Multi-country real-life experience of anti-vascular endothelial growth factor therapy for wet age-related macular degeneration

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ABSTRACT
Background/aims Real-life anti-vascular endothelial growth factor (VEGF) therapy use in patients with wet age-related macular degeneration (wAMD) was assessed in a retrospective, observational study in Canada, France, Germany, Ireland, Italy, the Netherlands, UK and Venezuela.

Methods Medical records of patients with wAMD, who started ranibizumab treatment between 1 January 2009 and 31 August 2009, were evaluated. Data were collected until the end of treatment and/or monitoring or until 31 August 2011.

Results 2227 patients who received ≥1 anti-VEGF injection with a baseline visual acuity assessment and ≥1 postbaseline visual acuity assessment for the treated eye were evaluated. Visual acuity improved until about day 120; thereafter, visual acuity gains were not maintained. Mean change in visual acuity score from baseline to years 1 and 2 was +2.4 and +0.6 letters, respectively. Patients received a mean of 5.0 and 2.2 injections in the first and second year, respectively. There were substantial differences in visual outcomes and injection frequency between countries. More frequent visits and injections were associated with greater improvements in visual acuity.

Conclusions In clinical practice, fewer injections are administered than in clinical trials. Anti-VEGF treatment resulted in an initial improvement in visual acuity; however, this was not maintained over time.

Trial registration number NCT01447043.

INTRODUCTION
Neovascular (wet) age-related macular degeneration (wAMD) is a progressive degenerative disease of the central retina.1 2 The vascular endothelial growth factor (VEGF) antibody fragment ranibizumab was one of the first pharmacological agents proven to be beneficial in the treatment of wAMD.

Ranibizumab was approved for the treatment of wAMD based on results from two phase III trials: ANCHOR (patients with predominantly classic choroidal neovascularisation (CNV)) and MARINA (patients with minimally classic or occult CNV).3 4 Ranibizumab treatment resulted in improvements in visual acuity which were maintained with monthly treatment, resulting in visual acuity gains at month 12 of 8.5–11.3 letters in ANCHOR and 6.5–7.2 letters in MARINA.3 4 Monthly intravitreal injections are associated with a significant treatment burden for patients, caregivers and physicians, often making such a regimen unachievable in clinical practice. To reduce management burden, less frequent dosing regimens of ranibizumab (quarterly or pro re nata (PRN; as needed)) have been evaluated, but these usually have been associated with slightly less favourable outcomes than monthly dosing.5–7 In the HARBOR1 study, patients who received PRN treatment after three initial monthly doses had a numerically smaller gain in vision at month 12 than those continuing monthly treatment.3 However, the SUSTAIN8 and IVAN9 studies indicated that efficacy outcomes could be achieved with less than monthly dosing.9 10 A treat-and-extend regimen has also been used.11 12 In Europe, ranibizumab is licensed for monthly dosing until visual acuity is stable, followed by monitoring and resumption of treatment as needed.13 In the USA, ranibizumab once monthly is recommended; however, patients may receive three or four monthly doses followed by less frequent dosing with regular assessments.14

We report results from AURA8*, an international, retrospective study that assessed management of patients with wAMD receiving anti-VEGF treatment in clinical practice between 2009 and 2011.

METHODS
Study design
AURA was a retrospective, observational, multicentre study conducted in Canada, France, Germany, Ireland, Italy, the Netherlands, UK and Venezuela.

Patients with wAMD, who started treatment with ranibizumab between 1 January 2009 and 31 August 2009, were consecutively screened for eligibility. Written consent was obtained from each patient prior to inclusion where applicable. Approval from the relevant independent ethics committees (or their equivalents) was obtained prior to study commencement outside of any country where such approval is required by law. Patients who started treatment with an alternative anti-VEGF agent were also included. Patients could have a second wet AMD lesion if the first lesion was not treated before the start of treatment.

The study was conducted according to the tenets of the Declaration of Helsinki and Good Clinical Practice guidelines. Medical records were collected until the end of treatment and/or monitoring or until 31 August 2011.

