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1 **Lifetime exposure to ambient ultraviolet radiation and the risk for cataract**
2 **extraction and age-related macular degeneration: the Alienor Study**

3

4 Cécile Delcourt^{1,2}, Audrey Cougnard-Grégoire^{1,2}, Mathieu Boniol³, Isabelle Carrière^{4,5}, Jean-François
5 Doré⁶, Marie-Noëlle Delyfer^{1,2,7}, Marie-Bénédicte Rougier⁷, Mélanie Le Goff^{1,2}, Jean-François
6 Dartigues^{1,2}, Pascale Barberger-Gateau^{1,2}, Jean-François Korobelnik^{1,2,7}.

7

8

9 1 : Univ. Bordeaux, F-33000 Bordeaux, France ;

10 2 : INSERM, ISPED, Centre INSERM U897-Epidemiologie-Biostatistique, F-33000 Bordeaux, France

11 3 : International Prevention Research Institute, Lyon, France

12 4 : Inserm, U1061, Montpellier, F-34093 France

13 5 : University Montpellier I, Montpellier, F-34000 France

14 6 : Centre de Recherche en Cancérologie de Lyon, UMR Inserm U1052 - CNRS U5286, Lyon, F-
15 69008, France

16 7 : CHU de Bordeaux, Service d'Ophtalmologie, Bordeaux, F-33000, France

17

18

19 Corresponding author/reprint request : Cécile Delcourt, Inserm U897, ISPED, Université Bordeaux
20 Segalen, 146 rue Léo Saignat, 33076 Bordeaux Cedex. Tel : +33 5 47 30 42 04; Fax : +33 5 57 57 14
21 86 ; email : cecile.delcourt@isped.u-bordeaux2.fr

22

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29

30 **ABSTRACT**

31 Purpose: While exposure to ultraviolet radiation is a recognized risk factor for cataract, its association
32 is more controversial with age-related macular degeneration (AMD). We report the associations of
33 lifetime exposure to ambient ultraviolet radiation (UVR) with cataract extraction and AMD.

34 Methods: The Alienor Study is a population-based study of 963 residents of Bordeaux (France), aged
35 73 years or more. Lifetime exposure to ambient UVR was estimated from residential history and
36 Eurosun satellite-based estimations of ground UVR. It was divided in 3 groups (lower quartile,
37 intermediate quartiles, upper quartile), using the intermediate quartiles as the reference. Early and late
38 AMD were classified from retinal colour photographs. Cataract extraction was defined as absence of
39 the natural lens at slit lamp.

40 Results: After multivariate adjustment, subjects in the upper quartile of lifetime ambient UVR exposure
41 were at increased risk for cataract extraction (OR=1.53, 95 % confidence interval (CI): 1.04-2.26,
42 p=0.03) and for early AMD (OR= 1.59, 95 % CI: 1.04-2.44, p=0.03), by comparison with subjects in the
43 intermediate quartiles. Subjects in the lower quartile of UVR exposure were also at increased risk for
44 early AMD (OR=1.69, 95 % CI: 1.06-2.69, p=0.03), by comparison with those with medium exposure.
45 Associations of late AMD with UVR exposure was not statistically significant.

46 Conclusions: This study further confirms the increased risk for cataract extraction in subjects exposed
47 to high ambient UVR. Moreover, it suggests that risk for early AMD is increased in subjects exposed to
48 high UVR, but also to low UVR, by comparison with medium exposures.

49

50 Keywords: macular degeneration, cataract, light exposure, ultraviolet radiation, risk factors,
51 epidemiology

52

53

54 Cataract and age-related macular degeneration (AMD) are leading causes of blindness and visual
55 impairment worldwide ¹. Both of these diseases are multifactorial, involving non-modifiable (e.g. age,
56 gender, genetics) and modifiable factors ^{2,3}. The control of these modifiable factors may represent a
57 preventive strategy for decreasing the incidence of these diseases and of the related visual
58 impairment. Among other modifiable factors (in particular smoking and nutrition) ^{2,3}, the potential role
59 of sunlight exposure in the aetiology of these diseases has been investigated ^{4 5}. The absorption of
60 solar radiations by biological tissues results in photochemical reactions and the formation of reactive
61 oxygen species (including singlet oxygen) which may damage all cellular components (lipids, proteins,
62 DNA)⁶. The part of solar radiations that interacts with the eye is known as “optical radiations” and
63 includes wavelengths from ultraviolet (100-400 nm), visible light (400-760 nm) to infrared (760-10,000
64 nm). Ultraviolet radiations represent the most energetic part of optical radiations, and are thus
65 responsible for a large part of photochemical damage.

