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Screening older people for impaired vision in primary care: cluster randomised trial

Liam Smeeth, Astrid E Fletcher, Smita Hanciles, Jennifer Evans, Richard Wormald

Abstract

Objective To determine the effectiveness of screening for visual impairment in people aged 75 or over as part of a multidimensional screening programme.

Design Cluster randomised trial.

Setting General practices in the United Kingdom participating in the MRC trial of assessment and management of older people in the community.

Participants 4340 people aged 75 years or over randomly sampled from 20 general practices, excluding people resident in hospitals or nursing homes.

Intervention Visual acuity testing and referral to eye services for people with visual impairment. Universal screening (assessment and visual acuity testing) was compared with targeted screening, in which only participants with a range of health related problems were offered an assessment that included acuity screening.

Main outcome measures Proportion of people with visual acuity less than 6/18 in either eye; mean composite score of 25 item version of the National Eye Institute visual function questionnaire.

Results Three to five years after screening, the relative risk of having visual acuity < 6/18 in either eye, comparing universal with targeted screening, was 1.07 (95% confidence interval 0.84 to 1.36; P = 0.58). The mean composite score of the visual function questionnaire was 85.6 in the targeted screening group and 86.0 in the universal group (difference 0.4, 95% confidence interval −1.7 to 2.5, P = 0.69).

Conclusions Including a vision screening component by a practice nurse in a pragmatic trial of multidimensional screening for older people did not lead to improved visual outcomes.

Introduction

In population based surveys undertaken in the United Kingdom, visual acuity of less than 6/12 (below the United Kingdom driving requirement) has been found in around 20% of people aged 75 or older.1–3 A recent survey estimated that more than 70% of visual impairment in people aged 65 or older was potentially reme-

Baseline screening occurred between 1995 and 1999. Practices were randomised to two forms of screening: targeted or universal. Randomisation was performed centrally at the London School of Hygiene and Tropical Medicine, using a computer generated randomisation list stratified by thirds of Jarman score and of standardised mortality ratio. Because of the nature of the intervention, participants or researchers could not be blinded to the group assignment.
Interventions

**Universal screening group**

All participants in the trial were invited to complete a brief assessment followed by a detailed health assessment by a trained nurse that included measurement of visual acuity. Acuity was measured using a Glasgow acuity chart, which measures vision on the logMAR scale. For ease of interpretation, we give the Snellen equivalent acuity. Binocular vision was measured first, followed by vision in each eye. Measurements were repeated in people with visual acuity less than 6/18 in either eye, using a pinhole occluder to provide an estimate of acuity corrected for refractive error.

For people with a pinhole vision of less than 6/18 in either eye, referral to an ophthalmologist was recommended unless they had been seen by an ophthalmologist in the previous year, or were registered blind or partially sighted. People with presenting vision of less than 6/18 in either eye that improved with pinhole to better than 6/18 were advised to see an optician. People with visual acuity less than 6/18 who could not complete a pinhole assessment were recommended for referral to an ophthalmologist.

**Targeted screening group**

At the time the trial was started, the contractual obligation on general practitioners to offer all patients aged over 75 years an annual health assessment meant it was not possible to have a control group that was offered no screening. In practices randomised to the targeted screening strategy, participants were invited to complete a brief screening assessment. This assessment covered all areas specified in the general practitioner contract, including a question about difficulty seeing, but did not include measurement of visual acuity. Only people found to have a specified range and level of problems during the brief assessment were invited to have a detailed assessment. Although reporting a lot of difficulty seeing was one possible factor contributing to eligibility for a more detailed assessment, it was not on its own sufficient to lead to an offer of a detailed assessment. The trial was designed such that about 20% of people in the targeted group would be eligible for a more detailed assessment.

Sampling

We randomly sampled 10 practices from each arm of the trial and 220 participants from each sampled practice and asked them to participate in the collection of visual outcomes (fig 1).

At baseline the prevalence of visual acuity < 6/18 in one or both eyes was 30% with an intraclass correlation coefficient of 0.022. We estimated that outcome data for 2000 participants would provide over 80% power significant at the 5% level to detect a 30% difference in the prevalence of visual impairment. To allow for deaths and other losses to follow up we randomly sampled 220 eligible participants from each practice. Two practices in the universal screening group had fewer than 220 eligible people: 170 in one practice and 210 in the other. The final sample therefore included 2140 people in the universal screening group and 2200 people in the targeted screening group.

Outcome assessment

Three to five years after the baseline screening, sampled participants in the 20 practices included were invited to have an assessment of their vision. Research nurses assessed visual acuity and visual function and asked participants about contact with any eye services. In addition they searched medical records for information relating to vision. The two primary outcomes were the proportion of people with presenting visual acuity < 6/18 in one or both eyes, and the composite score of the 25 item version of the National Eye Institute visual function questionnaire. The questionnaire has been shown to be a reliable and valid measure of visual function; the highest available score is 100 representing no disability related to vision.

Analysis

All analyses used the "survey" commands in Stata 7.0 and took account of the cluster design. The intention to treat principle was applied as far as possible. Participants with outcome data available were analysed in the group to which they were randomised, regardless of whether they actually had a screening assessment at baseline.

