Prospective audit of exudative age-related macular degeneration: 12-month outcomes in treatment-naive eyes

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Abstract: Purpose: To report the 12-month outcomes of 1,140 treatment-naïve eyes with exudative age-related macular degeneration (wet AMD) who were treated for 12 months with intravitreal anti-VEGF drugs in routine clinical practice. Methods: Index visit characteristics, such as lesion type and size, visual acuity (VA in logMAR [Logarithm of the Minimal Angle of Resolution] letters), as well as treatments, outcomes (VA, lesion activity status) and ocular adverse events were recorded in a prospectively designed electronic database. Index visit characteristics associated with the 12-month VA outcome were identified using mixed effects linear regression. Results: Mean change in VA in the cohort after 12 months was +4.7 logMAR letters (95%CI: 3.4 to 6.1) with a mean of 7.0 injections. No significant difference was found in change in VA or number of injections by type or size of the lesion. Median time to inactivation of lesions was 194 days. VA at the index visit was the strongest predictor for the 12-month outcomes. Infectious endophthalmitis occurred in 2 cases, retinal detachment in 1 case from a total of 9,162 injections. Conclusions: These findings indicate that VEGF inhibitors can achieve reasonably good outcomes for wet AMD when used in routine clinical practice.

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Prospective Audit of Exudative Age-Related Macular Degeneration: 12-month Outcomes in Treatment-Naïve eyes.

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Running head: 12-month outcomes of the FRB! Project wet AMD audit

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ABSTRACT

Purpose
To report the 12-month outcomes of 1,140 treatment-naïve eyes with exudative age-related macular degeneration (wet AMD) who were treated for 12 months with intravitreal anti-VEGF drugs in routine clinical practice.

Methods
Index visit characteristics, such as lesion type and size, visual acuity (VA in logMAR [Logarithm of the Minimal Angle of Resolution] letters), as well as treatments, outcomes (VA, lesion activity status) and ocular adverse events were recorded in a prospectively designed electronic database. Index visit characteristics associated with the 12-month VA outcome were identified using mixed effects linear regression.

Results
Mean change in VA in the cohort after 12 months was +4.7 logMAR letters (95%CI: 3.4 to 6.1) with a mean of 7.0 injections. No significant difference was found in change in VA or number of injections by type or size of the lesion. Median time to inactivation of lesions was 194 days. VA at the index visit was the strongest predictor for the 12-month outcomes. Infectious endophthalmitis occurred in 2 cases, retinal detachment in 1 case from a total of 9,162 injections.

Conclusion
These findings indicate that VEGF inhibitors can achieve reasonably good outcomes for wet AMD when used in routine clinical practice.
While the efficacy of ranibizumab (Lucentis®, Novartis, Switzerland), bevacizumab (Avastin®, Roche, Switzerland) and aflibercept (Eylea®, Bayer, Switzerland) for exudative age-related macular degeneration (wet AMD)\(^1\) has been convincingly demonstrated by tightly controlled phase 3 clinical trials,\(^2-7\) it is still not certain that the results of these studies will be replicated in the real world after the new drugs have been approved for general use. Many patients being treated for wet AMD in the general community may not have met inclusion criteria of the clinical trials. Even if they had, a heavy treatment burden on all involved in routine retinal practice has led to dosing regimens that are less intensive than those used in the pivotal trials, such as the pro re nata (PRN) and treat-and-extend regimens.\(^8,9\)

The Fight Retinal Blindness! (FRB) Project has established a prospective audit system that can anonymously track outcomes of treatment of retinal disease, such as wet AMD, in large numbers of patients treated in routine retinal treatment centres.\(^10\) Here we describe the 12-month outcomes, including visual acuity, grading of lesion activity and adverse events, for 1,140 treatment-naïve participants in the FRB! Project wet AMD audit.
METHODS

Study design & Setting

This is an observational study utilising anonymised longitudinal data from the FRB registry that were captured during routine clinical practice. All treatment decisions and visit schedules were entirely at the discretion of the treating physician and patient. Details of the FRB project data tracking system have been published. The research followed the tenets of the declaration of Helsinki. Patients were given information regarding the project and given the opportunity to opt out of the project. Each of the three academic core centres from the Universities of Sydney, Melbourne and Western Australia obtained approval from their respective Human Research Ethics Committees (HREC) to conduct the project as a quality assurance activity. Overarching ethical approval for the other centres was obtained from the Royal Australian and New Zealand College of Ophthalmologists’ HREC.