66 The crystalline lens is particularly exposed to phototoxic damage, because it absorbs most of
67 ultraviolet radiations (UVR), together with the cornea. This has been confirmed by epidemiological
68 studies, which have shown consistent associations of cataract (resulting from lens opacification) with
69 sunlight exposure and, in particular, UVR exposure ⁷⁻¹³. Ultraviolet exposure has been consistently
70 associated with the risk for cataract in numerous studies, performed in different continents with
71 different methodologies, showing dose-dependent relationships, and specific associations with cortical
72 cataracts. It is now a recognized risk factor for cataract ².

73 By contrast, epidemiological data regarding the associations of light exposure with the risk for AMD
74 remain scarce and inconsistent. Several studies have suggested an increased risk for AMD in subjects
75 highly exposed to sunlight ¹⁴⁻¹⁷, but others showed no significant associations ¹⁸⁻²⁰ and some even
76 suggested a decreased risk for AMD in the most exposed subjects ^{21, 22}.

77 In the present study, we report the associations of lifetime exposure to ambient UVR with the risk for
78 cataract extraction and AMD, in the framework of a population-based cross-sectional study of elderly
79 subjects from the South of France.

80

81 **Subjects and Methods**

82 ***Study aims***

83 The Alienor (Antioxydants, Lipides Essentiels, Nutrition et maladies OculaiRes) Study is a population-
84 based prospective study aiming at assessing the associations of age-related eye diseases (AMD,
85 glaucoma, cataract, dry eye syndrome) with nutritional factors (in particular antioxidants, macular
86 pigment and fatty acids).²³ It also takes into account other major determinants of eye diseases,
87 including gene polymorphisms, lifestyle and vascular factors.

88

89 ***Study sample***

90 Subjects of the Alienor Study were recruited from an ongoing population-based study on the vascular
91 risk factors for dementia, the Three City (3C) Study²⁴. The 3C Study included 9,294 subjects aged 65
92 years or more from three French Cities (Bordeaux, Dijon and Montpellier), among which 2,104 were
93 recruited in Bordeaux. Subjects were contacted individually from the electoral rolls. They were initially
94 recruited in 1999-2001 and followed-up about every two years since. The Alienor study consists in eye
95 examinations, which are offered to all participants from the third follow-up (2006-2008) of the 3C
96 cohort in Bordeaux. Among the 1,450 participants re-examined in 2006-2008, 963 (66.4 %)
97 participated in the first eye examination of the Alienor Study.

98 This research followed the tenets of the Declaration of Helsinki. Participants gave written consent for
99 the participation in the study. The design of this study has been approved by the Ethical Committee of
100 Bordeaux (Comité de Protection des Personnes Sud-Ouest et Outre-Mer III) in May 2006.

101

102 ***Eye examination***

103 The eye examination took place in the Department of Ophthalmology of the University Hospital of
104 Bordeaux. It included a recording of ophthalmological history, measures of visual acuity, refraction,
105 two 45° non mydriatic color retinal photographs (one centred on the macula, the other centred on the
106 optic disc), measures of intraocular pressure and central corneal thickness and tear film break-up time
107 test. A self-completed questionnaire on risk factors specific to the eye (including residential history for

108 estimation of light exposure) and dry eye symptoms was filled at home and brought back on the day of
109 the eye examination.

110 Retinal photographs were performed using a high-resolution digital non mydriatic retinograph (TRC
111 NW6S, Topcon, Japan). Photographs were interpreted in duplicate by two specially trained
112 technicians. Inconsistencies between the two interpretations were adjudicated by a retina specialist for
113 classification of AMD and other retinal diseases, or by a glaucoma specialist for classification of
114 glaucoma. All cases of late AMD, other retinal diseases and glaucoma were reviewed and confirmed
115 by specialists. Graders had no access to light exposure variables, or any of the potential confounders.

116

117 ***Cataract extraction***

118 In each eye, cataract extraction was defined as the absence of the natural crystalline lens at slit lamp.

119

120 ***Classification of AMD***

121 Retinal photographs were interpreted according to the international classification²⁵ and to a
122 modification of the grading scheme used in the Multi-Ethnic Study of Atherosclerosis for drusen size,
123 location and area²⁶. Eyes were classified according into one of the three exclusive groups: no AMD,
124 early AMD, late AMD.

