Impact of intravitreal aflibercept dosing regimens in treatment-naïve patients with neovascular age-related macular degeneration in routine clinical practice in France: results from the RAINBOW study

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ABSTRACT

Objective To evaluate 12-month outcomes in treatment-naïve patients with neovascular (wet) age-related macular degeneration (AMD) stratified by intravitreal aflibercept (IVT-AFL) regimen.

Methods and analysis Patients included in the 12-month interim analysis of Real life of intravitreal Aflibercept in FraNce: observational Study in Wet AMD (RAINBOW), a 4-year, ongoing observational study conducted in France, were stratified by IVT-AFL dosing regimen. Safety (n=593) and effectiveness (n=428) data were analysed. Regimens included a regular cohort (three initial monthly IVT-AFL injections and ≥6 injections) and irregular cohorts (<6 injections) with and without three initial monthly injections. The main outcome measure was mean gain in best-corrected visual acuity (BCVA) at 12 months.

Results Mean number of IVT-AFL injections was 6.0 (all patients, n=513), 7.2 (regular cohort, n=102), 6.1 (irregular cohort with three initial monthly injections, n=266) and 5.2 (irregular cohort without three initial monthly injections, n=60). Overall mean gain in BCVA at 12 months was 5 letters; +7.1 letters (regular cohort) and +5.6 letters (irregular cohort with three initial monthly injections), both p<0.001 versus baseline, and −1.1 letters (irregular cohort without three initial monthly injections), p=0.669. Improvements in BCVA were also significantly greater in the regular cohort (p<0.001) and irregular cohort with three initial monthly injections (p=0.003) compared with the irregular cohort without three initial monthly injections. Ocular and non-ocular adverse events were reported in 14.7% and 17.4% of all patients, respectively.

Conclusion Treatment-naïve patients with neovascular AMD receiving three initial monthly injections followed by regular or irregular injections over 12 months experienced better visual acuity outcomes than those receiving irregular treatment without three initial monthly injections.

Trial registration number NCT02279537.

Key messages

What is already known about this subject?
▸ An initial analysis of the RAINBOW real-world study, reported improved visual and anatomical outcomes in treatment-naïve patients with neovascular age-related macular degeneration (AMD) treated with intravitreal aflibercept over a 12-month period.

What are the new findings?
▸ This real-world study shows that treatment-naïve patients with neovascular AMD receiving three initial monthly injections of intravitreal aflibercept followed by regular or irregular injection intervals over 12 months experienced better visual acuity outcomes than those receiving irregular treatment without three initial monthly injections.

How might these results change the focus of research or clinical practice?
▸ These findings suggest that intravitreal aflibercept treatment with three initial monthly injections, followed by regular treatment, offers the best possible outcomes for patients with neovascular AMD in the real-world setting.

INTRODUCTION

Age-related macular degeneration (AMD) is a common degenerative disease of the retina and the leading cause of legal blindness in the elderly worldwide.1 There are two forms of AMD: dry and wet. The dry form, which is characterised by drusen, pigmentary abnormalities and atrophy, accounts for 90% of cases and 10% of AMD-related blindness.2 The wet (neovascular) form is more aggressive, and accounts for 10% of cases and 90% of AMD-related blindness and severe visual impairment.2 Neovascular AMD has a significant impact on the quality of life of patients.1 Choroidal neovascularisation (CNV) is the most common cause of vision loss due to neovascular AMD; it is triggered...
by angiogenic factors such as vascular endothelial growth factor-A (VEGF-A) and placental growth factor (PGF). The anti-VEGF agent ranibizumab is a monoclonal antibody that inhibits VEGF-A. Randomised studies conducted between 2003 and 2004 have demonstrated the benefits of monthly ranibizumab (0.5 mg) dosing in patients with neovascular AMD. However, such frequent dosing is a burden for patients, caregivers and the healthcare system. Treatment regimens with fewer ranibizumab injections have subsequently been evaluated in studies where ranibizumab was given as needed (pro re nata (PRN)). The Prospective OCT study with Lucentis for neovascular AMD (PrONTO Study; 2004–2005), study of ranibizumab in patients with subfoveal CNV secondary to AMD (SUSTAIN; 2006–2008) and Inhibition of VEGF in Age-related CNV (IVAN; 2008–2010) studies showed that efficacy outcomes could be achieved with less than monthly dosing. Others, such as the study of ranibizumab administered monthly or on an as-needed basis in patients with subfoveal neovascular AMD (HARBOR; 2009–2010), showed that a PRN regimen is slightly less favourable than a monthly dosing regimen. In a real-world setting, the ranibizumab PRN regimen was associated with undertreatment and a decline in visual acuity (VA) over 12 months.

