Bilateral involvement by age related maculopathy lesions in a population

Jie Jin Wang, Paul Mitchell, Wayne Smith, Robert G Cumming

Abstract

Aims—To describe the influences of age and sex on the frequency of bilateral age related macular degeneration (AMD) and age related maculopathy (ARM) lesions.

Methods—The Blue Mountains Eye Study examined 3654 older Australians, 82% of permanent residents living in an area west of Sydney. Stereo macular photographs were graded for AMD (neovascular maculopathy and geographic atrophy) and early ARM lesions (soft drusen, reticular drusen, hyperpigmentation, and hypopigmentation).

Results—Among 230 gradable cases of AMD or early ARM, 183 (80%) were bilateral. For AMD, 39/69 cases (57%) were bilateral, while for early ARM, 123/161 cases (77%) had signs in both eyes. The individual lesions, reticular drusen (91%) and indistinct soft drusen (79%) were most frequently present in both eyes. Geographic atrophy was bilateral in 56%, neovascular AMD in 40%, and distinct soft drusen in 47%, while hyperpigmentation was bilateral in 38% and hypopigmentation in only 28% of cases. A consistent age related increase in bilateral distribution was observed for most lesions. After adjusting for effects of age, current smoking, and AMD family history AMD and ARM component lesions, except for soft drusen, were more frequently bilateral in women. This sex difference was significant only for neovascular AMD, odds ratio 7.7 (95% confidence intervals 1.3–46.7). An AMD family history was more frequently reported in cases with bilateral involvement.

Conclusions—This study has documented differences in the age related bilaterality of individual ARM components with higher bilateral rates for reticular or indistinct soft drusen compared with other lesions. The increased bilaterality of most ARM lesions among women is likely to contribute to the increased age adjusted risk of AMD blindness found in women.

Materials and methods

STUDY POPULATION

The Blue Mountains Eye Study is a population based survey of vision and common eye diseases in an urban elderly population (aged 49 years and older) comprising two postcode areas in the Blue Mountains region, west of Sydney, Australia. Details of the survey method and procedures have been previously described. Of 4433 eligible residents, 3654 (82.4%) were examined during 1992–4. Family
history of AMD was assessed using an interviewer administered questionnaire, which included the following questions: “We want to know if eye disease runs in families. Have either of your parents, or any brothers or sisters or children had macular degeneration diagnosed?” An AMD family history was defined from a positive answer to this question; subjects with missing family history data were excluded from analyses. Smoking status was defined as “current” if the participant currently smoked or had stopped smoking for less than 1 year before the examination. Subjects who had ever smoked but had stopped for more than 1 year were defined as “past” smokers.

All participants underwent a detailed eye examination. This included Zeiss FF3 30° stereo photographs (Carl Zeiss, Oberkochen, Germany) of Diabetic Retinopathy Study fields one (disc) and two (macula), with non-stereo photographs of fields three (temporal to macula), four and five (upper and lower vascular arcades) and a field nasal to the optic disc. Two graders performed a masked grading of the stereo fundus photographs to assess presence of AMD or ARM lesions. The Wisconsin Age Related Maculopathy Grading System was closely followed with minor modifications. A transparent grid, supplied by Dr Ronald Klein, University of Wisconsin, Madison, including three concentric circles, was placed over one of the stereo photos of field two for each eye. This divided the macular area into three zones: central, inner, and outer circles, corresponding to a radius of 500, 1500, and 3000 µm from the foveolar centre. All photographs were graded initially by either one of the two graders and all advanced cases were assessed by PM, who also adjudicated any queries or discrepancies. Intergrader and intra-grader reliability, assessed on a random subsample of gradable eyes, showed good agreement for all lesions. In the assessment of lesion bilaterality, all cases with ARM lesions had a detailed regrading by the two graders.