125 Late AMD was defined by the presence of neovascular AMD or geographic atrophy within the grid
126 (3000 microns from the foveola). Neovascular AMD included serous or hemorrhagic detachment of the
127 retinal pigment epithelium (RPE) or sensory retina, subretinal or sub-RPE hemorrhages and fibrous
128 scar tissue. Geographic atrophy was defined as a discrete area of retinal depigmentation, 175 microns
129 in diameter or larger, characterized by a sharp border and the presence of visible choroidal vessels.
130 Five cases of late AMD had no gradable photographs and were classified using ophthalmological
131 history of AMD and AMD therapy (in particular intravitreal antiangiogenic agents and photodynamic
132 therapy), and confirmed by their treating ophthalmologist.

133 Early AMD was defined by the presence of soft distinct drusen and/or soft indistinct drusen and/or
134 reticular drusen and/or pigmentary abnormalities. Soft distinct and indistinct drusen were larger than

135 125 microns in diameter and with uniform density and sharp edges or decreasing density from the
136 center outwards and fuzzy edges, respectively. Pigmentary abnormalities were defined as areas of
137 hyperpigmentation and/or hypopigmentation (without visibility of choroidal vessels).

138

139 ***Ambient UVR exposure***

140 For each participant, average annual ambient UVR exposure was estimated using the residential
141 history, by weighting annual ambient UVR at each location by the time spent at that location.

142 Residential history from birth (locations, time spent at each location) was self-declared up to the first
143 eye examination (2006-2008). In France, locations were divided in 101 geographical areas,
144 corresponding to the 101 administrative « departments » (95 in metropolitan France and 6 overseas
145 departments). Concerning foreign countries, ambient UVR was generally estimated for the capital of
146 the country, except when the capital was very off-centered, in which case a more central location was
147 chosen. Very large countries (United States, China, etc) were excluded from this analysis, since an
148 estimation of solar radiation in a single geographic location is meaningless.

149 Average annual ambient solar radiation was assessed for the first 65 years of life, since almost all
150 subjects (97.3 %) lived in Bordeaux area beyond this age.

151

152 *Ultraviolet radiation*

153 In each location, UVR was extracted from Eurosun UV database (www.eurosun-project.org). Briefly,
154 UV irradiation levels were initially extracted from surface solar irradiance derived from the Meteosat
155 satellite's images. From these irradiation levels, the UV component was computed by a model which
156 exploits the algorithm set up by the Royal Institute of Meteorology, Belgium, published in the European
157 Solar Radiation Atlas (spectral model of Joukoff ESRA). The algorithm converts the total irradiance (E)
158 into its spectral distribution $E(\lambda)$, every 10nm, and gives estimates for total UV (280-400 nm), UVA
159 (315-400 nm) and UVB (280-315 nm). The calculation of individual exposure assumes that the
160 irradiation levels in different regions remained constant over the years. The information provided by
161 Eurosun corresponds to the average of daily UV radiation over the period 1988-2007 for each location.

162 We conducted analysis of change in UV radiation over the period 1988-2007 in Eurosun maps and
163 found only minor changes of less than 1% yearly change. This observations were in line with the
164 results obtained from COST 726 which also reported a stability of UV irradiation in Europe for a period
165 of 50 years of data up to 2002 (the project COST 726's report and details are available at [http://www-
166 med-physik.vu-wien.ac.at/uv/cost726/cost726.htm](http://www-med-physik.vu-wien.ac.at/uv/cost726/cost726.htm)). So these analyses from Eurosun data and
167 observation from COST 726 action are suggesting a stability of UV which let us assume no change
168 over time. These estimates were available for European and North African countries, with a resolution
169 of 5km. However, they are not available for other countries (Americas, Sub-saharian Africa, Asia,
170 Oceania).

171

172 *Estimation of missing ultraviolet radiation data using global solar radiation*

173 In each location, global ambient annual solar radiation (a measure of solar energy including all
174 wavelengths) was estimated using astronomical equations and the statistics of sunshine hours, using
175 the same methodology as in the POLA Study¹³. Overall, global ambient annual solar radiation
176 estimates were available in 116 locations (101 French departments, 7 European countries, 3 North
177 African countries, 5 other countries). In 105 locations (95 French departments, 7 European countries,
178 3 North African countries), both UVR and global solar radiation (GSR) estimates were available.

179 Pearson's correlation coefficient between these two variables was 0.952. For the 11 areas with
180 missing UVR but available solar radiation (6 French overseas departments and 5 countries), we thus
181 estimated ambient UVR from global solar radiation, using linear regression modeling. The regression
182 equation derived from the 105 locations with both UVR and GSR was: $UVR = 5613.105 + 0.0729 \times$
183 GSR ($r^2=0.91$). The same analyses were performed for UVA and UVB, leading to the following
184 equations: $UVA = 5467.398 + 0.0710 \times GSR$; $UVB = 145.708 + 0.00189 \times GSR$.