Results

**Response rates and baseline characteristics**

In the targeted screening group, 76.5% (1684/2200) of participants completed a brief screening assessment; 150 (6.8%) had sufficient number or severity of problems to be eligible for a more detailed assessment including visual acuity; and 120 (5.5%) completed the detailed assessment. In the universal screening group, 73.1% (1565/2140) completed a detailed assessment including visual acuity (fig 2). Table 1 shows the characteristics of participants at baseline.

**Vision screening and eligibility for referral**

In the universal screening group, 451/1565 people (28.8%) had a visual acuity < 6/18 in either eye. Of these, 79 (18%) were eligible for referral to an optician and 220 (49%) were eligible for referral to an ophthalmologist. Among the 120 people in the targeted
screening group who had a visual acuity test, the proportion of people with reduced visual acuity was higher than in the universal screening group (43% v 29%), as were the proportions eligible for referral to the eye services (7% v 5% for referral to optician, 24% v 14% for referral to ophthalmologist) (table 2), reflecting the fact that eligibility for vision screening was based on having a specified level of problems detected at the brief assessment.

Outcome assessment

A total of 1807 outcome assessments were completed, 90.4% of the target of 2000 (fig 2). Excluding the 1465 people who died before assessment of outcome, the response rate was 62.8% (1807/2875). The response rate was 5.7% lower in the universal screening group (829/1432 v 978/1443), largely because two practices in the universal screening arm had low response rates because of problems with nurses’ workload. The median time between baseline screening and outcome assessment was 3.9 years (interquartile range 3.3-4.5 years) in the targeted screening group and 3.9 (3.4-4.3) years in the universal group.

Primary outcomes

Comparing universal with targeted screening, we found that the relative risk of visual acuity less than 6/18 in either eye was 1.07 (95% confidence interval 0.84 to 1.36; P = 0.58) (table 3). The mean composite score of the National Eye Institute visual function questionnaire was 85.6 in the targeted screening group and 86.0 in the universal group, a difference of 0.4 (−1.7 to 2.5; P = 0.69). The standard deviation for the composite score was 17.8 overall and was similar in both groups.

Adjustment for age, sex, level of self reported visual difficulty at baseline, and time until follow up had no impact on the results. Stratification by age, sex, level of self reported visual difficulty at baseline, social isolation, or time until follow up produced no evidence of subgroup effects. For the subgroup with time to fol-

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Table 1 Baseline characteristics of participants. Values are percentages (numbers)

<table>
<thead>
<tr>
<th></th>
<th>Targeted screening group (n=1684)</th>
<th>Universal screening group (n=1662)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Men</strong></td>
<td>36.8 (582)</td>
<td>39.8 (661)</td>
</tr>
<tr>
<td><strong>Median age (years)</strong></td>
<td>79.9</td>
<td>80.2</td>
</tr>
<tr>
<td>Reporting a lot of difficulty seeing newsprint</td>
<td>9.7 (162/1672)</td>
<td>7.7 (127/1668)</td>
</tr>
<tr>
<td>Socially isolated*</td>
<td>15.1 (254/1657)</td>
<td>16.8 (278/1666)</td>
</tr>
<tr>
<td>Reporting often or always having difficulty making ends meet</td>
<td>2.1 (35/1672)</td>
<td>2.3 (39/1669)</td>
</tr>
<tr>
<td>Reporting one or more falls at home in the previous six months</td>
<td>17.7 (290/1665)</td>
<td>20.4 (334/1663)</td>
</tr>
<tr>
<td>Taking 5 or more drugs regularly</td>
<td>17.6 (278/1602)</td>
<td>18.5 (286/1547)</td>
</tr>
<tr>
<td>*Defined as reporting seeing friends, neighbours, or relatives less than twice a week.</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

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Table 2 Screening findings and eligibility for referral in people who had visual acuity screening. Values are percentages (numbers)

<table>
<thead>
<tr>
<th>Findings</th>
<th>Targeted screening (n=120)</th>
<th>Universal screening (n=1565)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Detailed assessment</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Visual acuity &lt;6/18 in either eye</td>
<td>43 (53)</td>
<td>29 (451)</td>
</tr>
<tr>
<td>Pinhole assessment carried out</td>
<td>51 (27/93)</td>
<td>36 (223/451)</td>
</tr>
<tr>
<td>Visual acuity &lt;6/18 in either eye that corrected to &gt;6/18 with pinhole correction</td>
<td>6 (8)</td>
<td>50 (79)</td>
</tr>
<tr>
<td>Seen by ophthalmologist in previous 12 months</td>
<td>7 (8)</td>
<td>6 (130)</td>
</tr>
<tr>
<td>Registered:</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Blind</td>
<td>3 (4)</td>
<td>1 (18)</td>
</tr>
<tr>
<td>Partially sighted</td>
<td>5 (6)</td>
<td>2 (35)</td>
</tr>
<tr>
<td>Data on registration status missing</td>
<td>3 (3)</td>
<td>1 (24)</td>
</tr>
<tr>
<td>Eligible for referral to optician</td>
<td>7 (8)</td>
<td>6 (179)</td>
</tr>
<tr>
<td>Eligible for referral to ophthalmologist</td>
<td>24 (29)</td>
<td>14 (229)</td>
</tr>
<tr>
<td><strong>Other visual findings</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Visual acuity &lt;6/18 binocular</td>
<td>22 (26)*</td>
<td>12 (179)*</td>
</tr>
<tr>
<td>Visual acuity &lt;6/12 in either eye</td>
<td>60 (72)</td>
<td>47 (736)</td>
</tr>
<tr>
<td>Visual acuity &lt;6/12 binocular</td>
<td>46 (54)*</td>
<td>34 (508)*</td>
</tr>
<tr>
<td>Owned glasses</td>
<td>88 (105)</td>
<td>88 (1382)</td>
</tr>
<tr>
<td>*Excluding two people with missing data.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>†Excluding 84 people with missing data.</td>
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</table>
Table 3  Visual acuity outcomes. Values are percentages (numbers)