Background
In 2009, ranibizumab was approved for the treatment of wAMD based on results from two phase III trials: ANCHOR (patients with predominantly classic choroidal neovascularisation (CNV)) and MARINA (patients with minimally classic or occult CNV).3 4 Ranibizumab treatment resulted in improvements in visual acuity which were maintained with monthly treatment, resulting in visual acuity gains at month 12 of 8.5–11.3 letters in ANCHOR and 6.5–7.2 letters in MARINA.3 4 Monthly intravitreal injections are associated with a significant treatment burden for patients, caregivers and physicians, often making such a regimen unachievable in clinical practice. To reduce management burden, less frequent dosing regimens of ranibizumab (quarterly or pro re nata (PRN; as needed)) have been evaluated, but these usually have been associated with slightly less favourable outcomes than monthly dosing.5–7 In the HARBOR1 study, patients who received PRN treatment after three initial monthly doses had a numerically smaller gain in vision at month 12 than those continuing monthly treatment.3 However, the SUSTAIN8 and IVAN9 studies indicated that efficacy outcomes could be achieved with less than monthly dosing.9 10 A treat-and-extend regimen has also been used.11 12 In Europe, ranibizumab is licensed for monthly dosing until visual acuity is stable, followed by monitoring and resumption of treatment as needed.13 In the USA, ranibizumab once monthly is recommended; however, patients may receive three or four monthly doses followed by less frequent dosing with regular assessments.14

We report results from AURA8*, an international, retrospective study that assessed management of patients with wAMD receiving anti-VEGF treatment in clinical practice between 2009 and 2011.

*HARBOR (The PHase III, double-masked, multicenter, randomized, Active treatment-controlled study of the efficacy and safety of 0.5 mg and 2.0 mg Ranibizumab administered monthly or on an as-needed Basis (PRN) in patients with subfoveal neOvAscularAge-related macular degeneration).

SUSTAIN (Study of Ranibizumab in Patients With Subfoveal Choroidal Neovascularization Secondary to Age-Related Macular Degeneration).

IVAN (A randomised controlled trial of alternative treatments to Inhibit VEGF in Age-related choroidal Neovascularisation).

AURA (A retrospective non-interventional study to assess the effectiveness of existing Anti-vascular Endothelial growth factor [anti-VEGF] treatment Regimens in patients with wet Age-related macular degeneration).
committees or institutional review boards and other national health authorities was received where required by local law and/or regulations.

Patients participating in an investigational study of any other drug or device while using anti-VEGF therapy were excluded. Patients must have received ≥1 ranibizumab injection to be included and were followed to the end of their treatment and/or monitoring or until 31 August 2011.

Study endpoints
The primary aim was to evaluate changes in visual acuity after the start of anti-VEGF therapy. Visual acuity was measured using Early Treatment Diabetic Retinopathy Study (ETDRS; equivalent notations) or Snellen (true Snellen fractions; where the numerator equals the test distance), depending on the centre. Thereafter, this was converted to the visual acuity scoring system (letter count; see online supplementary table S1).

Secondary objectives included determining anti-VEGF treatment regimens and disease monitoring in real-life settings.

Patients’ medical records and results from examinations performed during routine practice were evaluated.

Statistical analysis
It was calculated that 399 subjects per country would be required to estimate the change from a baseline score in visual acuity based on the ETDRS letter score, with a 95% probability of obtaining a CI with a width of not more than three letters. To allow for a 10% dropout rate, a sample size of 444 subjects per country was required.

Due to the exploratory nature of the study, all analyses are descriptive. The overall (exposed) population consisted of those patients who additionally had ≥1 postbaseline assessment of visual acuity for the treated eye. The first-year and second-year completer analysis sets included those in the effectiveness analysis set for whom follow-up data for at least 1 and 2 years after first injection, respectively, were documented.

To account for missing data, mean change in visual acuity was assessed using a last observation carried forward (LOCF) analysis. Mean change in visual acuity from baseline was calculated at the first and second year of the study. Secondary outcomes were analysed by frequency tables or summary statistics by visit.

RESULTS
Patients
Overall, 2227 patients were included in the effectiveness analysis set (see online supplementary figure S1). Follow-up was documented for 1695 patients for at least 1 year after the first injection (first-year completers’ set) and 1184 for at least 2 years after the first injection (second-year completers’ set). Baseline characteristics were similar across the analysis sets (table 1).

