
Subtitle: Real world Outcomes in UK centres.

Gerald Liew PhD*, Aaron Y Lee MD*, Javier Zarranz-Ventura PhD, Irene Stratton FFPH, Catey Bunce PhD, Usha Chakravarthy PhD, Cecilia S Lee MD, Pearse A. Keane MD, Dawn Sim PhD, Toks Akerele MD, Martin McKibbin MD, Louise Downey MD, Salim Natha MD, Clare Bailey MD, Rehna Khan MD, Richard Antcliff MD, Stewart Armstrong MD, Atul Varma MD, Vineeth Kumar MD, Marie Tsaloumas MD, Kaveri Mandal MD, Catherine Egan PhD, Robert L Johnston** PhD, Adnan Tufail** PhD, on behalf of UK AMD EMR Users Group

* These authors contributed equally and GL and AYL should be considered joint first authors
** These authors contributed equally and RLJ and AT should be considered joint senior authors

Corresponding author – Adnan Tufail

Moorfields Eye Hospital NHS Trust, 162 City Road, London, EC1V 2PD

Adnan.Tufail@moorfields.nhs.uk

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Word count: 219
Complete UK AMD EMR Users Group (listed alphabetically by centre and within each centre)

28 Belfast Health and Social Care Trust, UK - Usha Chakravarthy

29 Calderdale and Huddersfield NHS Foundation Trust, UK - Rehna Khan

30 Countess of Chester Hospital NHS Trust, UK - Stewart Armstrong

31 Gloucestershire Hospitals NHS Foundation Trust, UK - Robert Johnston, Quresh Mohamed, Ahmed Sallam, Javier Zarranz-Ventura, Paul Donachie, Irene Stratton

32 Hinchingbrooke health care NHS Trust, UK - Toks Akerele

33 Hull and East Yorkshire Hospitals NHS Trust, UK - Seema Arora, Helen Cook, Louise Downey, Kala Gopalakrishnan, Fiona Lyon, Tahir Islam, Naeem Zaman.

34 Leeds Teaching Hospitals NHS Trust, UK - Oliver Backhouse, Tim Dabbs, Bryn Davies, Martin McKibbin, Bataung Mokete, Damian O'Neill

35 Mid Yorkshire Hospitals NHS Trust, UK - Atul Varma

36 Moorfields Eye Hospital NHS Trust, UK - Catey Bunce, Catherine Egan, Pearse A Keane, Gerald Liew, Praveen J Patel, Dawn A Sim, Adnan Tufail, Wen Xing

37 Royal United Hospital Bath NHS Trust, UK - Richard Antcliff

38 University Hospital Birmingham NHS Foundation Trust, UK - Marie Tsaloumas

39 University Hospitals Bristol NHS Foundation Trust, UK - Clare Bailey

40 University of Sydney, Sydney NSW, Australia – Gerald Liew

41 University of Washington, Washington, USA – Aaron Lee, Cecilia S Lee

42 University of Washington, Washington, USA – Kaveri Mandal

43 Wirral University Teaching Hospital NHS Foundation Trust, UK - Vineeth Kumar

44 Wrightington, Wigan and Leigh NHS Foundation Trust, UK - Salim Natha

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Abstract

**Aims:** International variations in visual acuity (VA) outcomes of eyes treated for neovascular age-related macular degeneration (nAMD) are well documented, but intra-country inter-centre regional variations are not known. This data is important for national quality outcome indicators. We aimed to determine intra-country, inter-centre regional variations in outcomes for treatment of nAMD.

**Methods:** Prospective multi-centre national database study of 13 UK centres that treated patients according to a set protocol (3 loading doses, followed by pro-re-nata re-treatment). 5,811 treatment naive eyes of 5,205 patients received a total of 36,206 ranibizumab injections over 12 months.

**Results:** Mean starting VA between centres varied from 48.9 to 59.9 ETDRS letters. Mean inter-centre VA change from baseline to 12 months varied from +6.9 letters to -0.6 letters (mean of +2.5 letters). The proportion of eyes achieving VA of 70 letters or more varied between 21.9% and 48.7% at 12 months. Median number of injections (visits) at each centre varied from 5 to 8 (9 to 12) with an overall median of 6 (11). Age, starting VA, number of injections and visits but not gender were significantly associated with variation in these VA outcomes (P<0.01). Significant variation between centres persisted even after adjusting for these factors.