Intravitreal aflibercept (IVT-AFL) is a fusion protein that consists of human VEGF receptors 1 and 2 combined with the fragment crystallizable (Fc) portion of IgG. It binds VEGF-A and PGF with a higher affinity than other naturally occurring VEGF receptors, so it has the potential for better efficacy. In the randomised VEGF Trap-Eye: Investigation of Efficacy and Safety in Wet AMD (VIEW) studies conducted between 2007 and 2011, IVT-AFL 2 mg administered every 8 weeks (2q8; after three initial monthly IVT-AFL injections) was non-inferior to the approved ranibizumab dosing schedule (0.5 mg every 4 weeks) in all clinical endpoints at week 52 without any marked difference in adverse events (AEs). Based on previous studies, IVT-AFL was approved with a dosing schedule in the EU of three initial monthly injections of 2 mg followed by 2q8 for the first 12 months. After 12 months, injection intervals can be extended based on visual and anatomical outcomes. Outcomes from clinical practice settings showed that using this practical IVT-AFL dosing regimen, a 12-month visual gain of five letters or more was achieved, which is better than many previous real-world data collections. Prior to the approval of IVT-AFL in Europe in 2012, PRN dosing was the most common regimen to treat neovascular AMD in France, based on the availability of ranibizumab since 2007. Approval of IVT-AFL may have led to variations in the IVT-AFL treatment protocol/regimen used in clinical practice as physicians adapted their routine clinical practices away from the previously reactive approach. Consequently, intervals between injections may have varied substantially. In addition, we have also previously shown that not all patients receive the approved three initial doses of IVT-AFL.

The ongoing Real life of intravitreal Aflibercept In FraNce: oBservatiOnal Study in Wet AMD (RAINBOW) study was initiated in 2014, with the objective of monitoring the outcomes achieved with IVT-AFL in treatment-naïve patients with neovascular AMD treated in routine clinical practice. The aim of this interim analysis of the RAINBOW study was to monitor treatment outcomes in patients with neovascular AMD stratified by IVT-AFL dosing regimen (identified retrospectively) over the first 12 months of treatment: a regular cohort of patients who received the indicated posology of three initial monthly injections of IVT-AFL followed by regular injections every 2 months, with 26 injections given during the 12 months; and two irregular cohorts who received <6 injections of IVT-AFL, either with or without three initial monthly injections. The aim of the stratification was to assess if there were any variations in the IVT-AFL treatment protocol and their impact on outcomes.

MATERIALS AND METHODS

Study design

RAINBOW is an ongoing, observational, multicentre, ambispective (retrospective and prospective) study to monitor the real-world effectiveness and safety of IVT-AFL in treatment-naïve patients with neovascular AMD enrolled in 55 centres across France. The study began in October 2014. Data from patients who started IVT-AFL treatment between 2 January and 13 October 2014 were retrospectively collected, and data from patients who started IVT-AFL treatment after 14 October 2014 were prospectively collected. This interim analysis used 12-month data collected up to 9 January 2017. However, patients will be followed for a period of 4 years or until study discontinuation.