185 Finally, estimates of UVR exposure still could not be estimated for some countries. When, for a given
186 subject, the number of years spent in such countries was less or equal to 3 years, these countries
187 were eliminated from the calculations. For 71 subjects, average ambient UVR was therefore calculated
188 on 62, 63 or 64 years instead of 65. In addition, 113 subjects were excluded from the analyses
189 because they had spent more than 3 years in countries where UVR could not be estimated.

190

191 ***Other variables***

192 The following potential confounders have been selected, based on literature results reporting
193 significant associations of AMD and/or cataract and/or light exposure with age, sex, educational level,
194 diabetes, hypertension, asthma, oral corticosteroid use, smoking, physical activity, body mass index
195 (BMI), plasma glucose and lipids, dietary intake of energy, antioxidants, lutein and zeaxanthin and
196 omega3 fatty acids^{2,3}. We also included the two major genetic polymorphisms associated with AMD
197 Complement Factor H (CFH) Y402H (rs1061170) and Age-Related Maculopathy Susceptibility 2
198 (ARMS2) A69S (rs10490924) polymorphisms²⁷⁻²⁹. These genetic polymorphisms were also strongly
199 associated with AMD in the Alienor study^{30 31}. By comparison with subjects homozygous for the most
200 frequent allele (TT for CFH Y402H and GG for ARMS2 A69S), subjects homozygous for the less
201 frequent allele (CC for CFH Y402H, TT for ARMS2 A69S) exhibited 5- to 15- fold increased risk for
202 AMD. Heterozygous subjects (TC for CFH Y402H and GT for ARM2 A69S) exhibit intermediate risks
203 (2- to 5- fold increased risks). Indeed, all of these factors may represent confounders in the
204 relationship of AMD with lifetime ambient UVR, since lifestyle (smoking, physical activity, diet), health
205 conditions and genetic characteristics have been reported to vary according to geographical location,
206 and thus with lifetime ambient UVR³²⁻³⁴.

207 Data were collected during a face-to-face interview using a standardized questionnaire administered
208 by a trained psychologist or nurse. At baseline (1999-2000), general data included: demographic
209 characteristics, educational level and smoking. BMI (kg/m²) was calculated as weight/height² using
210 weight and height. Physical activity was assessed by 2 questions: "Do you practice sports?" (yes/no)
211 and "Do you perspire when you practice sports?" (Never/sometimes/most of the time/always). A 3-
212 level variable was computed to describe intensity of physical activity, as already published.³⁵ Plasma
213 glucose and lipids were measured at the Biochemistry Laboratory of the University Hospital of Dijon
214 from baseline fasting blood samples. Dietary intake of energy, antioxidants, lutein and zeaxanthin and
215 omega3 fatty acids were estimated from a 24h recall performed through face-to-face interview by
216 specifically trained dietitians in 2001-2002^{35,36}. Genetic polymorphisms were determined by the Lille
217 Génopôle, from the DNA samples collected at baseline (1999–2001).

218

219

220 **Statistical methods**

221 First, associations of ambient solar radiation with socio-demographic factors, lifestyle, biological and
222 dietary parameters were performed using chi-square test or analysis of variance.

223 Associations of cataract extraction and AMD with ambient UVR were estimated using logistic
224 Generalized Estimating Equations (GEE) models³⁷, which allow taking into account the data from both
225 eyes and their intra-individual correlations. Adjusted odds-ratios were estimated using cataract
226 extraction or AMD as the dependent variable, and ambient UVR and potential confounders as the
227 independent variables. Ambient UVR was divided in three groups (lower quartile, intermediate
228 quartiles, upper quartile), the medium category being the reference. With regard to AMD, two models
229 were performed (for early and late AMD, respectively), subjects without any AMD being the reference
230 in both models. With regard to cataract, subjects without cataract extraction were the reference group.

231 With regard to cataract extraction, potential confounders retained in the analysis were known risk
232 factors for cataract² (age, gender, smoking, diabetes, oral corticosteroid use and asthma) and factors
233 significantly associated with ambient UVR (at $p < 0.05$). Similarly, with regard to AMD, potential
234 confounders retained in the analysis were well known risk factors, which are strongly associated with
235 AMD in our cohort (age, gender, smoking³⁰, CFH³⁰ and ARMS2³¹ polymorphisms, dietary intake of
236 omega3 fatty acids³⁶) and factors significantly associated with ambient UVR (at $p < 0.05$). No
237 collinearity was detected between the variables included in the final models. No major confounding
238 was detected (variation by more than 10 % of the estimates of the odds-ratios when deleting one
239 confounder from the models). All statistical analyses were performed using statistical software (SAS,
240 version 9.1; SAS Institute Inc., Cary, NC).