Patient data recorded from 27 retinal specialists located across Australia from January 2006 until September 2012 were aggregated for analysis. The project began collecting data from the core centres in Sydney, Melbourne and Perth then spread to non-academic retinal services in the capital cities of most Australian states.

Participants and Variables

Few eligibility criteria were applied beyond treatment-naïve eyes commencing treatment for wet AMD that had been diagnosed by their treating ophthalmologist with VA > 20 letters. All eyes in the database that commenced treatment between Jan 2004 and Nov 2011 were included in this analysis, so that all potentially had 12 months follow-up. At the index visit, i.e. the visit at which treatment was commenced, the study participants’ age, angiographic lesion criteria such as lesion type and greatest linear dimension (GLD), visual acuity (Logarithm of the Minimum Angle of Resolution – LogMAR, recorded as letters read), choroidal neovascularization (CNV) status (active, inactive), along with treatment history and
treatment decisions (treated or not treated and name of drug used) were recorded. Investigators were asked to enter whichever VA reading was best: uncorrected, corrected or pinhole. The best VA achieved during each visit was used for analysis. The judgement of “active” or “inactive” was left to the investigator’s discretion, thus reflecting real-world practice. It was suggested that Users should grade lesions as active if there was intra- or sub-retinal fluid, or any other feature, present that could be attributed to activity of the neovascular lesion. Follow-up visits recorded subsequent VA, CNV status, all treatment decisions and any ocular adverse events. Three subgroups of interest were pre-specified: occult lesions (OC), minimally classic lesions (MC) and predominantly classic lesions (PC).

Statistical methods

For continuous variables means or medians and interquartile range (Q1, Q3) were computed. Seventeen percent of patients contributed both eyes to the study database; when measuring variation and performing statistical tests at the index visit, fellow eyes were randomly removed to ensure any possible inter-eye correlation would not bias estimates. Formal comparisons were made using the non-parametric Kolmogorov-Smirnoff (KS) test, which is sensitive to any difference in the underlying distribution of two samples.

The outcomes analysis used data from all eyes that completed 12 months follow-up, while the safety analysis set included all available data over 12 months. We also examined outcomes for eyes that did not complete 12 months follow-up due to withdrawal from treatment or loss to follow-up. Study endpoints included 12-month longitudinal VA, time from first intravitreal injection to inactivation of CNV and change in CNV status over 12 months. Within-eye changes in VA over 12 months were tested using the paired t-test. Longitudinal VA data were plotted using a Lowess smoothed regression line. A mixed effects regression model was
fitted to the longitudinal VA data to examine the effects of lesion type, GLD and age on VA at 12 months.

Kaplan-Meier analysis\textsuperscript{13} was used to examine time from first injection to inactivation of CNV status. All observed adverse events were tabulated and reported. Analysis and plots were done using R version 2.15.0\textsuperscript{14}.
RESULTS

There were 1,140 eyes that completed 12 months follow-up (10,758 visits). The study population was 61% female and the mean age was 79.3 years (Q1,Q3: 75, 85). Mean visual acuity at the index visit was 57.1 letters (Q1,Q3: 45, 69) (Table 1). Owing to the quality assurance features of the FRB web-based data entry system, data quality was high for all variables (> 99.5% complete) with the exception of GLD (80% complete) and lesion type (88% complete).

