Systematic Review and Meta-Analysis

Lutein and zeaxanthin intake and the risk of age-related macular degeneration: a systematic review and meta-analysis

Le Ma1, Hong-Liang Dou2, Yi-Qun Wu3, Yang-Mu Huang1, Yu-Bei Huang3, Xian-Rong Xu1, Zhi-Yong Zou1 and Xiao-Ming Lin1*

1Department of Nutrition and Food Hygiene, School of Public Health, Peking University, 38 Xueyuan Road, Beijing 100191, People’s Republic of China
2Peking University Eye Center, Peking University Third Hospital, 49 North Garden Road, Beijing 100191, People’s Republic of China
3Department of Epidemiology and Biostatistics, School of Public Health, Peking University, 38 Xueyuan Road, Beijing 100191, People’s Republic of China

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Abstract

Lutein and zeaxanthin are thought to decrease the incidence of age-related macular degeneration (AMD); however, findings have been inconsistent. We conducted a systematic literature review and meta-analysis to evaluate the relationship between dietary intake of lutein and zeaxanthin and AMD risk. Relevant studies were identified by searching five databases up to April 2010. Reference lists of articles were retrieved, and experts were contacted. Literature search, data extraction and study quality assessment were performed independently by two reviewers and results were pooled quantitatively using meta-analysis methods. The potential sources of heterogeneity and publication bias were also estimated. The search yielded six longitudinal cohort studies. The pooled relative risk (RR) for early AMD, comparing the highest with the lowest category of lutein and zeaxanthin intake, was 0.96 (95% CI 0.78, 1.17). Dietary intake of these carotenoids was significantly related with a reduction in risk of late AMD (RR 0.74; 95% CI 0.57, 0.97); and a statistically significant inverse association was observed between lutein and zeaxanthin intake and neovascular AMD risk (RR 0.68; 95% CI 0.51, 0.92). The results were essentially consistent among subgroups stratified by participant characteristics. The findings of the present meta-analysis indicate that dietary lutein and zeaxanthin is not significantly associated with a reduced risk of early AMD, whereas an increase in the intake of these carotenoids may be protective against late AMD. However, additional studies are needed to confirm these relationships.

Key words: Lutein; Zeaxanthin; Age-related macular degeneration; Meta-analysis

Age-related macular degeneration (AMD), the leading cause of legal blindness in people aged over 65 years in industrialised countries, is a progressive disorder primarily affecting the macula, the central region of the retina involved with central vision(1). Early AMD is characterised clinically by yellowish deposits known as soft drusen accumulations and pigmentary abnormalities in the retinal pigment epithelium (RPE) and Bruch’s membrane, whereas late-stage manifestations encompass atrophy of photoreceptors and the RPE underlying it, choroidal neovascularisation, subretinal haemorrhage, detachment of RPE and retinal scarring(2). Currently, it has been reported that more than ten million people in the USA and approximately fifty million worldwide suffer from AMD(3). In the UK, almost 200 000 people aged 75 years or older were visually impaired due to AMD(4). Owing to the sharp rise in the elderly population, the disease has brought a huge burden for the health care system and had a profound impact on the quality of life and independence of older individuals. It is estimated that by the year 2020 the number of patients with late AMD will be increased by more than 50% to almost three million in the USA alone(5).

Abbreviations: AMD, age-related macular degeneration; AREDS, Age-Related Eye Disease Study; ICGS, International Classification and Grading System; RPE, retinal pigment epithelium; RR, relative risk; WARMGS, Wisconsin Age-related Maculopathy Grading System.

* Corresponding author: X.-M. Lin, fax +86 10 62015583, email linbjmu@bjmu.edu.cn
To date, the exact pathogenic mechanism of AMD has not been completely identified, and AMD is regarded as a complicated disease caused by the actions and interactions of multiple environmental risk factors and genetic factors. Although new therapies with anti-vascular endothelial growth factor agents have been shown to slow progressive visual loss effectively in certain types of neovascular AMD, these agents are costly and there are no treatments for geographic atrophy at present. Thus, identifying modifiable risk factors that could prevent or delay the onset of early AMD is considered to be far more preferable and of critical importance. Apart from being related to ageing, AMD has been associated with several modifiable risk factors, including smoking, sunlight exposure and diet.