Visual acuity: change from baseline
In the effectiveness analysis set, an improvement in visual acuity was observed until about day 120 (figure 1). Thereafter, visual acuity noticeably decreased. Similar results were observed in the first-year and second-year completers’ sets. Mean change in visual acuity score from baseline to years 1 and 2 was +2.4 and +0.6 letters, respectively (effectiveness analysis set; LOCF analysis). In the first-year completers’ set, changes in visual acuity from baseline to years 1 and 2 were +2.7 and +0.3 letters, respectively. In the second-year completers’ set, changes in visual acuity from baseline to years 1 and 2 were +4.0 and +1.2 letters, respectively.

Patients who received an initial loading scheme (first three injections within 90 days) had greater improvements in visual acuity versus those who did not (figure 1), but the rate of decline in visual acuity thereafter did not seem associated with administration of loading doses. Baseline demographics were similar between patients who received loading doses and those who did not, including baseline visual acuity (data not shown).

<table>
<thead>
<tr>
<th>Table 1</th>
<th>Patient demographics at baseline (all analysis sets)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Overall (exposed set)</td>
</tr>
<tr>
<td>N</td>
<td>2609</td>
</tr>
<tr>
<td>Age at diagnosis, years (mean±SD)</td>
<td>76.8±8.4</td>
</tr>
<tr>
<td>Age at treatment start, years (mean±SD)</td>
<td>76.9±8.4</td>
</tr>
<tr>
<td>Women, n (%)</td>
<td>1593 (61.1)</td>
</tr>
<tr>
<td>Insurance*, n (%)</td>
<td>Public (2214 (84.9))</td>
</tr>
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<td></td>
<td>Private (183 (7.0))</td>
</tr>
<tr>
<td></td>
<td>None (186 (7.1))</td>
</tr>
<tr>
<td></td>
<td>Other (6 (0.2))</td>
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<tr>
<td></td>
<td>Missing (29 (1.1))</td>
</tr>
<tr>
<td>Angiographic findings</td>
<td>n=1951</td>
</tr>
<tr>
<td>No CNV</td>
<td>42 (2.4)</td>
</tr>
<tr>
<td>Classic CNV</td>
<td>550 (28.2)</td>
</tr>
<tr>
<td>Classic and occult</td>
<td>226 (11.6)</td>
</tr>
<tr>
<td>Occult</td>
<td>762 (39.1)</td>
</tr>
<tr>
<td>Disciform scar</td>
<td>11 (0.6)</td>
</tr>
<tr>
<td>Unknown</td>
<td>340 (17.4)</td>
</tr>
<tr>
<td>Missing</td>
<td>15 (0.8)</td>
</tr>
<tr>
<td>Visual acuity, letters (mean±SD)</td>
<td>NA</td>
</tr>
</tbody>
</table>

*Multiple responses possible.
CNV, choroidal neovascularisation; NA, not available.
Visits, visual acuity tests and optical coherence tomographies (OCTs)

The mean number of patient visits was lower in the second year than in the first year (effectiveness analysis set: 8.6 and 4.9 visits, respectively). Similar results were seen in the first-year completers’ set (9.5 and 6.4 visits, respectively) and second-year completers’ set (9.8 and 7.5 visits, respectively).

This was also reflected in the number of performed visual acuity tests and OCTs. In the effectiveness analysis set, the mean number of visual acuity tests per patient was 6.0 and 3.8 in the first and second years, respectively. Similar results were seen in the first-year completers’ set (6.7 and 5.0, respectively) and second-year completers’ sets (7.2 and 5.9). The mean number of OCTs was 4.5 in the first year and 3.2 in the second year of the effectiveness analysis set. Similar results were seen in the first-year completers’ set (5.2 and 4.3, respectively) and second-year completers’ sets (5.8 and 5.1).

Anti-VEGF injections

Most patients in the effectiveness analysis set did not have any treatment changes during the study and received ranibizumab throughout. A minority of patients also received treatment with bevacizumab (141 patients) and pegaptanib (7 patients). In the full 2 years, patients received a mean of 7.2 anti-VEGF injections each in the effectiveness analysis set. In the first-year and second-year completers’ sets, 8.4 and 9.1 injections, respectively, were received.