Table 1: Index visit characteristics of eyes that completed 12 months follow-up and those that did not

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>12-month Completers</th>
<th>Non-completers</th>
</tr>
</thead>
<tbody>
<tr>
<td>Eyes</td>
<td>1,140</td>
<td>230</td>
</tr>
<tr>
<td>Visits</td>
<td>10,758</td>
<td>1,496</td>
</tr>
<tr>
<td>Median days follow-up (Q1, Q3)</td>
<td>-</td>
<td>210 (111, 302)</td>
</tr>
<tr>
<td>Mean index VA (Q1, Q3)</td>
<td>57.1 (45, 69)</td>
<td>52.5 (40, 65)</td>
</tr>
<tr>
<td>Mean age (Q1, Q3)</td>
<td>79.3 (75, 85)</td>
<td>79.9 (75, 85)</td>
</tr>
<tr>
<td>Female</td>
<td>61.3%</td>
<td>58.3%</td>
</tr>
<tr>
<td>Median GLD (Q1, Q3)</td>
<td>2,000 (1,300, 3,050)</td>
<td>2,315 (1,500, 3,390)</td>
</tr>
</tbody>
</table>

Lesion type

- Occult n (%) 529 (53.6 %) 109 (51.4 %)
- Minimally classic n (%) 211 (21.4 %) 50 (23.6 %)
- Predominantly classic n (%) 171 (17.3 %) 43 (20.3 %)
- Other n (%) 76 (7.7 %) 10 (4.7 %)
- Unclassified n 163 18

GLD: Greatest Linear Dimension; VA: LogMAR Visual Acuity
Treatment administered

A total of 8,013 injections were given to the 1,140 eyes that completed 12 months, a mean \((Q_1,Q_3)\) of 7.0 (5, 9) injections per eye (Table 2). The mean number of injections by lesion type was similar. The majority of injections administered were ranibizumab (91%) irrespective of lesion type, with the remainder being bevacizumab. For all lesion types the interval between injections was greater when lesions were graded as inactive than when they were graded as active.

Table 2: Injection frequency and type over 12 months follow-up

<table>
<thead>
<tr>
<th></th>
<th>Occult</th>
<th>Min class</th>
<th>Pred class</th>
<th>All</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mean ((Q_1,Q_3)) number of injections</td>
<td>7.0 (5, 9)</td>
<td>6.8 (5, 9)</td>
<td>7.1 (5, 9)</td>
<td>7.0 (5, 9)</td>
</tr>
<tr>
<td>Median ((Q_1,Q_3)) days between injections when active</td>
<td>35 (28, 52)</td>
<td>41 (29, 55)</td>
<td>35 (28, 56)</td>
<td>36 (28, 56)</td>
</tr>
<tr>
<td>Median ((Q_1,Q_3)) days between injections when inactive</td>
<td>43 (35, 63)</td>
<td>49 (36, 63)</td>
<td>42 (33, 56)</td>
<td>42 (35, 63)</td>
</tr>
<tr>
<td>% Ranibizumab injections</td>
<td>91.3%</td>
<td>92.3%</td>
<td>91.4%</td>
<td>91.4%</td>
</tr>
</tbody>
</table>

Nin Class = minimally classic; predom = predominantly

VA and GLD at the index visit

There were notable differences in the distributions of VA and GLD among the lesion type subgroups (Table 3, Figure 1). Visual acuity when starting treatment was lower for the PC classic group than the OC subgroup \((P<0.0001;\ KS\text{-}test)\) and the MC subgroup \((P=0.01;\ KS\text{-}test)\).
GLD was lower in the PC group than either OC or MC: OC vs. PC (P=0.002); MC vs. PC (P=0.005); OC vs. MC (P=0.5), (Figure 1).