**Conclusions:** There are modest differences in VA outcomes between centres in the UK. These differences are influenced, but not completely explained, by factors such as patient age, starting VA, number of injections and visits. These data provide an indication of the VA outcomes that are achievable in real world settings.
Introduction

Age-related macular degeneration (AMD) is one of the leading causes of irreversible blindness in patients aged over 60 years. Since the introduction of anti-Vascular Endothelial Growth Factor (VEGF) agents to treat neovascular AMD (nAMD) in 2006, rates of blindness and visual impairment from AMD have declined dramatically.\(^1\)\(^-\)\(^3\) Data from several randomized clinical trials suggest patients gain on average 6-11 letters using the most aggressive monthly dosing posology in the first year of treatment.\(^4\)\(^-\)\(^7\) A less resource intensive Pro-Re-Nata (PRN) dosing posology has been found in the Comparison of Age-related macular degeneration Treatment Trials (CATT)\(^6\)\(^,\)\(^8\) and Inhibit VEGF in Age-related choroidal Neovascularisation (IVAN)\(^7\)\(^,\)\(^9\) trials to produce outcomes similar to monthly dosing. This has provided a sound empirical basis for use of PRN posology in the National Health System (NHS) in the United Kingdom (UK).

Although PRN dosing posology in theory is able to achieve excellent outcomes, ‘real-world’ studies have not matched the outcomes achieved in clinical trials.\(^10\)\(^-\)\(^13\) We have previously reported longitudinal results from a cohort of patients recruited from hospitals in the UK.\(^10\)\(^,\)\(^11\)\(^,\)\(^14\) Patients had standardised data recorded at the point of care into an Electronic Medical Record (EMR) system and the grouped national findings showed that PRN retreatment after 3 loading doses resulted in moderately stable vision.\(^11\) National data have also been reported from Australia for a treat and extend posology.\(^15\) International multi-country real life comparisons have reported some international differences in VA outcomes and treatment patterns,\(^16\) which may be influenced by different reimbursement and health system structures. However, what remains unclear is how much, if any, intra-country inter-centre regional variation occurs. This data is obscured by aggregate national data, and is important as it permits understanding of the factors that influence outcomes in a real world setting and allows national quality benchmarks to be set. In this report we examine inter-centre variations in patient characteristics, the number of treatments delivered and their impact on visual outcomes in 13 geographically distinct UK centres.

Methods
Study Design

The study design is described in detail in previous reports from the UK-nAMD Database Study group.\textsuperscript{10,11,14} In brief, sites known to make comprehensive use of Electronic Medical Records (EMR) systems were contacted and requested to contribute data. Patient identifiers were completely removed and site and clinician data were pseudo-anonymised. On this basis an ethics committee determined that formal ethics approval was not required. The study was conducted in accordance with the declaration of Helsinki and the UK’s Data Protection Act.

Study Centres

Thirteen NHS hospitals that deliver ranibizumab AMD treatment services in England and Northern Ireland submitted data to this study. Each site is the sole NHS provider of nAMD care to their local population and very few patients switch between providers. Following NICE approval for the use of ranibizumab for nAMD in the NHS in August 2008 all sites used this drug almost exclusively, although prior to this date some sites offered limited treatment with bevacizumab. The study was initiated on 1st Feb 2012, all approvals and data extraction was performed by 02 April 2012. Data was delivered to the analysis team by the end of April 2012.

Data variables

Analysis was restricted to treatment-naïve eyes undergoing ranibizumab therapy conforming to 3 monthly loading dose followed by PRN posology for nAMD that completed at least 12 months of follow up. In this report the ‘best-measured VA’ was the best VA with refraction or habitual correction and/or pinhole as measured on an ETDRS chart and expressed as LogMAR vision and ETDRS letters. The vast majority of sites measured VA with habitual correction rather than best-corrected refracted VA at all time points and used ETDRS charts. Analysis for eyes with very low VA was undertaken by substituting counting fingers (CF), hand movement (HM), and perception of light (PL) with 2.0, 2.3, and 2.7, respectively. We examined the outcome measures commonly reported in clinical trials (VA change from 0 to 12 months, proportion of eyes gaining and losing 15 ETDRS letters etc) as well as other measures such as VA change from 3 to 12 months and proportion of eyes achieving 70 ETDRS letters at 12 months (Snellen equivalent of 20/40 or 6/12, driving equivalent in many jurisdictions).
Statistical methods

Medisoft Ophthalmology (Medisoft Limited, Leeds, UK) was the EMR system used for data extraction. Data for right and left eyes of patients who had had at least one intravitreal injection of ranibizumab for nAMD were extracted. Both STATA version 11 and SPSS version 19 were used to analyse data. Perl and R package ggplot2 was used for multivariable analyses, construction of generalised linear models and creation of funnel plots.