Participants

Patients aged ≥50 years diagnosed with neovascular AMD and treated with IVT-AFL by their physician were eligible. Prior or current treatment with any anti-VEGF agent or macular laser in the study eye was not allowed. For patients receiving treatment only in one eye, the study eye was defined as the worst-seeing eye of each patient, and the second eye was considered as treatment naïve. For patients receiving treatment in both eyes, the first eye to be treated was considered the study eye; if both eyes were treated at the same time, the worst-seeing eye or right eye (if similar) was considered the study eye. Patients were excluded if they did not meet the indication criteria for IVT-AFL, if they had another retinal disease (diabetic retinopathy, diabetic macular oedema, myopic CNV, retinal vein occlusion, central serous chorioretinopathy, or angioid streaks), or if they were participating in any other interventional study.

Outcomes

This is an interim analysis to evaluate 12-month outcomes in treatment-naïve patients with neovascular AMD.
stratified by IVT-AFL regimen. The primary objective was to evaluate the difference in best-corrected visual acuity (BCVA), assessed preferentially by the early treatment diabetic retinopathy study (ETDRS) protocol or any visual logarithmic scale, from baseline to 12 months (range: 11–13). We converted VA readings to a standardised score using a conversion chart (online supplementary table 1).

Other assessments evaluated included the percentage of patients who gained ≥20, ≥5, ≥10, or ≥15 letters, or lost >15 letters at 12 months; the proportion of patients with BCVA ≥70 letters at 12 months; and mean change in central retinal thickness (CRT) at month 12. All AEs occurring after the first injection of IVT-AFL (up to the 30-day period after the last IVT-AFL injection) were documented.

Data sources and measurements
For retrospectively enrolled patients, we obtained treatment(s) and outcomes from medical records. For prospectively enrolled patients, we recorded this information at routine follow-up visits. We recorded data in an electronic case report form from the inclusion date prospectively enrolled patients, we recorded this information at routine follow-up visits. We recorded data in an electronic case report form from the inclusion date.

Statistical analyses
Approximately 600 patients needed to be enrolled to achieve around 390 usable patient data sets at 48 months, with an expected precision of 1.5 letters. This included an estimated 10% loss to follow-up per year with 5% unevaulable data. We based these assumptions on the results from the VIEW studies, in which the mean (SD) difference in BCVA (letters) from baseline to 12 months was 7.9 (15.0) letters (VIEW 1) and 8.9 (14.4) letters (VIEW 2). The formula we used to derive these estimates was:

\[ n = \frac{(1.96)^2 \times SD^2}{i^2} \]

where \( i^2 = \) estimated precision

\[ \sqrt{n} = 1.96 \times \frac{15}{13} = 19.6 \rightarrow n \approx 384 \]

The study was not designed to confirm or reject any hypotheses. Interim analyses were predefined in the study protocol at yearly intervals. Statistical analyses were exploratory and descriptive in nature. We used summary statistics for categorical and quantitative (continuous) data. We described continuous data by non-missing values, mean, SD and 95% CI. We generated frequency tables for categorical data.

All AEs were coded using the latest Medical Dictionary for Regulatory Activities V.21. We performed the analysis of the primary assessment without replacement of missing values. However, we performed two sensitivity analyses with two imputation methods for missing data (last observation carried forward (LOCF) approach, and imputation of missing data by the median value of the population). These sensitivity analyses were devised to deal with any potential bias due to premature dropouts.

The LOCF method was assumed to be more conservative as premature dropouts were expected to have poorer results than patients completing study treatment.

Safety analyses were based on data from patients who received at least one IVT-AFL injection (safety analysis set (SAS)). Patients with two treated eyes were included twice in this population. Effectiveness analyses were based on patients who also had documented assessments in the study eye at baseline and at least one follow-up visit (full analysis set (FAS)), which included one eye per patient. The primary outcome analyses were based on patients who had documented VA assessments at baseline and month 12 (FAS targeted).