Table 3: Index visit visual acuity and greatest linear dimension with visual acuity change after 12 months

<table>
<thead>
<tr>
<th></th>
<th>Occult Mean (Q1, Q3) VA Index visit</th>
<th>Min class Mean (Q1, Q3) VA Index visit</th>
<th>Pred class Mean (Q1, Q3) VA Index visit</th>
<th>All Mean (Q1, Q3) VA Index visit</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mean 12 month VA change (95% CI*)</td>
<td>4.9 (2.1 to 7.1)</td>
<td>4.5 (1.9 to 7.1)</td>
<td>5.1 (1.9 to 8.2)</td>
<td>4.7 (3.4 to 6.1)</td>
</tr>
<tr>
<td>Median (Q1, Q3) Index GLD</td>
<td>2,080 (1,255, 3,200)</td>
<td>2,015 (1,525, 3,030)</td>
<td>1,740 (1,065, 2,555)</td>
<td>2,000 (1,300, 3,050)</td>
</tr>
</tbody>
</table>

* Bias corrected and accelerated bootstrap 95 % confidence intervals

Unadjusted 12 month VA outcomes

The mean within-eye change in visual acuity was a +4.7 letter improvement (95%CI: 3.4 to 6.1) for the study population as a whole. Similar clinically relevant mean improvements were observed for all subgroups (Figure 2A): OC +4.9 letters (95%CI: 2.1 to 7.1), MC +4.5 letters (95%CI: 1.9 to 7.1) and PC +5.1 letters (95%CI: 1.9 to 8.2). The Lowess lines indicate that all three groups exhibited a monotonic improvement throughout 12 months (Figure 2B).

Modelled 12 month VA outcomes

Given the observed imbalance at the index visit in VA and GLD for the 3 subgroups, a mixed effects regression model was fitted to the longitudinal VA measurements to mitigate potential confounding influences (Table 4). The model coefficients for the MC and PC lesions (relative to OC) of -1.3 and -0.5 respectively indicate that lesion subgroup had very little effect (less than 1.5 LogMAR letters) on visual acuity outcomes. The coefficient for Age of -0.03 indicated slightly worse outcomes with increasing age: a three decade increase in age was associated with a decreased gain of 1 LogMAR letter after 12 months of treatment. A 1mm
(1,000 µm) increase in GLD was associated with a reduced gain of 0.5 letters. The coefficient for Time indicated an annual mean improvement of 3.1 letters. Visual acuity at the index visit was a highly significant predictor of outcome.

Table 4: Coefficients from mixed effects model fit to 12-month longitudinal VA data.

<table>
<thead>
<tr>
<th>Model coefficient</th>
<th>t value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Index Visual Acuity</td>
<td>0.9</td>
</tr>
<tr>
<td>Index Visit Age</td>
<td>-0.03</td>
</tr>
<tr>
<td>MC (relative to OC)</td>
<td>-1.3</td>
</tr>
<tr>
<td>PC (relative to OC)</td>
<td>-0.5</td>
</tr>
<tr>
<td>GLD 1000µm</td>
<td>-0.5</td>
</tr>
<tr>
<td>1 Year Follow-up</td>
<td>3.1</td>
</tr>
</tbody>
</table>

Lesion activity over 12 months

The median time from first intravitreal injection to lesions being graded as “inactive” was 194 days (95%CI: 174 to 216, Figure 3). Thirty seven percent of eyes were persistently graded as active during the 12 months of treatment. The median time between injections was 36 days (Q1,Q3: 28, 56) while the lesions were graded as “active” and 42 days (Q1,Q3: 35, 63) while graded “inactive”.

Eyes that did not complete 12 months follow-up

Two hundred and thirty eyes (17%) either withdrew from treatment or were lost to follow-up over the observed 12-month interval (non-completers). Median follow-up time for these eyes was 210 days (Q1,Q3: 111, 302). At the index visit non-completers were similar to completers in most respects except for lower VA (mean 57.1 vs. 52.5; P=0.0004, KS-test). The outcomes for non-completers are shown in longitudinal profiles in Fig 4.
Safety

Ocular adverse events observed over 12 months follow-up are summarised in Table 5. The most common adverse event was patient-reported post-injection pain (45 instances). Two instances of infectious endophthalmitis were reported out of a total of 9,162 injections.