Dietary antioxidants, particularly lutein and zeaxanthin, are hypothesised to have the capacity to modulate defence and repair systems that operate in response to oxidative damage and inflammation on the basis of the notion that the retina is highly susceptible to oxidative stress because of its high concentration of oxygen and PUFA, in combination with an intense exposure to light, which have been implicated in the development of AMD. As the major components of macular pigment, lutein and its structural isomer, zeaxanthin, are thought to have the beneficial effects on preventing the onset and progression of AMD, Laboratory data suggested an important role for these two carotenoids in protecting the neural retina from photo-oxidative damage and the development of common visually disabling disorders by absorbing blue light and by quenching reactive oxygen species through powerful antioxidant activity. Currently, a number of epidemiological studies have evaluated the relationship between dietary intake of these xanthophyll carotenoids and the risk of AMD. Although some studies have suggested a possible protective effect with the consumption of lutein and zeaxanthin in diet, others failed to show this benefit.

Given the public health importance of clarifying the potential role of lutein and zeaxanthin in the prevention of AMD, we conducted a systematic review of the evidence to evaluate the relationship between dietary intake of lutein and zeaxanthin and the risk of AMD. We also investigated whether this association was differed by subtype of AMD.

All retrieved articles were examined by performing an initial screen of identified abstracts and titles. Any studies that did not address the association between lutein and zeaxanthin intake and AMD were excluded, and the full texts of the remaining articles were further checked for their suitability for the present meta-analysis. The full-text articles of all references selected after application of these criteria were reviewed by using the same criteria. The present systematic review and meta-analysis was limited to cohort studies. Randomised clinical trials would have been considered, but none were found. For inclusion, a study had to meet the following criteria: (1) the primary outcome was clearly defined as AMD; (2) dietary intake of lutein and zeaxanthin was quantified; (3) relative risks (RR) or relative odds and their CI (or sufficient data to calculate these) were reported; (4) potential confounders were controlled for by matching or multivariable analysis in the studies. If a series of articles was published from the same study, the report with the most updated data was selected for analysis, although previous articles could be reviewed to supplement missing data where applicable; in the case of duplicate publication, only one publication was included. All literature search and article review were conducted independently in a standardised manner by two investigators (L. M. and Y.-Q. W.). Wherever discrepancies between investigators occurred for inclusion or exclusion, a third investigator (X.-M. L.) was involved to adjudicate disagreements or uncertainties.

Methods

Selection of studies

The electronic databases MEDLINE, EMBASE, ISI Web of Science, CINAHL and the Cochrane Central Register of Controlled Trials in the Cochrane Library were searched for publications through April 2010 using the keywords lutein, zeaxanthin, xanthophyll or carotenoid in conjunction with each of the following words: AMD, age-related maculopathy, neovascular AMD, exudative AMD, choroidal neovascularisation and geographic atrophy. Language restrictions were not imposed. We also performed a manual search of references cited by the published original studies and relevant review articles on the topic for additional studies, and contacted authors and experts in the field to identify any ongoing studies.

Data abstraction and study quality assessment

All data were independently abstracted in duplicate by two investigators (L. M. and Y.-Q. W.) using a standardised protocol. All disagreements between investigators were resolved by referencing the original publication and by discussion with a third investigator (X.-M. L.). We also contacted the primary authors to request additional information. Study characteristics recorded were as follows: first author's name, year of publication, country of origin, study design, characteristics of the study population (sample size, sampling methods, source of population and distribution of age, sex, ethnicity and BMI), diagnosis method of AMD, classification and grading systems for AMD, dietary assessment tool, follow-up duration, the RR or relative odds overall and in each subgroup and the corresponding CI or standard errors, and the confounding factors matched or adjusted in the studies. If a study provided several risk estimates, the most completely adjusted estimate was extracted.

The outcomes of interest were early AMD and late AMD. The early AMD was defined by the presence of drusen, pigmented abnormalities in RPE or both, whereas the late AMD, including neovascular AMD and geographic atrophy, was defined by the presence of choroidal neovascularisation, detachment of RPE or geographic atrophy. The AMD cases were ascertained in the studies retrieved using fundus photograph or medical record review of visual acuity, on the basis of the criteria from the International Classification and Grading System (ICGS), the Wisconsin Age-related Maculopathy...
Grading System (WARMGS) or the modified Age-Related Eye Disease Study (AREDS).

Information on key indicators of study quality was also extracted, and assessment of methodological quality of each study was carried out by two of the investigators (L. M. and Y.-Q. W.) independently, according to the Newcastle–Ottawa Scale\(^2\). The Newcastle–Ottawa Scale judges the quality of studies based on three aspects of the study: selection of study groups (four criteria), comparability of study groups (one criterion) and assessment of the outcome (three criteria). Studies that fulfilled five or more of the Newcastle–Ottawa Scale criteria were considered to be categorised as good quality. Studies that met four or fewer of this criteria were considered fair quality or poor quality. Discrepancies regarding quality parameters were also decided by discussion and consensus.