Results

Participants

Over the 1 year of follow up analysed in this study, 36,206 ranibizumab injections were performed in 5,811 eyes. Table 1 shows the baseline demographics of each of the 13 centres. The number of eyes treated ranged from 39 (Centre M) to 923 (Centre C). Patients treated at each centre were of similar age, ranging from a mean of 78.3 (J) to 81.7 years at the time of the first injection (K).

Treatment and outcome characteristics of each centre are shown in Table 2. The median number of injections in each centre was 6 with one centre providing a median of 8 injections (J) and 2 centres providing a median of 5 (L, M). Mean starting VA across all centres was lowest in centre F (48.9 ETDRS letters) and highest in centre L (59.9 letters). VA after 12 months of treatment was highest in centre I (63.2 letters) and lowest in centre M (52.9 letters). The centres with the higher mean starting VA did not necessarily finish 12 months with the highest VA at 1 year, nor did the centres with the lowest starting VA finish with the lowest VA at 1 year. Centres that saw patients the most frequently (B) or injected most frequently (J) achieved the 5th and 2nd best VA at 1 year respectively and showed the least variation between 3 and 12 months.

At 12 months, the mean change in VA across 13 centres varied from +6.9 letters (centre I) to -0.6 letters (centre L), a difference of 7.5 letters, with a mean VA gain of +2.5 letters. Figure 1 shows these results graphically, with further details in Table 2. The funnel plot in Figure 1 shows the distribution of centres and 95% and 99% confidence intervals. Two centres (L, D) were slightly outside the 99% confidence intervals; these were the only centres reporting a slight reduction in VA. One of these centres had the lowest number of injections (5, L); however the other centre had the median number of injections (6, D). Starting VA in
centre D was near the middle of the distribution, at 53.7 letters, while in centre L, eyes commenced
treatment with the best starting VA at 59.9 letters. In contrast, Figure 2 (supplement) shows that there was
less variation in VA change from 0 to 3 months, with a tighter clustering of results. Figure 3 (supplement)
shows the VA change from 3 months to 12 months, with most centres clustered within the 95% confidence
intervals, and the previous 2 centres that were outside the 99% CI when considering change from 0-12
months are now either within or very close to the limits (L, D).

The proportion of eyes that gained 15 ETDRS letters or more at 12 months ranged from 7.7% (A) to 29.5%
(K). When examined on the funnel plot (Figure 4 supplement), these proportions showed little variation
from centre to centre, with all centres within or above the 99% CI. The proportion of eyes that lost 15
ETDRS letters or more at 12 months showed an even tighter distribution, with all centres within the 99% CI,
and all except one (L) within the 95% CI (Figure 5 supplement). The actual proportions ranged from 4.6% (I)
to 11.7% (L). We also examined the proportion of eyes maintaining or achieving driving vision of 70 ETDRS
letters or more and found all centres performed above the lower 99% CI limit (Figure 6).

It should be noted that the centre with the lowest number of visits and injections (M, median 9 visits,
median 5 injections, Table 2) had outcomes in the middle of the distributions for all the VA measures
studied (Figures 1-6). Similarly, the centres with the highest number of visits (B, 12 visits) and highest
number of injections (J, 8 injections), generally had outcomes either in the middle of the distribution or
higher than average but still within the 95% CI (Figures 1-6).

We performed multivariable analyses to determine which factors were associated with better visual
outcomes. Younger age, worse starting VA, and higher number of injections and visits but not gender were
significantly associated with variation in these VA outcomes (P<0.01). Significant variation between centres
persisted even after adjusting for these factors.

**Discussion**

There is considerable published data on VA outcomes derived from clinical trials but limited data describing
real-world outcomes. Real-world outcomes indicating what is possible when trial results translate to clinical
practice are ultimately the most important measure as they reflect what happens to whole populations of
patients rather than the rarefied cohorts included in trials. They are also important for establishing
benchmarks standards that are achievable in busy public systems, and for defining measures of quality care
that take into account the heterogeneity of patient populations and care delivery systems. This study
provides some of the first real-world outcomes from a single national health system, namely 13 UK public
hospital centres using a PRN treatment posology. We report that there was some inter-centre variation in
VA outcomes up to a maximum difference of 7.5 letters between the highest and lowest VA achieved from
0-12 months. Age, starting VA, number of injections and visits but not gender were significantly associated
with variation in these VA outcomes.

The median performance of these 13 centres is comparable to results from clinical trials, once the lower
starting VA and lower number of injections is taken into account. Table 3 compares findings from this study
with clinical trial results. The CATT6 and IVAN7-9 studies achieved mean improvement from 0-12 months of
6.8 and 5.0 letters, respectively with 7 injections, while the Groupe d’Etude Français Avastin versus Lucentis
dans la DMLA néovasculaire (GEFAL)17 and Multicentre Anti-VEGF Trial in Austria (MANTA)18 studies
showed mean improvement of 2.9 and 4.1 letters respectively with 6 injections, as compared with mean
improvement of 2.5 letters with 6 injections in this study. These results suggest that a similar benchmark of
+2.5 letters improvement (0-12 months) with 6 injections represents quality ongoing care that is achievable
in a real-world, public hospital setting.