241

242 **RESULTS**

243 Figure 1 presents the distribution of annual ambient UVR exposures (total UV, UVA and UVB),
244 estimated from residential history. These variables showed important inter-individual variability,
245 ranging respectively, from 32.39 kJ/cm² to 50.93 kJ/cm², 31.55 kJ/cm² to 49.61 kJ/cm² and 0.81
246 kJ/cm² to 1.32 kJ/cm². The central peaks correspond to subjects who spent all their life in the

247 Bordeaux area. As shown in Table 1 for total UVR, these subjects constitute the middle group
248 (intermediate quartiles), who were born and lived all their life in the South of France. By contrast, two
249 thirds of subjects from the lower quartile of ambient solar radiation were born in Central or Northern
250 France, where they spent about 23 years on average. Among subjects from the upper quartile, more
251 than a quarter were born in Northern Africa and remained there for 32 years on average
252 (corresponding to the independence of these countries in the 1960s) and 8.13 % were born in
253 Southern Europe. Although the definition of the intermediate quartiles is narrow (39.649-40.173
254 kJ/cm²), there is significant variability, with a mean of 38.375 kJ/cm² in the lower quartile, compared
255 with 42.718 kJ/cm² in the upper quartile. Similar findings were found for UVA and UVB (data not
256 shown).

257 Socio-demographic, lifestyle and biological characteristics of participants appear different according to
258 their lifetime solar exposure. As shown in Table 2, compared with subjects from the middle quartiles,
259 subjects in the higher quartile were more often males (42.58 % versus 31.74 %) and tended to have a
260 higher frequency of the TC genotype of CFH Y402H (47.12 % versus 37.43 %) and lower frequency of
261 TT genotype of CFH Y402H (45.03 % versus 49.21 %), while subjects from the lower quartiles had a
262 higher educational level (48.80 % with University degree versus 26.97 %) and tended to have a higher
263 frequency of the TC genotype of CFH Y402H (47.15 % versus 37.43 %) and lower frequency of TT
264 genotype of CFH Y402H (40.93 % versus 49.21%). As shown in Table 3, both subjects from the upper
265 and lower quartiles had more smoked during their life (18.36 % and 22.93 % having smoked more
266 than 20 pack-years respectively), by comparison with intermediate exposures (12.50 %). Subjects in
267 the upper quartile also had higher total energy intake (1835.7 Kcal/day versus 1678.2 Kcal/day).
268 These variables (education, smoking, CFH Y402H genotypes and total energy intakes) were therefore
269 considered as potential confounders in the analyses of association of solar radiation with cataract
270 extraction and AMD.

271 Cataract extraction status was available in at least one eye of 958 subjects (99.5%). Of those, 837 had
272 complete data for light exposure variables. In addition, 143 subjects had missing values in potential
273 confounders, leaving 694 subjects in the fully adjusted models, representing 1388 eyes among which
274 542 had undergone cataract surgery. Participants with missing data were not significantly different
275 from those without missing data, with regard to age, gender, ambient UVR or genetic polymorphisms
276 (data not shown). As shown in Table 4, after adjustment for confounders, by comparison with

277 participants exposed to medium ambient total UVR, participants exposed to high ambient total UVR
278 were at increased risk for cataract extraction (OR=1.53, 95 % CI: 1.04-2.26, p=0.03), while participants
279 with low exposure had similar risk as those with medium exposures (0.75 (0.51-1.12)). Similar results
280 were observed for UVA and UVB. Among the potential confounders included in the multivariate
281 models, those significantly associated with cataract surgery were age (OR=1.18 for 1-year increase,
282 95 % CI: 1.13-1.22, p<0.0001), gender (OR=1.86 for females versus males, 95 % CI: 1.23-2.80,
283 p=0.003), having smoked 20 pack-years or more (OR= 2.23, 95 % CI: 1.37-3.64, p=0.001), history of
284 asthma (OR=1.99 1.15-3.42, p=0.01).