**Statistical analyses**

For all studies, RR or relative odds and their 95 % CI were extracted or derived from data reported in articles. Because the absolute risk of AMD was low, relative odds were considered an approximation of RR. Pooled RR estimates were calculated using both fixed effects and DerSimonian and Laird random effects models, weighting individual study results by the inverse of their variances. Forest plots were used to visually assess the RR estimates and corresponding 95 % CI across studies. Heterogeneity across studies was assessed by conducting \( Q \) tests (significance level of \( P<0.10 \) and quantifying the degree of heterogeneity by estimating the \( I^2 \) statistic\(^2\). \( I^2 \) values of less than or equal to 25, 50 and \( \geq 75 \) % represent low, moderate and high heterogeneity, respectively. In case of significant heterogeneity, the sources of heterogeneity were explored and sensitivity analyses were performed. Variables included in the subgroup analyses were population source (population based v. volunteer based), country of origin (USA v. not USA), mean age of participants (\( \leq 65 \) v. \( > 65 \) years), method of diagnosis (fundus photography v. visual acuity criteria) and classification criteria of AMD (ICGS v. WARMGS v. AREDS modified). Sensitivity analyses were conducted to examine the contribution of each individual study by iteratively eliminating each study from the meta-analysis and comparing the point estimates including and excluding the study. To assess the potential of publication bias, we performed both the Egger test and Begg test and examined relative symmetry of individual study estimates around the overall estimate using a funnel plot in which log RR were plotted against their corresponding standard errors\(^2\)). Additionally, we initially intended to conduct the dose–response meta-analysis to evaluate the relationship between dietary intake of lutein and zeaxanthin and AMD risk. Nevertheless, most studies had to be excluded for no CI, no distribution of cases and control subjects by exposure level or insufficient dose data on each category of lutein and zeaxanthin intake. Thus, we were unable to evaluate their associations in the dose–response meta-analysis. All statistical analyses were conducted by using RevMan version 5.0 (Cochrane Collaboration; Oxford, UK) and Stata version 8.2 (Stata Corporation, College Station, TX, USA). All \( P \) values were two-sided, with statistical significance set at a level of 0.05.

**Results**

**Literature search**

The search strategy yielded a total of 3465 citations. After excluding duplicates and screening the titles or abstracts, full-text versions of the remaining seventy-three articles were then retrieved for detailed evaluation. Of these seventy-three articles, five articles (six studies) were included in the systematic review and meta-analysis\(^2\) (Fig. 1).

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References identified through literature search (n 3465)

Excluded (n 903)
- Duplicate references

References screened based on titles or abstracts (n 2562)

Excluded (n 2489)
- Review article or technique summary
- Correspondence or editorial
- Not original human research
- Did not address the association between lutein and zeaxanthin intake and AMD

References retrieved for full-text review (n 73)

Excluded (n 68)
- Cross-sectional studies: 10
- Case–control study: 3
- Outcome was not AMD: 7
- Studies on AMD progression: 3
- Non-dietary exposures: 21
- No relevant comparisons: 15
- Multiple publications: 4
- Conference abstracts with insufficient data: 5

Articles included in meta-analysis (n 5) (six studies)

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Fig. 1. Flow diagram of study selection process. AMD, age-related macular degeneration.
Table 1. Characteristics of the cohort studies of dietary intake of lutein and zeaxanthin and risk of age-related macular degeneration (AMD)