<table>
<thead>
<tr>
<th></th>
<th>Targeted screening group (n=978)</th>
<th>Universal screening group (n=829)</th>
<th>Risk ratio (95% CI)</th>
<th>P value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Visual acuity &lt;6/18 in either eye</td>
<td>34.7 (339/978)</td>
<td>37.0 (307/829)</td>
<td>1.07 (0.84 to 1.36)</td>
<td>0.58</td>
</tr>
<tr>
<td>Visual acuity &lt;6/18 binocular vision</td>
<td>16.0 (160/962)</td>
<td>14.0 (114/817)</td>
<td>0.84 (0.64 to 1.10)</td>
<td>0.20</td>
</tr>
<tr>
<td>Visual acuity &lt;6/12 in either eye</td>
<td>59.7 (584/978)</td>
<td>58.8 (486/829)</td>
<td>0.98 (0.82 to 1.17)</td>
<td>0.83</td>
</tr>
<tr>
<td>Visual acuity &lt;6/12 binocular vision</td>
<td>36.5 (351/962)</td>
<td>31.3 (256/817)</td>
<td>0.80 (0.65 to 1.13)</td>
<td>0.21</td>
</tr>
</tbody>
</table>

Discussion

Even though this trial overcame possible reasons for the lack of effectiveness of screening seen in previous trials (by using measured visual acuity rather than self-reported problems, and by having a clear protocol for intervention), it showed that screening older people for visual impairment as part of a multidimensional screening assessment gave no benefit.

Possible explanations for lack of effect

A "no screening" control arm was not possible, but this feature of the design is unlikely to explain the lack of effect: in the targeted screening group, only 2.2% of people were eligible for referral to the eye services, compared with 18.0% in the universal screening group.

Although a high prevalence of visual impairment was identified at screening, many affected people were already in contact with the eye services. In addition, the period between screening and assessment of outcome was quite long (median 3.9 years). Improvement or deterioration in vision between screening and follow-up could have been missed. The intervention was, however, no more effective in a subgroup with a shorter time to assessment (1.6 to 3 years).

Response to outcome assessments seems low because just over a third of participants had died by the time of the assessment. The proportions who died were similar in the two groups. The follow up rate was 5.7% lower in the universal screening group. Although loss to follow up was not associated with baseline vision, a small effect on the results of the trial cannot be excluded.

Around half the people who had been advised to see an ophthalmologist obtained new lenses. However, uncorrected refractive error was underdetected because a minority of participants completed the pinhole assessment, largely because of difficulties with using a pinhole occluder. Only 52% of people eligible for a referral had evidence of a new referral to an ophthalmologist; in previous trials of multidimensional screening, rates of clinicians' adherence to recommendations for referral have been in the range of 49% to 79%.

Attendance at clinic of patients with evidence of a new referral to an ophthalmologist was 88%, higher than the range observed in previous trials (46% to 70%).

Conclusion

Some people obtained beneficial interventions from vision screening, but the number of people benefiting was small in the context of a population-based screening programme and was not sufficient to affect the prevalence of visual impairment overall. The evidence from randomised controlled trials undertaken to date does not support the inclusion of a vision screening component in multidimensional screening programmes for older people in a community setting.
Barriers to improving vision among older people will need to be overcome before screening is likely to be effective

The recommendation that an assessment of vision be included in the single assessment process proposed in the national service framework for older people is not supported by the evidence. Given the importance of visual impairment among older people, other strategies to improve their vision is needed. The effectiveness of an optimised primary care based screening intervention that overcomes possible factors contributing to the observed lack of benefit in trials to date warrants assessment.

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Contributors: LS, AF, RW, and JE designed the baseline vision screening. SH coordinated the collection of visual outcomes. LS undertook the sampling, analysed the data, and wrote the paper. All authors commented on drafts of the paper and approved the final version. LS and AF are the guarantors.

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Ethical approval: Ethical approval for all aspects of the study was obtained from relevant ethics committees both for the original trial and for the practices involved in the repeat vision assessment.