There are few other real-world studies with which to compare our results. A large database observational
study from Australia, the Fight Retinal Blindness Study,19 reported mean VA gains of 5.3 letters after 2 years
and 13 injections of a treat and extend posology. These results are superior to those achieved in this report
but direct comparisons are difficult due to differences in health systems, patient mix, and different follow-
up periods.

It should be noted that the difference in 0-12 month VA change between the highest and lowest scoring
centres was 7.5 ETDRS letters, a difference that is only marginally beyond what some studies have
considered non-inferior. The CATT6 and GEFAL17 considered a difference of 5 EDTRS letters to represent
noninferiority, while the MANTA18 considered a difference of 7 letters to be noninferior.18
Due to the ‘ceiling effect’ whereby eyes starting treatment with good VA have little room for further improvement, many measures are dependent on the starting VA.\textsuperscript{11;12;20} Adjusting for age, starting VA, number of injections and visits reduced, but did not eliminate the significant variation between centres, suggesting there are other unmeasured factors that contribute to these variations in outcomes.

This study has several strengths including a large sequential sample, collection of a standardised minimum dataset as mandated by the use of an EMR reflecting routine, real-world clinical practice and the large number of centres involved. A weakness of this study is the loss to follow up of a number of patients over time, as is inevitable in a real-world clinical setting. Although there were differences in baseline demographics between patients lost to follow-up and those who completed follow-up, we have previously shown that VA changes are similar in both groups.\textsuperscript{11} Best-corrected VAs were not routinely measured, instead the VAs with habitual correction were reported in this study which may underestimate absolute VA measurements compared with clinical trials. It should be noted that clinical treatment decisions were based on these VAs and we believe these represent real-world outcomes and may better reflect patients’ visual experience than protocol determined best-corrected VAs. A possible reason for differences in injection numbers may be individual differences in centre / physician thresholds for retreatment using a PRN posology.

In summary, we report that 13 UK centres using a PRN treatment posology for managing neovascular AMD achieved broadly similar VA outcomes with modest variability in outcomes. The difference between the highest and lowest VA gain at 12 months was 7.5 letters with a mean of +2.5 letters gained. Age, starting VA, number of injections and visits but not gender were significantly associated with variation in these VA outcomes. These data may be considered as establishing an achievable benchmark for the quality of PRN posology in real-world settings, and is likely to be relevant to many sites worldwide.
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Financial Disclosures

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GL and AF had full access to all the data in the study and take responsibility for the integrity of the data and the accuracy of the data analysis.
Reference List


Figure Legends.

**Figure 1**: Funnel plot showing the change in ETDRS letters from baseline to 12 months by centre. Solid black lines represent the 95% confidence intervals, dashed lines the 99% confidence intervals. ETDRS refers to Early Treatment Diabetic Retinopathy Study.

**Figure 2 (supplement)**: Funnel plot showing the change in ETDRS letters from baseline to 3 months by centre. Solid black lines represent the 95% confidence intervals, dashed lines the 99% confidence intervals. ETDRS refers to Early Treatment Diabetic Retinopathy Study.

**Figure 3 (supplement)**: Funnel plot showing the change in ETDRS letters from 3 months to 12 months by centre. Solid black lines represent the 95% confidence intervals, dashed lines the 99% confidence intervals. ETDRS refers to Early Treatment Diabetic Retinopathy Study.

**Figure 4 (supplement)**: Funnel plot showing the proportion of eyes gaining 15 ETDRS letters or more from baseline to 12 months by centre. Solid black lines represent the 95% confidence intervals, dashed lines the 99% confidence intervals. ETDRS refers to Early Treatment Diabetic Retinopathy Study.

**Figure 5 (supplement)**: Funnel plot showing the proportion of eyes losing 15 ETDRS letters or more from baseline to 12 months by centre. Solid black lines represent the 95% confidence intervals, dashed lines the 99% confidence intervals. ETDRS refers to Early Treatment Diabetic Retinopathy Study.

**Figure 6**: Funnel plot showing the proportion of eyes maintaining driving vision of 70 ETDRS letters or better at 12 months by centre. Solid black lines represent the 95% confidence intervals, dashed lines the 99% confidence intervals. ETDRS refers to Early Treatment Diabetic Retinopathy Study.