Table 5: Adverse Events

<table>
<thead>
<tr>
<th>Event</th>
<th>Frequency</th>
<th>Injections per AE</th>
</tr>
</thead>
<tbody>
<tr>
<td>Post injection pain reported</td>
<td>45</td>
<td>204</td>
</tr>
<tr>
<td>Haemorrhage reducing VA &gt; 15</td>
<td>5</td>
<td>-</td>
</tr>
<tr>
<td>Retinal detachment</td>
<td>1</td>
<td>9162</td>
</tr>
<tr>
<td>Non-infectious endophthalmitis</td>
<td>2</td>
<td>4581</td>
</tr>
<tr>
<td>Infectious endophthalmitis</td>
<td>2</td>
<td>4581</td>
</tr>
<tr>
<td>RPE tear</td>
<td>12</td>
<td>764</td>
</tr>
<tr>
<td>Cataract extraction / other surgery</td>
<td>15</td>
<td>611</td>
</tr>
</tbody>
</table>

VA = Visual Acuity; RPE = Retinal Pigment Epithelium
DISCUSSION

This analysis of outcome data that were collected prospectively and continuously from patients receiving treatment for exudative AMD has produced a number of observations on the use and outcomes of intravitreal therapy in routine practice. Mean visual acuity of the main cohort improved significantly by +4.7 logMAR letters over the first 12 months of treatment with a mean of 7 injections. The mean visual acuity of predominantly classic lesions improved slightly more than that of the minimally classic or occult groups, although eyes with predominantly classic lesions had lower visual acuity at the index visit. Otherwise, lesion type and size made little difference to the pattern of treatment outcomes, of which the strongest predictor was visual acuity at the first treatment visit. The median time to first grading of lesions as inactive was 194 days, with 37% still active at 12 months. Safety findings were similar to previous reports. These findings indicate that VEGF inhibitors achieve good outcomes for wet AMD when used in routine clinical practice.

Several other observational studies of intravitreal therapy for neovascular AMD have recently been published. The Swedish Lucentis Quality Registry found a good improvement in visual acuity after 3 injections of ranibizumab, but this subsequently dropped back to pre-treatment levels. Patients in that study received a mean of only 4.8 injections over 12 months, fewer than in the present study. Similar results were found by the WAVE study and an analysis of the German reinjection scheme. These studies that recorded lower gains in mean visual acuities also had a lower mean number of injections.

An improvement in mean visual acuity after the first 12 months of treatment that was more similar to our results has been reported by two other observational studies. A gain of 3.2 LogMAR letters was found with a mean of 5.1 injections in the French Lumiere study of 551 patients. Menghini et al. reported a mean improvement of 5 letters with a mean of 4
injections in 204 eyes.\textsuperscript{20}

An overall mean improvement of 4.7 logMAR letters in the current report is still somewhat less than was reported in phase 3 clinical trials of ranibizumab.\textsuperscript{2,3} However the improvements in these studies were primarily measured against the change of vision in the control groups. Verteporfin-treated eyes had lost a mean of 9.5 letters by 12 months in ANCHOR, while sham-treated eyes had lost 10.4 letters in MARINA. Seen in this light, the increase in visual acuity found in the present analysis of outcomes of treated eyes in routine practice is reassuring. This was achieved with a mean of 7.0 injections, significantly more than was given in previously reported observational studies,\textsuperscript{16-20} out of potentially 13 that would be given with a strict monthly regimen. This frequency is similar to that of the CATT study, in which a mean of 6.9 injections were given to the ranibizumab PRN group and 7.7 to the bevacizumab PRN group.\textsuperscript{5}

Median time to grading the lesion as “inactive” was 194 days. Thirty seven percent of lesions were consistently graded as active throughout the first year of the study. As might be expected, these eyes received more injections. A related variable, presence of fluid at the 1-year visit, was reported in 81% of bevacizumab PRN and 56% of ranibizumab monthly groups of the CATT study.\textsuperscript{5} It appears that reasonably good visual acuity outcomes can be obtained despite many eyes remaining active much or all of the time.