<table>
<thead>
<tr>
<th>Source (year)</th>
<th>Study participants</th>
<th>Exposure assessment</th>
<th>Follow-up (years)</th>
<th>Diagnosis method</th>
<th>Classification criteria</th>
<th>No. of early AMD cases</th>
<th>No. of late AMD cases</th>
<th>Controlled variables</th>
<th>Study quality*</th>
</tr>
</thead>
<tbody>
<tr>
<td>Van den Langenberg et al. (1998)(^{(29)})</td>
<td>1709 men and women aged 43–84 years in the USA</td>
<td>FFQ (100 items)</td>
<td>5</td>
<td>Colour fundus photographs</td>
<td>WARMGS</td>
<td>103</td>
<td>NR</td>
<td>Age, sex, smoking, energy intake, alcohol intake, CVD, and history of diabetes</td>
<td>Good</td>
</tr>
<tr>
<td>Van Leeuwen et al. (2005)(^{(23)})</td>
<td>4170 men and women aged 55–95 years in the Netherlands</td>
<td>FFQ (170 items)</td>
<td>8</td>
<td>Colour fundus photographs</td>
<td>ICGS</td>
<td>518</td>
<td>42</td>
<td>Age, sex, smoking, BMI, alcohol intake, SBP, atherosclerosis composite score, and serum total cholesterol</td>
<td>Good</td>
</tr>
<tr>
<td>Moeller et al. (2006)(^{(30)})</td>
<td>1787 female healthy volunteers aged 50–79 years in the USA</td>
<td>FFQ (122 items)</td>
<td>7</td>
<td>Colour fundus photographs</td>
<td>AREDS modified</td>
<td>327</td>
<td>34</td>
<td>Age, smoking, family history of AMD, iris colour, history of diabetes and CVD, and use of hormone replacement therapy use</td>
<td>Fair</td>
</tr>
<tr>
<td>Tan et al. (2008)(^{(24)})</td>
<td>2454 men and women aged 45–93 years in Australia</td>
<td>FFQ (145 items)</td>
<td>10</td>
<td>Colour fundus photographs</td>
<td>WARMGS</td>
<td>266</td>
<td>72</td>
<td>Age, sex, smoking, white cell count, family history of AMD, and job prestige</td>
<td>Good</td>
</tr>
<tr>
<td>Cho et al. (2008)(^{(31)})†</td>
<td>41 564 male health professionals aged 50–79 years in the USA</td>
<td>FFQ (130 items)</td>
<td>16</td>
<td>Signs of AMD, BCVA loss ≤ 20/30</td>
<td>ICGS</td>
<td>210</td>
<td>162</td>
<td>Age, smoking, BMI, energy intake, alcohol intake, and fish intake</td>
<td>Fair</td>
</tr>
<tr>
<td>Cho et al. (2008)(^{(31)})‡</td>
<td>71 494 female registered nurses aged 50–79 years in the USA</td>
<td>FFQ (130 items)</td>
<td>18</td>
<td>Signs of AMD, BCVA loss ≤ 20/30</td>
<td>ICGS</td>
<td>463</td>
<td>280</td>
<td>Age, smoking, BMI, energy intake, alcohol intake, fish intake, and postmenopausal hormone use</td>
<td>Fair</td>
</tr>
</tbody>
</table>

WARMGS, Wisconsin Age-related Maculopathy Grading System; NR, not reported; ICGS, International Classification and Grading System; SBP, systolic blood pressure; AREDS, Age-Related Eye Disease Study; BCVA, best corrected visual acuity.

* Study quality was judged based on Newcastle–Ottawa Scale\(^{(25)}\).
† The Health Professionals’ Follow-Up Study conducted by Cho et al.\(^{(31)}\).
‡ The Nurses’ Health Study conducted by Cho et al.\(^{(31)}\).
Study characteristics

The characteristics of these studies are presented in Table 1. Of the six longitudinal cohort studies, four were conducted in the USA (29–31), one in Australia (24) and one in the Netherlands (23). The number of subjects ranged from 1709 to 71,494, comprising a total of 2477 incident AMD cases and 0.12 million participants. The study population in three studies included both men and women (23,24,29), two consisted entirely of women (30,31) and one consisted of only men (31). The duration of follow-up ranged from 5 to 18 years. Three were population-based studies (24,29,30), whereas three studies consisted of volunteers (30,31). Most of the studies had follow-up rates of 76.6% or more, with the exception of the studies by Cho et al. (31), which did not report them. The association between dietary intake of lutein and zeaxanthin and the risk of early AMD was the primary outcome of interest for all studies, whereas four studies also reported late AMD (24,30,31).

Diagnosis of AMD was based on fundus photography in four of the studies (23,24,29,30) and on review of medical records in two (31). Three studies used the ICGS criteria to establish AMD (23,31), whereas the WARMGS criteria and the AREDS modified criteria were applied in two studies (24,29) and one study (30), respectively. Dietary intake information was collected using a FFQ in all the studies. All of the studies adjusted for age and smoking, fewer adjusted for alcohol intake (four studies) (23,24,29,31), sex (three studies) (23,24,29), energy intake (three studies) (29,31), BMI (three studies) (23,31), history of AMD (two studies) (24,30), CVD (two studies) (23,30) or postmenopausal hormone use (two studies) (24,31).