GRADING DEFINITIONS
Age related macular degeneration (AMD) included the two end stage lesions, neovascular maculopathy and geographic atrophy involving the foveal centre. Neovascular lesions included serous or haemorrhagic detachment of the RPE or sensory retina, or subtretinal fibrous tissue. Geographic atrophy was defined as a discrete area of retinal depigmentation at least 175 µm in diameter, characterised by a sharp border and the presence of visible choroidal vessels. Eyes with minimal subtretinal fibrosis and widespread surrounding atrophy were classified as having both neovascular AMD and geographic atrophy for the assessment of bilaterality. Cases with neovascular AMD in one eye and geographic atrophy in the fellow eye were graded as unilateral for both neovascular AMD and geographic atrophy, but bilateral for “any AMD”.

Early ARM was defined as the absence of AMD and either (1) indistinct soft or reticular drusen or (2) both distinct soft drusen and retinal pigmentary abnormalities, within the area of the grid. Hyperpigmentation was defined as definite clumps of dark brown or dark green pigment beneath the retina. Hypopigmentation was defined as discrete areas of retinal depigmentation without visible choroidal vessels, often with adjacent pigment clumping. We included cases only in which large drusen (maximum size larger than 125 µm diameter) were present in at least one eye. Subjects with AMD in either eye were excluded from the assessment of bilateral involvement by early ARM lesions.

We examined the following lesion groups for bilaterality:
1. Any ARM—either AMD or early ARM present in both eyes, including subjects with AMD in one eye and early ARM in the fellow eye
2. Any AMD—
   a. neovascular AMD
   b. geographic atrophy
3. Any early ARM—early ARM in one or both eyes without AMD in either eye
   a. reticular drusen
   b. indistinct soft drusen
   c. distinct soft drusen
   d. retinal pigmentary abnormalities (hyper- or hypopigmentation)
   e. hyperpigmentation
   f. hypopigmentation
4. Symmetrical soft drusen characteristics
   a. type: at least one soft drusen type in common between the eyes
   b. size: maximum drusen size no more than one grading step different between the two eyes
   c. location: majority of drusen located either within or outside the grid for the two eyes
   d. area: soft drusen area within the central and inner circles no more than one grading step different between the two eyes.

Cases with data available for only one eye were excluded from this study (three AMD cases and 10 early ARM cases, because of coexisting dense cataract, corneal, or retinal lesions).

DATA HANDLING AND STATISTICAL METHODS
Statistical Analysis System (version 6.11) (SAS Institute Inc, Cary, NC, USA) was used for statistical analyses including $\chi^2$ statistic, Mantel–Haenszel $\chi^2$ trend statistic and logistic regression analyses. In the logistic regression analyses, bilateral (compared with unilateral) involvement by individual ARM lesions or lesion groups, were the dependent variables and were assessed separately in different models. Age was used continuously while sex, current smoking, and AMD family history were included in this study. No statistical adjustment has been made for multiple comparisons.
Bilateral involvement by age related maculopathy lesions in a population

Table 1  Bilaterality of selected age related macular degeneration (AMD) and age related maculopathy (ARM) lesions

<table>
<thead>
<tr>
<th>Lesions</th>
<th>Total no of cases</th>
<th>No (%) with bilateral involvement</th>
</tr>
</thead>
<tbody>
<tr>
<td>Any ARM (AMD + early ARM)</td>
<td>230</td>
<td>183 (80)</td>
</tr>
<tr>
<td>Any ARM</td>
<td>69</td>
<td>39 (57)</td>
</tr>
<tr>
<td>neovascular AMD</td>
<td>48</td>
<td>19 (40)</td>
</tr>
<tr>
<td>geographic atrophy</td>
<td>39</td>
<td>22 (56)</td>
</tr>
<tr>
<td>Any early ARM</td>
<td>161</td>
<td>123 (76)</td>
</tr>
<tr>
<td>reticular drusen</td>
<td>35</td>
<td>32 (91)</td>
</tr>
<tr>
<td>indistinct soft drusen</td>
<td>118</td>
<td>93 (79)</td>
</tr>
<tr>
<td>distinct soft drusen</td>
<td>184</td>
<td>87 (47)</td>
</tr>
<tr>
<td>pigmentary abnormality</td>
<td>374</td>
<td>140 (37)</td>
</tr>
<tr>
<td>hyperpigmentation</td>
<td>368</td>
<td>138 (38)</td>
</tr>
<tr>
<td>hypopigmentation</td>
<td>262</td>
<td>73 (28)</td>
</tr>
</tbody>
</table>