Patients were retrospectively stratified by IVT-AFL dosing regimen over the first 12 months as follows: (1) regular cohort: patients who received three initial monthly IVT-AFL injections separated by 23–44 days and regular (≥6) injections separated by ≥39 to <88 days; (2) irregular cohort with three initial monthly injections: patients who received three initial monthly IVT-AFL injections separated by 23–44 days and irregular (<6) injections separated by <39 to ≥88 days; or (3) irregular cohort without three initial monthly injections: patients who received irregular (<6) injections separated by <39 to >88 days without three initial monthly IVT-AFL injections. The three treatment cohorts evaluated in this study were derived retrospectively from the data generated within it, to assess the uptake among clinicians of the newly approved dosing regimen compared with the previous more commonly used PRN treatment modality used in the management of neovascular AMD. We compared these three treatment regimens using an analysis of variance model with treatment regimen as a fixed factor and BCVA at baseline as a covariate. The statistical analysis was performed with the software package SAS release V.9.4 (SAS Institute Inc).

RESULTS
Patient disposition and baseline characteristics
A total of 608 patients were enrolled; 593 patients (737 treated eyes) were included in the SAS and 513 patients were included in the FAS, of whom 428 comprised the FAS targeted population (see figure 1). At the time of the interim analysis, 120 patients had prematurely withdrawn. The baseline characteristics were generally similar between treatment groups (table 1).

Treatments
The mean (SD) number of IVT-AFL injections over 12 months was 6.0 (2.1) in the FAS population (n=513). A total of 428 patients had BCVA assessments at baseline and 12 months, and 368 (86%) received three initial monthly IVT-AFL injections. These patients were stratified by IVT-AFL regimen into the regular cohort (n=102), irregular cohort with three initial monthly injections (n=266) and irregular cohort without three initial monthly injections (n=60). The mean (SD) number of injections was higher in the regular cohort (7.2 (0.8)) versus the irregular
cohort with three initial monthly IVT-AFL injections (6.1 (2.2)) and irregular cohort without three initial monthly IVT-AFL injections (5.2 (1.8)), although the duration of follow-up was comparable between the three treatment groups (online supplementary table 2).

**Visual acuity outcomes**
Mean (SD) BCVA increased from 57.5 (18.3) letters at baseline to 62.5 (18.8) letters at 12 months, resulting in a statistically significant mean (SD) gain of 5.0 (16.0) letters in all patients (p<0.001 vs baseline; figure 2). The results were confirmed by two sensitivity analyses conducted in the FAS population (n=513).

Mean (SD) BCVA increased from 59.8 (16.5) letters at baseline to 67.0 (14.7) letters at 12 months in the regular cohort with three initial monthly injections, resulting in a mean (SD) gain of 7.1 (11.8) letters and from 56.5 (18.5) to 62.1 (18.9) letters for patients from the irregular cohort with three initial monthly injections at 12 months, resulting in a gain of 5.6 (16.4) letters (both p<0.001 vs baseline). There was a gradual decline from baseline in mean (SD) BCVA to −1.1 (18.9) letters (from 57.6 (20.0) to 56.5 (22.9) letters) at 12 months in the irregular cohort without three initial monthly injections (p=0.669). Between-group comparisons showed that the improvements in BCVA were significantly greater in the regular cohort (p<0.001) and irregular cohort with three initial monthly injections (p=0.003) compared with the irregular cohort without three initial monthly injections (figure 2).

Gains of ≥15 letters were observed in 25.2% (all patients), 22.5% (regular cohort), 28.6% (irregular cohort with three initial monthly injections) and 15.0% (irregular cohort without three initial monthly injections) at 12 months, respectively. Losses of >15 letters were observed in 6.3% (all patients), 1.0% (regular cohort), 6.8% (irregular cohort with three initial monthly injections) and 13.3% (irregular cohort with three initial monthly injections) at 12 months. At 12 months, 45.6% (all patients), 54.9% (regular cohort), 43.6% (irregular cohort with three initial monthly injections) and 38.3% (irregular cohort without three initial monthly injections) could read ≥70 letters (figure 3).

**Anatomical outcomes**
The mean reductions in CRT from baseline to 12 months were −110.4 µm (all patients: baseline, 400.5 µm; 12 months, 290.1 µm), −133.7 µm (regular cohort: baseline, 400.8 µm; 12 months, 267.1 µm) and −113.9 µm (irregular cohort with three initial monthly injections: baseline, 403.2 µm; 12 months, 289.3 µm; all p<0.001 vs baseline). There was a smaller reduction in the irregular cohort without three initial monthly injections (−46.2 µm; baseline, 381.6 µm; 12 months, 335.4 µm; p=0.036) at 12 months.