Lutein and zeaxanthin intake and early age-related macular degeneration

All studies (23,24,29–31) reporting on the relationship between dietary lutein and zeaxanthin intake and early AMD risk were considered for inclusion in meta-analysis. Among the selected studies, all but one (31) found an association between intake of lutein and zeaxanthin and a reduced risk of early AMD, and only one study was statistically significant (24).

To explore the study heterogeneity, we performed stratified analyses across a number of participant characteristics (Table 2). These factors did not significantly alter the shape of association between lutein and zeaxanthin and early AMD risk, although population source and definition of AMD seemed to be slightly related with the results. Population-based studies tended to report a slightly stronger association of lutein and zeaxanthin intake with early AMD incidence, whereas the pooled estimate of volunteer-based studies was larger in magnitude (pooled RR 1.07; 95% CI 0.80, 1.45).

Likewise, inconsistencies in diagnosis method and separate criteria of AMD might somewhat affect the findings for lutein and zeaxanthin intake and early AMD risk. In sensitivity analyses, exclusion of any single study from the analyses did not markedly influence the overall results.

Visual inspection of the funnel plot for the studies evaluating lutein and zeaxanthin intake and its association with early AMD revealed symmetry (Fig. 3(a)). The Egger test (P = 0.76) and Begg test (P = 0.85) suggested no significant asymmetry of the funnel plot, indicating the absence of substantial publication bias.

Lutein and zeaxanthin intake and late age-related macular degeneration

Four studies (24,30,31) were included in the analysis of the relationship between dietary lutein and zeaxanthin intake and late AMD risk. The point estimates of the RR were consistently less than 1 in all studies, and none of studies reported statistically significant associations. A forest plot showing results of the comparison between the highest and lowest categories of lutein and zeaxanthin intake is shown in Fig. 4.

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**Table 1.** Study characteristics

<table>
<thead>
<tr>
<th>Study (year)</th>
<th>RR (95% CI)</th>
<th>Weight (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>van den Langenberg et al. (1998) (29)</td>
<td>0.93 (0.49, 1.76)</td>
<td>7.5</td>
</tr>
<tr>
<td>van Leeuwen et al. (2005) (23)</td>
<td>1.00 (0.77, 1.29)</td>
<td>21.5</td>
</tr>
<tr>
<td>Moeller et al. (2006) (30)</td>
<td>0.98 (0.77, 1.26)</td>
<td>22.3</td>
</tr>
<tr>
<td>Tan et al. (2008) (24)</td>
<td>0.66 (0.48, 0.92)</td>
<td>17.7</td>
</tr>
<tr>
<td>Cho et al. (2008) (31)*</td>
<td>1.66 (1.04, 2.64)</td>
<td>11.9</td>
</tr>
<tr>
<td>Cho et al. (2008) (31)†</td>
<td>0.89 (0.66, 1.20)</td>
<td>19.1</td>
</tr>
<tr>
<td>Pooled RR</td>
<td>0.96 (0.78, 1.17)</td>
<td>100.0</td>
</tr>
</tbody>
</table>

Fig. 2. Forest plot of relative risk (RR) and 95% CI for highest v. lowest category of dietary lutein and zeaxanthin intake and early age-related macular degeneration risk. * The Health Professionals’ Follow-Up Study conducted by Cho et al. (31). † The Nurses’ Health Study conducted by Cho et al. (31).
There was little evidence of heterogeneity among studies, with an $I^2$ of 0% ($P$ value for heterogeneity $= 0.86$). Overall, compared with individuals in the lowest intake of lutein and zeaxanthin, those in the highest intake had a significantly reduced risk of developing late AMD, with a fixed effects pooled RR of 0.74 (95% CI 0.57, 0.97). In a subgroup analysis of the three studies (24,31) that also reported on the RR of neovascular AMD, we estimated a reduction of 32% in neovascular AMD risk associated with high lutein and zeaxanthin intake (pooled RR 0.68; 95% CI 0.51, 0.92), and there was no statistically significant heterogeneity among these studies ($I^2 = 0$%; $P$ for heterogeneity $= 0.38$). In addition, stratified analyses showed consistency in the direction of effect when studies were grouped by the characteristics of participants. The sensitivity analyses showed minimal influence on the overall pooled estimate and heterogeneity for any single study. Fig. 3(b) displays a funnel plot for the visual assessment of publication bias. The funnel plot was symmetrical, and neither the Egger test ($P = 0.97$) nor the Begg test ($P = 1.00$) suggested publication bias.