Lesion characteristics, particularly lesion size (GLD) and type, did not significantly affect the outcomes of this study. Lesion type also had little effect on outcomes in retrospective analyses of MARINA and ANCHOR, in which mixed lesions had similar outcomes to purely classic or purely occult lesions.\textsuperscript{21,22} Menghini et al. also found no effect of lesion type on visual outcome after 24 months treatment in another observational study.\textsuperscript{20} In a recent report
from Comparison of AMD Treatment Trials, predominantly or minimally classic vs. occult CNV was not included in the final multivariate model of change in VA at 1 year because it was not statistically significant. Predominantly or minimally classic lesions were independently associated with less improvement in VA at 1 year in that study. Similarly, another recent report found no difference in VA outcome for occult, minimally classic or predominantly classic lesions in the PIER study.

The rate of serious adverse events was consistent with previous experience. Infectious endophthalmitis occurred in 2 patients, an incidence of 2.2 per 10,000 injections. Non-infectious endophthalmitis was reported in 2 more cases. Retinal detachment occurred in 1 eye, an incidence of 1.1 per 10,000 injections; this is similar to the rate at which retinal detachments are reported to occur in the general population. Mild adverse events appear to be under-reported, since there were only 45 episodes of post-injection pain. This indicates that registries may not accurately track outcomes that clinicians do not believe are clinically significant.

This study, like all observational studies, has some limitations arising from the way in which data were collected. Subjective criteria such as lesion activity or lesion type may not be uniformly graded in observational studies since they are reported by the treating physicians rather than a centralised Reading Centre. Thus these determinations may have lower internal validity than in a phase 3 clinical trial, but perhaps they are still meaningful since this is how these clinically important determinations are actually being made in the real world. The measurement of LogMAR visual acuity, the main outcome, is reasonably objective. Also, case selection and treatment regimens in observational studies may be very different to clinical trials and among different ophthalmologists. Nevertheless, the data presented show
generally consistent outcomes of treatment regimens, which appeared to be similar across the
different centres (data not shown).

There are a number of further analyses that can be performed on observational data that we
present here. A study of the efficacy of different treatment intensities will need to take into
account “treatment by time” interactions (a treatment in the first 3 months is likely to have a
greater affect than a treatment in the last 3 months) and the possibility that the outcome of
treatment drives treatment intensity, with eyes responding poorly receiving more treatments
than these that respond well, rather than vice versa. A study of poor responders would need to
include not just the proportion of patients who, for example, lose 15 letters, but also analysis
of their baseline characteristics, how the loss evolved over time and whether the causes could
be identified by a case by case analysis referring back to the clinic notes in a selected
subgroup. Treatment patterns and their different efficacies can also be identified: a pro re
nata regimen will be revealed when treatments are given only when the lesion is graded as
active, while a treat and extend regiment will have most treatments given when the lesion is
graded as inactive.

The significance of data from observational studies is that they provide an indication of what
is happening in routine clinical practice, in contrast to results of phase 3 clinical trials, which
may or may not be achievable in general. The results we present of intravitreal therapy for
wet AMD are reasonably good, at least in the Australian centres that chose to participate.
Further research is warranted to determine the functional implications of persistent activity
and whether cohorts of patients receiving routine treatment do as well as those in phase 3
studies when they are more closely matched to participants in those studies.
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REFERENCES


Figure 1: Density plots of visual acuity (A) and greatest linear dimension (B) at the index visit by lesion type.

Figure 2: Density plot of within group changes at 12 months (A) and fitted Lowess lines showing subgroup changes in VA over 12 months (B)

Figure 3: Kaplan-Meier plot of time from active lesion first being graded as inactive.

Figure 4: Fitted Lowess lines showing changes in VA over 12 months for non-completers (A,B,C) and completers (D).