**Safety**
Safety outcomes are summarised in table 2. One-hundred-and-seventy patients (28.7% of the treated population) declared at least one AE. In total, 287 AEs were reported during the period of interest: 144 ocular events were experienced by 14.7% (n=87) of patients and 143 non-ocular events by 17.4% (n=103) of patients. Thirty-five events were considered related to IVT-AFL by the investigator: 31 ocular events were experienced by 4.6% (n=27) of patients and four non-ocular events by 0.7% (n=4) of patients.
patients. Twenty-three ocular events and 17 non-ocular events were responsible for a treatment discontinuation in 31 patients; 17 of these events, all ocular, were considered to be related to the treatment according to the physician. A total of 60 serious AEs occurred in 7.8% (n=46) of patients when treated with IVT-AFL and one patient presented one serious AE when receiving other treatments. Overall, the most common treatment-related ocular AEs were vitreous floaters (1.9% of patients), lack of effectiveness (1.3%), eye pain (1.2%) and increased lacrimation (1.2%). Four patients died during the follow-up period.

**DISCUSSION**

The ongoing RAINBOW study is monitoring resource use and outcomes with IVT-AFL in treatment-naïve patients with neovascular AMD in France. The aim of this interim analysis was to monitor 12-month outcomes by evaluating three different IVT-AFL regimens. Patients were treated in accordance with the investigator’s normal clinical practice; these patients were then retrospectively divided into three cohorts based on the treatment protocol utilised (as described). Although some variability in the IVT-AFL treatment protocol was observed in this real-world setting, of the 593 patients who received at least one injection of IVT-AFL, 368 of 428 (86%) received three initial monthly IVT-AFL injections. Variability in the treatment protocols used with IVT-AFL is likely due to the option of anti-VEGF administration using either a reactive (PRN) or proactive treatment approach. Although both approaches incorporate three initial monthly injections, our observation that this does not occur in all patients, is not unique and has been seen previously with IVT-AFL and ranibizumab (with both a fixed-dose or treat-and-extend dosing).23–25 This, therefore, provides the rationale to evaluate the impact of any differences in treatment protocols on patient outcomes in clinical practice.
In the pooled analysis of the VIEW studies, the mean change in BCVA was 8.4 letters, the proportion of patients gaining ≥15 letters was 30.9% and the percentage of patients maintaining VA (losing <15 letters) was 95.3% at month 12 after a mean of 7.5 IVT-AFL injections in the combined IVT-AFL 2q8 group after three initial monthly injections.17 18 In this study, the overall results showed a mean visual gain of five letters, and were closer to those seen in the VIEW studies when the regimen included three monthly injections and a regular follow-up with a gain of 7.1 letters. Furthermore, the secondary endpoints of our entire cohort were very similar to the ones of the VIEW studies, with gains of ≥15 letters observed in 25.2% of patients and losses of >15 letters observed in only 6.3%.

When examining BCVA at 12 months for the overall population (62.5 letters), it is noticeable how similar this is to those values observed at 12 months in the VIEW 1 (63.6 letters) and VIEW 2 (60.5 letters) studies.