### Discussion

The present meta-analysis involved data on evaluating the effects of lutein and zeaxanthin on AMD prevention published in six cohort studies. Results from the present study suggested that high intake of lutein and zeaxanthin was significantly associated with a reduction in risk of late AMD. No significant relationship was found for dietary intake of these carotenoids and early AMD. We conducted some stratified analyses across participant characteristics, with essentially no change in the findings of the present study.

Several biological mechanisms have been proposed for the potential protective effect of lutein and zeaxanthin on preventing the onset of AMD. As the major components of macular pigment, lutein and zeaxanthin are uniquely concentrated at the macula, indicating that these carotenoids may exert their effects on protecting the macula from age-related loss of visual function and macular disease (32). Both lutein and zeaxanthin, possessing a series of unconjugated double bonds, are believed to be very effective antioxidants (33).

### Table 2. Stratified analysis of the association between dietary intake of lutein and zeaxanthin and age-related macular degeneration (AMD)

<table>
<thead>
<tr>
<th>Subgroup</th>
<th>n</th>
<th>Pooled RR</th>
<th>95% CI</th>
<th>Heterogeneity</th>
<th>Meta-regression</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Early AMD</strong></td>
<td></td>
<td></td>
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<td></td>
<td></td>
</tr>
<tr>
<td>Population source</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Population based</td>
<td>3</td>
<td>0.85</td>
<td>0.63, 1.13</td>
<td>0.16</td>
<td>0.29</td>
</tr>
<tr>
<td>Volunteer based</td>
<td>3</td>
<td>1.07</td>
<td>0.80, 1.45</td>
<td>0.08</td>
<td></td>
</tr>
<tr>
<td>Country of origin</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>USA</td>
<td>4</td>
<td>1.04</td>
<td>0.82, 1.33</td>
<td>0.16</td>
<td>0.34</td>
</tr>
<tr>
<td>Not USA</td>
<td>2</td>
<td>0.82</td>
<td>0.55, 1.23</td>
<td>0.06</td>
<td></td>
</tr>
<tr>
<td>Age (years)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>$\leq 65$</td>
<td>4</td>
<td>0.95</td>
<td>0.66, 1.38</td>
<td>0.02</td>
<td>0.86</td>
</tr>
<tr>
<td>$&gt; 65$</td>
<td>2</td>
<td>0.99</td>
<td>0.83, 1.18</td>
<td>0.95</td>
<td></td>
</tr>
<tr>
<td>Method of diagnosis</td>
<td></td>
<td></td>
<td></td>
<td></td>
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</tr>
<tr>
<td>Fundus photography</td>
<td>4</td>
<td>0.89</td>
<td>0.74, 1.09</td>
<td>0.35</td>
<td>0.17</td>
</tr>
<tr>
<td>Visual acuity criteria</td>
<td>2</td>
<td>1.18</td>
<td>0.64, 2.17</td>
<td>0.03</td>
<td></td>
</tr>
<tr>
<td>Classification criteria*</td>
<td></td>
<td></td>
<td></td>
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</tr>
<tr>
<td>ICGS</td>
<td>3</td>
<td>1.08</td>
<td>0.80, 1.46</td>
<td>0.08</td>
<td>0.29</td>
</tr>
<tr>
<td>WAMGS</td>
<td>2</td>
<td>0.71</td>
<td>0.53, 0.95</td>
<td>0.36</td>
<td></td>
</tr>
<tr>
<td>AREDS modified</td>
<td>1</td>
<td>0.98</td>
<td>0.77, 1.26</td>
<td>NA</td>
<td></td>
</tr>
<tr>
<td><strong>Late AMD</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Population source</td>
<td></td>
<td></td>
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</tr>
<tr>
<td>Population based</td>
<td>1</td>
<td>0.72</td>
<td>0.34, 1.50</td>
<td>NA</td>
<td>NP</td>
</tr>
<tr>
<td>Volunteer based</td>
<td>3</td>
<td>0.75</td>
<td>0.56, 1.00</td>
<td>0.69</td>
<td></td>
</tr>
<tr>
<td>Country of origin</td>
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<tr>
<td>USA</td>
<td>3</td>
<td>0.75</td>
<td>0.56, 1.00</td>
<td>0.69</td>
<td>NP</td>
</tr>
<tr>
<td>Not USA</td>
<td>1</td>
<td>0.72</td>
<td>0.34, 1.50</td>
<td>NA</td>
<td></td>
</tr>
<tr>
<td>Age (years)</td>
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<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>$\leq 65$</td>
<td>3</td>
<td>0.72</td>
<td>0.54, 0.97</td>
<td>0.78</td>
<td>NP</td>
</tr>
<tr>
<td>$&gt; 65$</td>
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<td>0.89</td>
<td>0.43, 1.81</td>
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<td>Fundus photography</td>
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<td>0.89</td>
<td>0.43, 1.81</td>
<td>NA</td>
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</table>

ICGS, International Classification and Grading System; WAMGS, Wisconsin Age-related Maculopathy Grading System; AREDS, Age-Related Eye Disease Study; NA, not applicable because only one study; NP, meta-regression was not possible.