Results

BILATERAL INVOLVEMENT OF ANY ARM, AMD, AND EARLY ARM

After excluding subjects who could not be graded for bilaterality, 230 subjects had any ARM (AMD or early ARM), of whom 183 (80%) were bilateral. Taking the two subgroups separately, 69 cases had AMD, including 39 (57%) who had bilateral AMD lesions, 161 cases had early ARM, including 123 (76%) who had bilateral lesions. In 20 of the 30 unilateral AMD cases, early ARM was present in the fellow eye. Among people with AMD, 19 of the 48 (40%) neovascular AMD cases were bilateral and 22 of the 39 (56%) geographic atrophy cases were bilateral (Table 1). In seven of the 29 cases (24%) with unilateral neovascular AMD, geographic atrophy was present in the fellow eye, while eight of the 17 unilateral geographic atrophy cases (53%) had neovascular AMD in the fellow eye. Seven cases had coexisting neovascular AMD and GA in the same eye.

For individual early ARM lesions graded, reticular drusen (91%) and indistinct soft drusen (79%) had the highest frequency of bilateral involvement. Distinct soft drusen were bilateral in 47% and retinal pigmentary abnormalities in 37% of cases (Table 1). Signs of hyperpigmentation and hypopigmentation were bilateral in 38% and 28% of cases, respectively. Graded soft drusen characteristics (type, size, location, and area covered) were also symmetrical between the two eyes in around 60% of cases (results not shown).

Figure 1 Influence of age on bilateral involvement by early and late lesions among people with signs of age related maculopathy.

Table 2  Effect of age on bilateral involvement by lesions of age related macular degeneration (AMD) and early age related maculopathy (ARM)

<table>
<thead>
<tr>
<th>Outcomes</th>
<th>Total no of cases</th>
<th>Percentage with bilateral involvement</th>
<th>Mantel–Haenszel χ² for trend</th>
<th>p Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Any ARM</td>
<td>230</td>
<td>43/60 60–69/70–79/80+</td>
<td></td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Any AMD</td>
<td>69</td>
<td>0/17 60–69/70–79/80+</td>
<td></td>
<td>0.003</td>
</tr>
<tr>
<td>neovascular AMD</td>
<td>48</td>
<td>0/17 60–69/70–79/80+</td>
<td></td>
<td>0.03</td>
</tr>
<tr>
<td>geographic atrophy</td>
<td>39</td>
<td>0/17 60–69/70–79/80+</td>
<td></td>
<td>0.78</td>
</tr>
<tr>
<td>Any early ARM</td>
<td>161</td>
<td>43/60 60–69/70–79/80+</td>
<td></td>
<td>0.001</td>
</tr>
<tr>
<td>reticular drusen</td>
<td>35</td>
<td>0/17 60–69/70–79/80+</td>
<td></td>
<td>0.002</td>
</tr>
<tr>
<td>indistinct soft drusen</td>
<td>118</td>
<td>50/60 60–69/70–79/80+</td>
<td></td>
<td>0.02</td>
</tr>
<tr>
<td>distinct soft drusen</td>
<td>184</td>
<td>13/41 60–69/70–79/80+</td>
<td></td>
<td>0.002</td>
</tr>
<tr>
<td>pigmentary abnormality</td>
<td>374</td>
<td>16/44 60–69/70–79/80+</td>
<td></td>
<td>0.01</td>
</tr>
<tr>
<td>hyperpigmentation</td>
<td>368</td>
<td>16/31 60–69/70–79/80+</td>
<td></td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>hypopigmentation</td>
<td>262</td>
<td>14/29 60–69/70–79/80+</td>
<td></td>
<td>0.14</td>
</tr>
</tbody>
</table>