285 AMD status was available in at least one eye of 875 subjects (91 %). Of those, 769 had complete data
286 for light exposure variables. In addition, 172 subjects had missing values in potential confounders,
287 leaving 597 subjects in the fully adjusted models, representing 1154 eyes among which 238 with early
288 AMD and 49 with late AMD. Participants with missing data were not significantly different from those
289 without missing data, with regard to age, gender, ambient UVR or genetic polymorphisms (data not
290 shown). Participants exposed to high ambient total UVR tended to be at increased risk for early AMD
291 (OR=1.59, 95 % CI: 1.04-2.44, p=0.03), by comparison with participants with medium exposures
292 (Table 5). In addition, participants exposed to low ambient solar radiation were increased risk for early
293 AMD (OR=1.69, 95 %CI: 1.06-2.69, p=0.03), by comparison with medium exposures. By contrast, no
294 statistically significant associations of late AMD with UVR exposures were found. Overall, results were
295 very similar for total UV, UVA and UVB exposures. Among the potential confounders included in the
296 multivariate models, those significantly associated with early AMD were age (OR=1.06 for 1-year
297 increase, 95% CI: 1.02-1.10, p=0.008), gender (OR=1.95 for females versus males, 95% CI: 1.19-
298 3.21, p=0.01), ARMS2 A69S GT genotype (OR=1.62, 95 % CI: 1.12-2.35, p=0.01), ARMS A69S TT
299 genotype (OR=12.09, 95 % CI: 4.63-31.55, p<0.0001) and CFH Y402H CC genotype (OR=1.89, 95%
300 CI: 1.13-3.16, p=0.01). The factors significantly associated with late AMD were age (OR=1.24, 95%
301 CI: 1.14-1.35, p<0.0001), gender (OR=2.93, 95 % CI: 1.16-7.43, p=0.02), ARMS A69S TT genotype
302 (OR=59.71, 95 % CI: 15.53-229.57, p<0.0001), CFH Y402H TC genotype (OR=3.49, 95% CI: 1.31-
303 9.30, p=0.009), CFH Y402H CC genotype (OR=6.68, 95 % CI: 2.05-21.77, p=0.001), dietary omega3
304 fatty acids intake (OR=0.63, 95% CI: 0.41-0.96, p=0.03) and secondary education (OR=7.46, 95 % CI:
305 1.63-34.05, p=0.01). We detected no significant interactions of genetic polymorphisms with UVR
306 exposure.

307

308 **DISCUSSION**

309 While confirming the well-known association of cataract with high ambient UVR,⁷⁻¹³ the present study
310 suggests a U-shaped association of early AMD with ambient UVR, with increased risk both in low- and
311 high-UVR exposures. The effects of UVR on human health generally follows such a U-shaped (or J-
312 shaped) relationship³⁸. Indeed, on one hand, high exposures have well documented effects on skin
313 diseases (skin cancers, sunburns and chronic sun damage...) and several eye diseases (acute
314 photokeratitis and conjunctivitis, acute solar retinopathy, pterygium, cortical cataract...)³⁸. On the other
315 hand, cutaneous UVR exposure is necessary for endogenous production of vitamin D, so that low
316 UVR exposures are associated with an increased risk for osteo-muscular diseases and possibly of
317 other diseases related to vitamin D deficiency (in particular several cancers, cardiovascular disease,
318 autoimmune diseases)³⁸⁻⁴⁰. Indeed, besides its effects on bone health, vitamin D has many cellular
319 effects, including regulation of cellular differentiation, proliferation, apoptosis and angiogenesis⁴⁰, and
320 anti-inflammatory properties⁴¹. Many of these processes are implicated in the physiopathology of
321 AMD and some studies have suggested that low vitamin D status may be associated with an
322 increased risk for AMD⁴²⁻⁴⁵. Whether vitamin D deficiency may explain the higher risk for AMD
323 observed with low sunlight exposure in the present study will need to be determined in future studies.

324

325 Only few epidemiological studies have addressed the potential link of AMD with UVR exposure, with
326 inconsistent results. This is probably due to major methodological difficulties in the assessment of
327 UVR exposure. Indeed, UVR exposure and its health effects result from distal factors (such as latitude,
328 season, cloud cover, stratospheric ozone levels and lower atmospheric pollution), conditioning
329 ambient UVR, in addition to proximal factors (in particular occupational and leisure exposures to
330 sunlight, use of hats and sunglasses, skin pigmentation and sun sensitivity...)⁴⁶ conditioning individual
331 exposure and response to exposure. Moreover, previous studies have shown that health effects of
332 UVR exposure are cumulative over the lifetime. Thus, estimation of lifetime ocular UVR exposure
333 requires detailed questionnaires on lifetime distal and proximal factors, combined with complex
334 modeling of their effects on ocular UVR exposure. Since most epidemiological studies in the field of
335 AMD, including the present study, addressed not only the effect of sunlight exposure, but also of many

336 other factors (smoking, cardiovascular risk factors, genetics, nutrition...), only partial information on
337 distal and/or proximal factors was collected, limiting the validity of the estimates of UVR exposure, and
338 the comparability of studies.