* Meta-regression was performed for the first two categories.
shown to help quench singlet oxygen, scavenge reactive free radicals and inhibit lipid peroxidation of membrane phospholipids, and thus may prevent or delay the development of AMD\(^{34,35}\). In addition, the spectrum of lutein and zeaxanthin includes a broad absorption band, with a peak at approximately 450 nm; therefore, these carotenoids have an important role in absorbing and attenuating the damaging blue light before it reaches the photoreceptors\(^{30}\). Results from animal studies also showed that macular xanthophylls could prevent or retard some of the destructive processes that ultimately led to AMD\(^{19,37}\). Furthermore, vascular defect and inflammation may be partly responsible for the pathogenesis of AMD, particularly of neovascular AMD\(^{38,39}\). There was evidence that lutein and zeaxanthin had the capacity to reduce thickening of the arteries and maintain the normal vascular function of retina and choroid\(^{15,40}\).

As there were no randomised clinical trials regarding the effect of dietary lutein and zeaxanthin on AMD prevention at present, the greatest interpretative weight associated with results could be obtained from the cohort design. Several other types of observational studies had previously examined the association between the intake of these carotenoids and the risk of AMD\(^{41–45}\). In contrast with the results from the cohort studies, almost all case–control and cross-sectional studies reported statistically significant associations between lutein and zeaxanthin intake and AMD risk, indicating that the combination of different study designs in the present meta-analysis would bias the present results towards the positive outcomes. Furthermore, case–control and cross-sectional studies might be inherently biased by various factors. It is generally considered that cohort studies provide stronger evidence for evaluating a relationship than other observational studies, because the cohort studies could largely reduce the likelihood of selection bias and reverse causation. Therefore, only cohort studies were included in the present systematic review and meta-analysis.

Results from the present analysis showed that lutein and zeaxanthin intake was not significantly associated with a decrease in the risk of developing early AMD. Among the six available studies, only one\(^{24}\) found a significant association between dietary lutein and zeaxanthin and the incidence of early AMD, whereas the others found no associations that were consistent with the present finding. Given the evidence that the older participants, who had a poorer survival than

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**Fig. 3.** Funnel plots with 95% CI for (a) early age-related macular degeneration (AMD) risk and (b) late AMD risk. RR, relative risk; se, standard error.

**Fig. 4.** Forest plot of relative risk (RR) and 95% CI for highest v. lowest category of dietary lutein and zeaxanthin intake and late age-related macular degeneration risk. * The Health Professionals’ Follow-Up Study conducted by Cho et al\(^{31}\). † The Nurses’ Health Study conducted by Cho et al\(^{31}\).
the younger participants did, were more likely to be AMD cases, the potential for survival bias should be considered. Such bias would increase with the average age of the sample and tend to under-report the strength of the association. However, findings from previous studies were not consistent. Moeller et al. found that associations of dietary lutein with early AMD were strengthened in younger, compared with older subjects. The present results showed no significant differences in RR stratified by age of participants, suggesting that the age of samples was not likely to be a strong contributor to heterogeneity. It is worth noting that meta-analyses of published data do not permit an adequate evaluation of age interactions. Further evaluations using person-specific data pooling are needed to better evaluate this possibility. We also repeated some analyses stratified by other participant characteristics, with essentially no change in these findings. Moreover, we conducted a sensitivity analysis and also found that the shape of association remained unchanged. The consistencies between these findings indicated that the associations were robust and combining studies with different participant characteristics did not bias the present results. In contrast with the findings for early AMD, we found a statistically significant relationship between lutein and zeaxanthin intake and the risk of late AMD. These inconsistent relationships from different stages of AMD might be partly explained by differences in the degree of macular pigment damage and ascertainment of AMD. Previous studies had shown that no significant differences in macular pigment optical density were found between eyes with and without early AMD or between the various stages of early AMD. Results from studies that compared the macular pigment optical density of eyes with and without late AMD were not consistent; however, most indicated declines in the optical density of macular pigment among subjects with late AMD. Similar results had also been found in the peripheral retina of autopsy specimens from donor eyes with AMD, indicating that the loss of macular pigment might reflect the accumulation of damage accrued over an entire lifespan. As an indicator of the macular pigment status of the retina, dietary intake of lutein and zeaxanthin is more likely to be related with late AMD, rather than early AMD. On the other hand, an alternative explanation for the discrepancies between the relationships of early AMD and late AMD could be due to the variety of diagnosis methods and classification criteria of AMD in different studies. Most of the studies support the notion that fundus photography is highly more accurate for detection of AMD, whereas the definition of AMD based on visual acuity criteria is subject to inter-individual variability. As AMD often progresses without any symptoms, many patients with AMD at early stage are not easy to distinguish only by routine ophthalmic examination. Thus, the inconsistencies of AMD assessment methods in the original studies may cause random misclassifications, which would be more likely to underestimate the true association, in particular, of early AMD. In addition, the present finding was in agreement with the subgroup analyses by subtype of late AMD. Results from the present study showed that dietary intake of lutein and zeaxanthin had a significant inverse association with neovascular AMD risk and this pooled RR was even stronger, suggesting these carotenoids might serve as a function in modulating inflammatory response and promoting flow of blood to and from the macular region.