Patients received more anti-VEGF injections in the first year than the second year (effectiveness analysis set: mean of 5.0 and 2.2 injections, respectively). Similar results were observed in the first-year and second-year completer sets (5.5 and 2.9 injections in the first-year completers’ set; 5.6 and 3.5 injections in the second-year completers’ set).

Reasons for ending the study

A final visit before 31 August 2011 was documented in 68% of the effectiveness analysis set. The main reason for ending the study was discontinuation of anti-VEGF therapy within the observation period (15.7%). Other reasons included change of treating physician (7.8%), patient withdrawal (2.8%), loss to follow-up (3.1%), patient death (0.2%), unknown reasons (1.3%) or other reasons (1.1%).

For those who discontinued anti-VEGF therapy, the primary reasons, as determined by the treating physician, were stable disease within the observation period (31.5%) or treatment failure (23.2%). Other reasons included fibrosis/atrophic/scarring (8.9%), patient’s decision (4.0%), ocular adverse event (3.2%), cost/reimbursement issue (2.0%), death (0.3%) or other reasons (12.6%).
Analysis by country

Of the eight countries, five enrolled more than 400 patients and were evaluable for separate analysis (France, Germany, UK, Italy and the Netherlands).

Baseline characteristics were generally similar between countries; however, there were differences in baseline visual acuity (table 2 and see online supplementary figure S2).

Mean change in visual acuity score from baseline to years 1 and 2 differed between countries (table 3). In all countries, there was an initial improvement in visual acuity followed by a decrease in the visual acuity gains (figure 2). This decrease was least pronounced in the UK and the Netherlands, where there remained an improvement in visual acuity between baseline and year 2. In contrast, there was a decline in mean visual acuity from baseline to year 2 in patients in Germany, France and Italy. Improvements in visual acuity between baseline and year 2 were also observed in Canada, Ireland and Venezuela, although caution should be used when interpreting these results due to low patient numbers.

In all countries, the mean number of visits was lower in the second year than the first year. However, the number of visits differed between countries. Patients in the UK had the highest number of visits (18.4 in the full 2 years). For other countries, the number of visits ranged from 8.3 (Venezuela) to 13.8 (Canada; table 3).

The number of anti-VEGF injections received differed between countries. Of the countries enrolling more than 400 patients, patients in the UK and the Netherlands had the highest number of injections.

DISCUSSION

In this retrospective study of real-life anti-VEGF therapy for wAMD between 2009 and 2011, visual acuity improved from baseline initially. Thereafter, a decline in visual acuity was observed, and the initial gain was not maintained. Patients visited their ophthalmologists more frequently during the first year and received more injections in the first year than in the second year. The use of a loading scheme appeared to be important for maximising the initial gain in visual acuity but did not seem to influence the rate of visual acuity decline during the full 2 years.

Our results are in line with other observational studies of ranibizumab use in clinical practice. In the LUMIERE study, which observed patients with wAMD who had received ranibizumab treatment for at least 12 months in France between September 2006 and October 2009, the mean change in visual acuity from baseline to month 12 was 3.2 letters and patients received an average of 5.1 injections. A smaller, retrospective chart review of patients treated in a French tertiary care centre showed that ranibizumab was associated with stabilisation of visual acuity, not with visual improvement.

In the German WA VE study, a total of 3470 patients received ranibizumab. German guidelines recommend three monthly intravitreal injections, followed by a maintenance phase, where patients receive reinjections if signs of activity are noted. Patients received a mean of 4.34 ranibizumab injections in 12 months, and the mean change in visual acuity from baseline to month 12 was +0.02 logarithm of minimal angle of resolution.

In a retrospective case review of 50 consecutive English patients with wAMD receiving treatment with ranibizumab on a PRN basis, patients were seen on average 11 times during the mean follow-up period of 13.6 months. Visual acuity at the end of follow-up improved by 4.6 letters from baseline, achieved with a mean of 4.7 injections per 12-month period.