Furthermore, the present results were similar to other real-world series using IVT-AFL as first-line therapy. In a retrospective real-world analysis of 1840 treatment-naïve eyes diagnosed with neovascular AMD using the same IVT-AFL treatment protocol that was used in the VIEW studies, the mean BCVA gain at 12 months was 5.1 letters, with a mean of seven IVT-AFL injections.22 Similar improvements in BCVA were reported in another retrospective real-world analysis conducted in the United Kingdom: 109 eyes of 102 patients diagnosed with neovascular AMD and treated with aflibercept were analysed (fixed dosing year 1 followed by a treat-and-extend approach). The mean BCVA gain (letters) was 5.9 (year 1), 6.4 (year 2) and 6.6 (year 3), and the corresponding median number of injections was 8 in year 1, and 4 in years 2 and 3.20 A prospective analysis (prospective non-interventional study to assess the effectiveness of aflibercept in routine clinical practice in patients with neovascular AMD (PERSEUS)) conducted in Germany indicated that outcomes in treatment-naïve patients were also more favourable in those receiving regular IVT-AFL injections (mean VA gain: 8.0 letters) than the irregularly treated population at 12 months (mean VA gain: 4.0 letters),26 highlighting the need for consistent and regular treatment regimens to obtain optimal outcomes as seen in clinical studies.

Variations were observed in the administration of IVT-AFL treatment in France, which differed from the indicated treatment regimen. One possible explanation is that prior to the approval of IVT-AFL, PRN was the most common regimen to treat neovascular AMD in France, and some physicians may have continued to use this regimen. Of note, phase 3 studies have not assessed the efficacy of IVT-AFL without the three indicated initial monthly injections. Our study did not assess the reasons for the lack
of compliance with the initial three-monthly injections; however, a lack of adherence to, or persistence with, anti-VEGF therapy has been evaluated in other studies. These have identified various factors including system-related factors such as difficulties in planning monthly injections or scheduling patient visits, as well as patient and disease-related characteristics. For example, a multinational study reported that patients missed clinic visits as the caregiver was unable to take them to the appointment, because of fear about receiving an injection, or because of illness. A study in France reported patients frequently missed treatment appointments over the summer vacation period, to the detriment of outcomes, and highlighted the importance of explaining to patients the need for treatment adherence, despite their need for holidays.

Other real-world studies reported the results of treatment with ranibizumab in patients with neovascular AMD in Europe, using a PRN regimen in the years 2006–2011. In LUMIERE, an observational study of ranibizumab use in French clinical practice, conducted from 2006 to 2009,

<table>
<thead>
<tr>
<th>AE, n (%)</th>
<th>SAS (n=593)</th>
<th>Regular cohort (n=102)</th>
<th>Irregular cohort with treatment initiation (n=266)</th>
<th>Irregular cohort without treatment initiation (n=60)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Any AE</td>
<td>170 (28.7)</td>
<td>22 (21.6)</td>
<td>77 (28.9)</td>
<td>12 (20.0)</td>
</tr>
<tr>
<td>Ocular</td>
<td>87 (14.7)</td>
<td>15 (14.7)</td>
<td>48 (18.0)</td>
<td>4 (6.7)</td>
</tr>
<tr>
<td>Non-ocular</td>
<td>103 (17.4)</td>
<td>11 (10.8)</td>
<td>42 (15.8)</td>
<td>8 (13.3)</td>
</tr>
<tr>
<td>Any treatment-related AE</td>
<td>31 (5.2)</td>
<td>5 (4.9)</td>
<td>17 (6.4)</td>
<td>1 (1.7)</td>
</tr>
<tr>
<td>Ocular</td>
<td>27 (4.6)</td>
<td>4 (3.9)</td>
<td>16 (6.0)</td>
<td>1 (1.7)</td>
</tr>
<tr>
<td>Non-ocular</td>
<td>4 (0.7)</td>
<td>1 (1.0)</td>
<td>1 (0.4)</td>
<td>0</td>
</tr>
</tbody>
</table>