The present meta-analysis has several strengths. The present review included a large number of people from different studies. Most of the studies had a large sample size and a long duration of follow-up. Almost all the studies had adjusted for age and smoking in the analyses. These relatively high-quality studies combined should give more reliable assessment of the relation between lutein and zeaxanthin and risk of AMD. Additionally, the association was essentially consistent among subgroups stratified by characteristics of participants, indicating that the conclusions of the present study were not dependent on arbitrary decisions in the present meta-analysis. Finally, the present results were unlikely to result from publication bias, as indicated by the funnel plots and other analyses.

The present study also has several limitations that merit consideration. First, the present study was based on observational studies and might have the problems of potential bias and confounding effects associated with such studies. Second, inconsistencies in methods of diagnosing AMD might have contributed to the inconsistent results among studies. Although case diagnosis of AMD in all the studies had been validated on the basis of the generally accepted standard methods, the accuracy of visual acuity criteria for detection of AMD was limited compared with fundus photography, because early signs of AMD were asymptomatic in most people. In the present analysis, the RR of the both forms of AMD associated with lutein and zeaxanthin did not materially change after exclusion of studies that did not use fundus photography for diagnosis. However, such bias could not be ruled out completely, which might have led to underestimation or overestimation of the association. Furthermore, the validated classification and grading systems for AMD were inconsistently applied between studies because of the different countries and periods in which the studies were performed. The present results were also likely to be affected by different separate criteria of AMD. Third, another limitation of the present study concerned the assessment methods of dietary intakes of lutein and zeaxanthin. Although all studies included in the present meta-analysis stated that the FFQ used to assess dietary intake had been validated, dietary intake was measured only at baseline and might not reflect changes in intake of these carotenoids during follow-up. Differences in nutrient databases used across the studies could also constitute potential sources of variability in the values of lutein and zeaxanthin. Fourth, even though several confounding factors had been adjusted for in all the studies incorporated, the possibility of other uncontrolled or potential residual confounding could not be fully excluded in the present meta-analysis. Moreover, lutein and zeaxanthin intake tended to be associated with some healthy lifestyles or dietary patterns that might be protective against AMD. Consequently, it was difficult to separate beneficial effects of dietary lutein and zeaxanthin intake from effects due to healthy lifestyle or dietary habits. This might be a possible explanation.
for the inconsistent findings from different population sources. Fifth, it is possible that populations with specific genetic backgrounds or nutritional status may affect the analysis of the relationship\(^{(66)}\). The present results were mainly based on studies carried out in Western populations, which limited the generalisation of findings. More research is necessary to be conducted in different populations to examine variations between populations. Finally, as with any meta-analysis, the potential for publication bias is a concern. Despite no publication bias examined in the present study, it was still difficult to fully rule out such bias because there was not a sufficient number of studies to detect it adequately.

In conclusion, the present systematic review and meta-analysis demonstrates that, on the basis of evidence available to date, dietary intake of lutein and zeaxanthin is not significantly associated with a decrease in the risk of developing early AMD, whereas an increase in the intake of xanthophylls may have beneficial effects for late AMD. It should also be noted that there are only a few studies that have examined this association, which limits the power of meta-analysis. Therefore, further well-designed large studies with prospective cohort design are required before definitive conclusions can be drawn regarding the potential effect of dietary lutein and zeaxanthin on AMD prevention.

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References

Lutein and age-related macular degeneration