EFFECT OF AGE AND SEX ON THE BILATERAL INVOLVEMENT OF LESIONS

As expected, a consistent age related increase in bilateral involvement for most lesions was found. This was highly significant using the Mantel–Haenszel χ² test for trend and logistic regression analyses, for almost all lesions (Table 2 and Fig 1). Exceptions were the bilateral prevalence of the following three lesions, two of which were relatively infrequent. Bilateral geographic atrophy was not observed until age 70–79 and declined for people aged 80 or older. Reticular drusen were nearly always found to be present in both eyes. Hypopigmentation was also uncommonly present in both eyes and showed no significant increase in bilateral involvement with increasing age. For all four soft drusen characteristics graded, a similar and statistically significant age related increase in bilateral involvement was found (Mantel–Haenszel χ² for trend, p<0.001, results not shown).

All age related maculopathy lesions were more frequently bilateral in women than men, apart from indistinct and distinct soft drusen. This consistent difference remained after adjusting for age, current smoking, and family history of AMD (Table 3). A significantly higher rate of bilateral involvement in women than men was found for neovascular AMD, odds ratio (OR) 7.7, 95% confidence intervals (CI) 1.3–46.7. For all other lesions, non-significant increased odds ratios were found for bilateral involvement in women (OR 2.5 for any AMD, OR 2.4 for geographic atrophy, and OR 1.6 for early ARM). For individual or grouped retinal pigmentary abnormalities, non-significant odds ratios of 1.3:1.4 were found. All four soft drusen characteristics were also more frequently bilateral in females, with non-significant increased age adjusted odds ratios ranging from 1.2 to 1.4 (results not shown).

As current smoking and family history of AMD have been shown to have strong associations with AMD, we also tried to assess the relation between AMD family history and bilateral involvement by ARM lesions. For the following lesions: (1) any ARM, (2) early ARM, (3) indistinct soft drusen and (4) reticular drusen, all cases reporting an AMD family history had the lesions present bilaterally.
Table 3  Effect of sex on the frequency of bilateral involvement by age related macular degeneration (AMD) and age related maculopathy (ARM) lesions

| Outcomes                        | Female Total cases | Female No of bilateral (%) | Male Total cases | Male No of bilateral (%) | Multivariate adjusted odds ratio (95% CI)*  
|--------------------------------|-------------------|-----------------------------|------------------|--------------------------|------------------------------------------  
| Any ARM                         | 139               | 116 (83)                    | 91               | 67 (74)                  | 1.6 (0.8–3.2)†                           
| Any AMD                         | 47                | 30 (64)                     | 22               | 9 (41)                   | 2.3 (0.8–8.0)                            
| neovascular AMD                 | 33                | 17 (52)                     | 15               | 2 (13)                   | 7.7 (1.3–46.7)                           
| geographic atrophy              | 25                | 16 (64)                     | 14               | 6 (43)                   | 2.4 (0.6–10.0)                           
| Any early ARM                   | 92                | 74 (80)                     | 69               | 49 (71)                  | 1.6 (0.7–3.5)†                          
| reticular drusen                | 25                | 24 (96)                     | 10               | 8 (80)                   | 8.5 (0.5–133.1)‡                         
| indistinct soft drusen          | 63                | 50 (79)                     | 55               | 43 (78)                  | 1.0 (0.4–2.6)†                           
| distinct soft drusen pigmentary | 104               | 51 (49)                     | 80               | 36 (45)                  | 1.1 (0.6–2.0)                            
| abnormality                     | 190               | 78 (41)                     | 184              | 62 (34)                  | 1.3 (0.8–1.9)                            
| hyperpigmentation               | 186               | 77 (41)                     | 182              | 61 (34)                  | 1.3 (0.8–2.0)                           
| hypopigmentation                | 141               | 44 (31)                     | 121              | 29 (24)                  | 1.4 (0.8–2.5)                           

*Adjusted for age, family history of AMD and current smoking.
†Adjusted for age and current smoking only as no unilateral cases had a family history of AMD.
‡Adjusted for age only as no unilateral cases had a family history of AMD or currently smoked.