### Most common (>1%) treatment-related ocular AEs occurring in any treatment group

<table>
<thead>
<tr>
<th>AE</th>
<th>SAS (n=593)</th>
<th>Regular cohort (n=102)</th>
<th>Irregular cohort with treatment initiation (n=266)</th>
<th>Irregular cohort without treatment initiation (n=60)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Vitreous floaters</td>
<td>11 (1.9)</td>
<td>3 (2.9)</td>
<td>6 (2.3)</td>
<td>–</td>
</tr>
<tr>
<td>Drug ineffective</td>
<td>8 (1.3)</td>
<td>1 (1.0)</td>
<td>5 (1.9)</td>
<td>–</td>
</tr>
<tr>
<td>Eye pain</td>
<td>7 (1.2)</td>
<td>–</td>
<td>4 (1.5)</td>
<td>1 (1.7)</td>
</tr>
<tr>
<td>Lacrimation increased</td>
<td>7 (1.2)</td>
<td>2 (2.0)</td>
<td>5 (1.9)</td>
<td>–</td>
</tr>
<tr>
<td>Metamorphopsia</td>
<td>6 (1.0)</td>
<td>1 (1.0)</td>
<td>3 (1.1)</td>
<td>–</td>
</tr>
<tr>
<td>Visual acuity reduced</td>
<td>5 (0.8)</td>
<td>1 (1.0)</td>
<td>3 (1.1)</td>
<td>–</td>
</tr>
<tr>
<td>Vision blurred</td>
<td>5 (0.8)</td>
<td>1 (1.0)</td>
<td>3 (1.1)</td>
<td>–</td>
</tr>
<tr>
<td>Eye allergy</td>
<td>5 (0.8)</td>
<td>1 (1.0)</td>
<td>–</td>
<td>1 (1.7)</td>
</tr>
<tr>
<td>Injection-site pain</td>
<td>4 (0.7)</td>
<td>–</td>
<td>3 (1.1)</td>
<td>–</td>
</tr>
<tr>
<td>Photophobia</td>
<td>–</td>
<td>–</td>
<td>3 (1.1)</td>
<td>–</td>
</tr>
<tr>
<td>Choroidal neovascularisation</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>1 (1.7)</td>
</tr>
<tr>
<td>Retinal oedema</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>1 (1.7)</td>
</tr>
<tr>
<td>Retinal vein occlusion</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>1 (1.7)</td>
</tr>
<tr>
<td>Any treatment-emergent SAE</td>
<td>46 (7.8)</td>
<td>7 (6.9)</td>
<td>18 (6.8)</td>
<td>4 (6.7)</td>
</tr>
<tr>
<td>Ocular</td>
<td>4 (0.7)</td>
<td>2 (2.0)</td>
<td>2 (0.8)</td>
<td>0</td>
</tr>
<tr>
<td>Non-ocular</td>
<td>42 (7.1)</td>
<td>5 (4.9)</td>
<td>16 (6.0)</td>
<td>4 (6.7)</td>
</tr>
<tr>
<td>Treatment-related ocular SAE</td>
<td>1 (0.2)</td>
<td>1 (1.0)</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Traumatic cataract</td>
<td>4 (0.7)</td>
<td>–</td>
<td>3 (1.1)</td>
<td>–</td>
</tr>
<tr>
<td>Peripheral ischaemic attack</td>
<td>1 (0.2)</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Any AE leading to discontinuation</td>
<td>31 (5.2)</td>
<td>4 (3.9)</td>
<td>17 (6.4)</td>
<td>1 (1.7)</td>
</tr>
<tr>
<td>Ocular</td>
<td>20 (3.4)</td>
<td>2 (2.0)</td>
<td>12 (4.5)</td>
<td>1 (1.7)</td>
</tr>
<tr>
<td>Non-ocular</td>
<td>13 (2.2)</td>
<td>2 (2.0)</td>
<td>7 (2.6)</td>
<td>0</td>
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<tr>
<td>Any treatment-related AE leading to discontinuation</td>
<td>14 (2.4)</td>
<td>1 (1.0)</td>
<td>8 (3.0)</td>
<td>1 (1.7)</td>
</tr>
<tr>
<td>Ocular</td>
<td>14 (2.4)</td>
<td>1 (1.0)</td>
<td>8 (3.0)</td>
<td>1 (1.7)</td>
</tr>
<tr>
<td>Non-ocular</td>
<td>0 (0.0)</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Death (not treatment-related)</td>
<td>4 (0.7)</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
</tbody>
</table>

Values in the cohort columns are based on the FAS population.

*AE rates<0.5% are not shown.