The strengths of our study include the large sample size and the real-life assessment of anti-VEGF use, monitoring and visual

<table>
<thead>
<tr>
<th>Country</th>
<th>N</th>
<th>Mean overall visits in full 2 years</th>
<th>Mean performed VA tests in full 2 years</th>
<th>Mean performed OCTs in full 2 years</th>
<th>Mean injections in full 2 years</th>
<th>Change in VA score at day 90*</th>
<th>Change in VA score at year 1*</th>
<th>Change in VA score at year 2*</th>
<th>Mean VA score at year 2*</th>
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</thead>
<tbody>
<tr>
<td>UK</td>
<td>410</td>
<td>18.4</td>
<td>17.8</td>
<td>16.6</td>
<td>9.0</td>
<td>5.7</td>
<td>6.0</td>
<td>4.1</td>
<td>59.0</td>
</tr>
<tr>
<td>The Netherlands</td>
<td>350</td>
<td>12.7</td>
<td>7.0</td>
<td>5.9</td>
<td>8.7</td>
<td>4.0</td>
<td>1.6</td>
<td>52.7</td>
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<tr>
<td>France</td>
<td>398</td>
<td>13.4</td>
<td>9.2</td>
<td>9.1</td>
<td>6.3</td>
<td>4.1</td>
<td>0.8</td>
<td>41.4</td>
<td>54.4</td>
</tr>
<tr>
<td>Germany</td>
<td>420</td>
<td>10.8</td>
<td>7.7</td>
<td>3.4</td>
<td>5.6</td>
<td>3.3</td>
<td>1.1</td>
<td>−0.8</td>
<td>51.9</td>
</tr>
<tr>
<td>Italy</td>
<td>365</td>
<td>12.7</td>
<td>6.5</td>
<td>4.9</td>
<td>5.2</td>
<td>1.4</td>
<td>0</td>
<td>−2.9</td>
<td>47.8</td>
</tr>
</tbody>
</table>

* Last observation carried forward analysis.
NA, not available; OCT, optical coherence tomography; VA, visual acuity.
outcomes in consecutively enrolled patients across multiple centres and countries. The retrospective design prevented the investigator bias that may occur in prospective open-label studies. Our study is limited by the observational and uncontrolled nature of the design. In particular, our data reflect the documentation in the centres and therefore different patients contributed to the observed visual acuity data at each visit. The LOCF analysis that we, therefore, used to assess visual acuity changes over time led to a small downward shift of the visual acuity curves versus the observed values at each visit (see online supplementary Figure S3). Multiple centres per country were involved and included university hospitals and office-based physicians; however, it is uncertain whether the clinical centres included in AURA are fully representative of patient management in the entire country. For example, different retreatment criteria may have been used in different centres and there were some methodological limitations, such as different clinical centres used different OCT machines and the visual acuity score was translated into one common scale for analysis.

The number of visits, injections and visual acuity outcomes differed substantially between countries. Visual acuity outcomes, measured as change from baseline, may have been partly related to differences in baseline visual acuity and partly due to monitoring and treatment practices. For example, patients in the Netherlands had a lower baseline score (and thus more to gain), while patients in Italy had a higher baseline score and the low gain in visual acuity during the loading phase may have been due to a ceiling effect. However, maintenance of visual acuity gains was very different in countries with similar baseline visual acuity, for example, in the UK versus France. More visits and injections appeared to be correlated with more successful maintenance of visual acuity gains. The SEVEN-UP study, a non-interventional trial which evaluated outcomes 7–8 years after initiation of ranibizumab and included 65 patients originally treated with ranibizumab in the ANCHOR, MARINA and HORIZON studies had similar findings.20 A mean of 6.8 anti-VEGF injections had been received since exit from the HORIZON study (mean of 3.4 years) and while 43% of study eyes had stable or improved letter score versus ANCHOR/MARINA baseline measurements, 34% had declined by ≥15 letters. However, mean gain in letter score since exit from HORIZON was significantly better in patients who had received

Figure 2  Mean change in visual acuity score from baseline over time for all patients by country: Germany, France, UK, Italy and the Netherlands (A) and Canada, Ireland and Venezuela (B). Data based on effectiveness analysis set using a last observation carried forward (LOCF) approach.
11 anti-VEGF injections, suggesting vision may be partly related to injection frequency.20

The differences in disease management between countries are likely due to varying constraints and incentives associated with the healthcare systems within these countries, including reimbursement, selection of patients for treatment or the number of permitted injections. Further investigation of factors associated with improved visual outcome is required.