339 The Maryland Watermen Study included detailed questionnaires of ocular exposures, combined with
340 field and laboratory data, in order to assess ocular UVR exposure. In this study, ocular UVR exposure
341 was not significantly associated with prevalent AMD⁴⁷. In the Beaver Dam Eye Study, ambient UV-B
342 exposure, estimated from residential history, was not significantly associated with prevalent¹⁵ or
343 incident^{16,17} AMD, but subjects having spent more than 5 hours/day outside during summer in their
344 youth were more likely to have prevalent¹⁵ and incident AMD^{16,17}. In the POLA Study²², performed in
345 the South of France, the risk for prevalent early AMD was higher in subjects exposed to low ambient
346 solar radiation. In the Blue Mountains Eye Study, prevalent AMD was associated both with high and
347 low sun sensitivity index, by comparison with medium sun sensitivity index⁴⁸, although this was not
348 confirmed in the prospective analysis of the same cohort²⁰. Finally, an Australian case-control study,
349 AMD was associated with poorer tanning ability and lower time of ocular sunlight exposure²¹.

350 In the present study, only estimates of ambient UVR exposure were available, based on residential
351 history and satellite-based estimates of UVR, while no data were available on sun-related behaviors,
352 skin pigmentation or sun sensitivity. This may have led to significant misclassification of exposures,
353 since actual ocular UVR exposure may be quite different from ambient UVR exposure in those
354 subjects with very low time spent outdoors. However, it seems most likely that misclassification was
355 unrelated to ambient UVR (i.e. that there would be similar proportions of people with low/high outdoors
356 activities in areas with different ambient UVR), and was thus unlikely to have biased the associations
357 of eye diseases with UVR exposure. Moreover, the well known association of UVR exposure with
358 cataract was confirmed in the present study, suggesting that our measure of UVR exposure had some
359 validity.

360 While the lack of information on proximal factors is clearly a limitation, the use of satellite-based
361 estimates of UVR is a strength. Indeed, most previous studies relied on formulas, estimating UVR
362 mainly from latitude and altitude. While these are major determinants of UVR, weather conditions also
363 contribute to actual ground UVR, in particular because of attenuation by cloud cover. UVR estimations
364 in the present study rely on Meteosat satellite measurements of total solar irradiance by the earth's

365 surface by reflection, over thirty years.

366 Using detailed estimation of ocular lifetime exposure, the Maryland Watermen study and the EUREYE
367 study have suggested that AMD risk may be associated with blue light exposure^{14, 49}, which is also
368 supported by animal and laboratory studies^{6, 50}. In the present study, we could only estimate ambient
369 UVR, since no data on blue light are currently available in the Eurosun project. It is however difficult to
370 distinguish the effect of the different wavelengths, since they are naturally highly correlated. For
371 instance, in the present study, global solar radiation (a measure of solar energy including all
372 wavelengths) showed a correlation coefficient of 0.95 with UV radiation, measured in 105 geographical
373 areas.

374 The present study has several limitations. One limitation of our study could come from the
375 representativeness of the sample. The Alienor subsample tends to over-represent younger subjects
376 and high socioeconomic status, among subjects participating to the 3C Study²³. The individuals
377 included in this study may therefore be healthier and have different lifestyles, particularly concerning
378 sunlight exposure, than the general population. These differences may have affected the distribution
379 of sunlight exposure or the prevalence of eye diseases. However, subjects participating in the 3C
380 study and included in the Alienor study were not different from those who were not included for most
381 parameters of interest in our study²³. Furthermore, as described previously²³, the age-gender-
382 specific prevalence rates of AMD in the Alienor study were similar to that observed in other studies
383 performed in Europe^{51, 52} and other industrialized countries⁵³. Data collection was performed in the
384 same way in all individuals regardless of their AMD stage and photograph graders had no access to
385 sunlight exposure data. Therefore, we can assume that the error was not differential and was unlikely
386 to have biased the estimation of any of the associations of AMD with ambient solar radiation.