AE, adverse event; FAS, full analysis set; SAE, serious adverse event; SAS, safety analysis set.
the mean BCVA change from baseline to month 12 was 3.2 letters in patients receiving a mean of 5.1 injections, with fewer than 40% of patients receiving the recommended treatment of three initial monthly injections, suggesting that failure to follow the recommended dosing was associated with lower than anticipated efficacy. In TWIN, a follow-up conducted from 2010 to 2011, 56.6% of patients received the recommended treatment of three initial monthly injections (mean of 5.6 ranibizumab injections over 12 months), with a slightly better improvement in VA (gain of 4.3 letters). These studies, like ours, reinforce the importance of following the approved treatment protocol of three initial monthly IVT-AFL injections.

Analysis of UK medical records to compare the effectiveness of IVT-AFL in proactive regimens (fixed or treat-and-extend) versus ranibizumab in reactive regimen (PRN) found that patients treated with IVT-AFL achieved greater VA gains at 1 year (+6.1 ETDRS letters) than those treated with ranibizumab (+1.6 ETDRS letters, p=2.2e−16). The authors attributed the observed difference to the dosing regimen rather than the anti-VEGF agent used. However, it should be noted that, although the IVT-AFL group had significantly more injections than the ranibizumab group (7.0 vs 5.8; p<2.2e−16), they required fewer clinic visits (9.0 vs 10.8; p<2.2e−16).

The safety profile of IVT-AFL in the real-life RAINBOW study was consistent with that reported in previous studies. Of the 31 patients with treatment-related AEs, three were serious (transient ischemic attack, peripheral arterial occlusive disease and traumatic cataract). Four deaths were recorded; none were considered related to IVT-AFL. No new safety signal was seen.

There are some limitations to this observational study; the visit schedule was at the discretion of the treating physician, which resulted in missing data at time points throughout the study. To address this, we included some flexibility around the initial dosing period, dosing intervals and final BCVA assessment. In addition, two sensitivity analyses were conducted with two different imputation methods for missing values (LOCF and median); the results remained consistent across these analyses. This interim analysis was also exploratory, and the results should be interpreted with caution; notably, allocation to treatment schedules was not randomised and no analysis of disease severity that could influence treatment regimen was carried out. However, noting the earlier limitations, given that the data sources were from 55 ophthalmological centres utilising real-world practice, our overall findings could be generalised to wider practice.

In conclusion, this second interim analysis of the RAINBOW study confirms the preliminary results, which demonstrated that IVT-AFL was associated with visual and anatomical improvements over 12 months under real-world conditions in France. Our results emphasise the benefit that treatment-naïve patients with neovascular AMD obtained with a first-line treatment with IVT-AFL in real life, with a clinically meaningful mean VA gain of five letters at 1 year, and even greater gains when three initial monthly injections were given. Thus, even though the study was not randomised, the results suggest that better results can be obtained with a regular treatment in accordance with the summary of product characteristics (ie, three initial injections within 23–44 days and further injections spaced by no more than 88 days) for the first year. Results from this study should help physicians understand the overall importance of three initial monthly IVT-AFL injections followed by regular treatment during the first year with the goal of maximising patient outcomes.

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Data availability statement Availability of the data underlying this publication will be determined according to Bayer’s commitment to the EFPIA/PhRMA “Principles for responsible clinical trial data sharing”. This pertains to scope, time point and process of data access. As such, Bayer commits to sharing upon request from qualified scientific and medical researchers patients-level clinical trial data, study-level clinical trial data, and protocols from clinical trials in patients for medicines and indications approved in the United States (US) and European Union (EU) as necessary for conducting legitimate research. This applies to data on new medicines and indications that have been approved by the EU and US regulatory agencies on or after January 01, 2014. Interested researchers can use to request access will be granted to anonymized patient-relevant information is provided in the Study sponsors section of the portal. Data access will be granted to anonymized patient-level data, protocols and clinical study reports after approval by an independent scientific review panel. Bayer is not involved in the decisions made by the independent review panel. Bayer will take all necessary measures to ensure that patient privacy is safeguarded.

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