A substantial proportion of patients did not have complete follow-up for the full 2 years of the study, although this number varied considerably between countries. This highlights the need to ensure that patients continue to be monitored and receive appropriate treatment. The main reasons for discontinuing treatment in AURA were documented as stable disease or treatment failure, but as this was a retrospective analysis, detailed assessments of reasons for discontinuing treatment are limited.

Overall, we observed a good initial response to therapy, which then declined over time. Our results indicate that to achieve the best outcomes with ranibizumab, more frequent monitoring and injections may be needed. Ongoing research into treatment schedules and alternative therapies may help to reduce the trade-off between treatment burden and visual outcomes.

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Contributors FHG, RT, SB, AB, MGC, RC, CBH, PH, GS and SS are members of the Global Steering/Publication Committee for the AURA study and have been responsible for and/or involved in provision of patients/data acquisition, data analysis, data review and interpretation. SH was responsible for the global study management and for the supervision of the set-up and conduct of the study. TH was the sponsor study physician and has contributed to the study concept, data analysis and interpretation. TB has contributed to the study concept, data analysis and interpretation. All authors provided critical review of the manuscript and approved the final version for publication.

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Competing interests FHG is a consultant to Acurela, Genentech, Novartis, Bayer HealthCare, Alcon, OPTOS, Heidelberg Engineering, Carl Zeiss Meditec, Allergan and Pfizer and has received grants from OPTOS, Heidelberg Engineering, Carl Zeiss Meditec, Alcon, Genentech, Bayer HealthCare and Novartis. RT is a board member of Alcon, Novartis, Allergan, Bausch & Lomb, Pfizer, Alimera, Bayer HealthCare and FCI-Zeiss and a consultant for Allergan, DORC, Alcon, Novartis, Takeda, Bausch & Lomb and FCI-Zeiss. He has received payment for lectures from Alcon, Bausch & Lomb, Novartis, Allergan, Pfizer, Takeda, Bayer HealthCare and Alimera and has received payment for the development of educational presentations from Bausch & Lomb, Novartis, Zeiss, Sony, Alcon and Allergan. He has also received support for meeting expenses (travel and accommodation) from Alcon, Novartis, Allergan, Bausch & Lomb, Pfizer, Bayer HealthCare, DORC, Takeda, Servier and Alimera. SB is a consultant and board member for Bayer HealthCare and has received support for travel to meetings and fees for participation in review activities from Bayer HealthCare. AB is a consultant to Bayer HealthCare and has received travel support and fees for participation in review activities from Bayer HealthCare. He has also received payment for lectures from Bayer HealthCare. MGc is a consultant to Bayer HealthCare and has received support for travel to meetings from Bayer HealthCare. RC’s institution has received an honorarium from Bayer HealthCare. CBH is a consultant to Bayer HealthCare. PH has received grants from Bayer HealthCare, Alimera and Novartis and acts as a consultant to Bayer HealthCare, Novartis and Allergan. He has also received support for travel to meetings, for participation in review activities and provision of writing assistance, medicines, equipment or administrative support from Bayer HealthCare and Novartis, received payment for lectures from Bayer HealthCare, Alimera and Novartis and has received support for conference attendance. GS acts as a consultant to Novartis, Bayer HealthCare, Allergan, Genentech, Roche, Heidelberg Engineering, Alcon, QLT and OD-DS. He also received support for travel to meetings from Bayer HealthCare, payment for lectures from Zeiss and is a patent holder in conjunction with Ocular Instruments, Inc. SH, TH and TB are employees of Bayer HealthCare. SS has received grants from Bayer HealthCare, Alimera and Novartis and acts as a consultant to Bayer HealthCare, Novartis and Allergan. She has also received support for travel to meetings, for participation in review activities and provision of writing assistance, medicines, equipment or administrative support from Bayer HealthCare and Novartis, received payment for lectures from Bayer HealthCare, Alimera and Novartis and has received support for conference attendance.

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