Discussion

Although the prognosis of bilateral macular drusen has been reported from case series,13–14 few studies have examined the bilaterality of individual ARM lesions.12 27–28 with no data from a population reported to date. Population based data on the bilateral frequency of ARM and individual ARM lesions is valuable as it is not affected by the selection biases inherent in clinic based samples. Accurate estimates of the frequency of bilateral involvement by ARM lesions would assist clinicians to predict the cost of AMD and to plan eye care and visual rehabilitative services for this disease.

In our study, any ARM lesion was found in both eyes of 80% of gradable subjects, with both eyes affected in 57% of the AMD cases. This proportion of bilateral AMD is only slightly higher than the 42% rate (34/77) calculated from the Beaver Dam Eye Study report,29 and may be due to the older age profile of our study population. Our rates for any ARM are also comparable to the Framingham Eye Study report,30 despite the use of different diagnostic criteria and methods of ascertainment. In Framingham, the bilateral proportion of “senile macular degeneration” cases was 64%, 71%, and 79% for age groups 52–64, 65–74, and 75–85, respectively.30

As the major determinant of ARM is increasing age,21 29–31 it could be expected that increasing bilateral involvement by individual ARM lesions would also be strongly age dependent. While this was found for most ARM lesions, we found no significant age related increase in bilateral involvement by hypopigmentation, a relatively common lesion. The contrasting bilaterality rates found for individual ARM lesions may imply that indistinct or reticular soft drusen are highly specific markers for ARM, while hypopigmentation may be a less specific manifestation of this disease. However, we can not exclude the possibility that measurement error or grading difficulty could have affected the grading of hypopigmentation or that some findings may be due to chance as a result of the multiple analyses performed.

In the Beaver Dam Eye Study population, women 75 years of age or older had a significantly higher frequency of neovascular AMD27 as well as a higher 5 year incidence rate of early ARM (OR 2.2, compared with men).32 In our study population, women had higher age specific prevalence rates for AMD lesions and soft drusen than men.21 Furthermore, 15 of the 16 people bilaterally blind from AMD were women, which is reflected in the very high age adjusted odds ratio (7.7) found for bilateral neovascular AMD in women compared with men in our study. For AMD and all individual ARM lesions apart from soft drusen, bilateral involvement was more frequent in women than men, after adjusting for age, current smoking, and AMD family history. An increased life expectancy for women could partly explain the different rate of bilateral AMD between women and men, but not the observed differences in risk for bilateral early ARM or pigmentary abnormalities. Hormonal factors in women could explain these consistent differences, as the Rotterdam study has reported an increased risk of AMD for early menopause in women.33 A decreased risk of neovascular AMD has also been reported for women taking postmenopausal oestrogen therapy.34

AMD family history was associated with bilateral involvement of most lesions and was absent in all unilateral cases of any ARM, early ARM, indistinct soft drusen, and reticular drusen. Strong familial aggregation of AMD was recent reported from the Rotterdam Eye Study.20 A recent case-control study also reported significant familial aggregation at a lower strength.35 Our finding is consistent with both these findings and further emphasises the role of genetic influences in ARM.

In summary, the Blue Mountains Eye Study has provided age specific bilateral prevalence rates for AMD and ARM in older Australians. After adjusting for the effects of age, current smoking, and AMD family history, the Blue Mountains Eye Study suggests an increased risk of bilateral AMD and early ARM in women, statistically significant for bilateral neovascular AMD. An AMD family history may also predict bilateral involvement of most individual ARM lesions.

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References

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