387 In observational studies, confounding is always a concern. We therefore adjusted for potential
388 confounders, including all known risk factors for cataract and AMD. However, we cannot totally
389 exclude residual confounding. In the present study, it is in particular striking that not only socio-
390 demographic and lifestyle variables varied with ambient UVR exposure, but also genetic background
391 (mainly CFH Y402H). While there are major differences in distribution of CFH Y402H alleles among
392 ethnic groups (with low frequency of the at-risk C allele in non-Europeans^{54, 55}), to our knowledge,
393 there are no available data on its geographical distribution within the European continent (where most

394 of the Alienor participants were born). However, large variations of allele frequencies between
395 European countries have been reported, in particular for Apolipoprotein E³³. Since AMD has a strong
396 genetic component, and there may be geographical variation in the distribution of gene
397 polymorphisms, gene polymorphisms may be confounders in the relationship of AMD with any factor
398 affected by geographical variation (including UVR exposure, but also lifestyle, diet...). This underlines
399 the importance of taking into account both environmental and genetic confounders in studies of
400 associations of eye diseases with sunlight exposure. None of the previously published studies took
401 potential genetic confounders into account. In the present study, although we took into account the
402 two major genetic risk factors for AMD (CFH Y402H and ARMS A69S), we cannot exclude that these
403 groups of subjects differ for other genetic factors related to AMD (complement factors CF1, C2, C3
404 and CFB, LIPC, CETP...³).

405 Since this study was cross-sectional, recall bias may have affected the results. However, only few
406 prospective studies are available in this field. Our study included only a small number of cases of late
407 AMD, which induced low statistical power for detecting associations with ambient solar radiation and of
408 interactions of genetic polymorphisms with UVR exposure. Finally, cataract status was determined
409 only on the basis of cataract surgery, rather than on the presence of lens opacities, which were not
410 available in our study because of lack of pupil dilation. This may have caused misclassification of
411 cataract status and therefore may have biased the estimates.

412 In conclusion, our study confirms the high risk for cataract in subjects exposed to high ambient solar
413 radiation. It also suggests a U-shaped association of early AMD with UVR exposure, with an increased
414 risk for both low- and high-exposures. This will need to be confirmed in future, possibly prospective,
415 studies. The reasons for a potentially increased risk for early AMD with low solar radiation remain
416 unclear and need to be further studied.

417

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Table 1. Light exposure-related characteristics of participants according to lifetime ambient total UVR exposure of the Alienor Study (Bordeaux, France, 2006-2008) (N=837)

	Total UVR, kJ/cm ²			p-value
	≤ 39.649 (N=209)]39.649-40.173[(N=419)	≥40.173 (N=209)	
Mean total UVR, kJ/cm ² (sd)	38.375 (1.26)	39.887 (0.08)	42.718 (2.29)	
Place of birth (n (%)) (N=837)				
South of France	66 (31.58)	392 (93.56)	102 (48.80)	P<0.0001
Central France	52 (24.88)	9 (2.15)	14 (6.70)	
Northern France	88 (42.11)	12 (2.86)	14 (6.70)	
Southern Europe	0 (0.00)	4 (0.95)	17 (8.13)	
Northern Africa	0 (0.00)	1 (0.24)	57 (27.27)	
Other	3 (1.44)	1 (0.24)	5 (2.39)	
Time spent at place of birth, years (median (range)) (N=837)				
South of France	68 (39-89)	79 (55-94)	74 (35-92)	P<0.0001
Central France	22 (2-58)	13 (1-34)	20 (4-37)	P=0.11
Northern France	23.5 (3-69)	3.5 (1-42)	12.5 (1-31)	0.001
Southern Europe		2.5 (1-6)	17 (2-31)	0.03
Northern Africa		1 (1-1)	32 (17-48)	P<0.0001
Other	3 (1-6)	3 (3-3)	18 (1-30)	0.13

Table 2. Socio-demographic, medical and genetic characteristics of participants according to lifetime ambient total UVR exposure of the Alienor Study (Bordeaux, France, 2006-2008) (N=837)

	Ambient total UVR, kJ/cm ²			p-value
	≤ 39.649 (N=209)]39.649- 40.173[(N=419)	≥40.173 (N=209)	
Age, years (mean (sd)) (N=837)	80.50 (4.42)	80.21 (4.49)	79.96 (4.44)	0.46
Male gender (n (%)) (N=837)	87 (41.63)	133 (31.74)	89 (42.58)	0.008
Education (n (%)) (N=837)				
None or primary school	48 (22.97)	130 (31.00)	54 (25.84)	P<0.0001
Secondary school	34 (16.27)	137 (32.70)	63 (30.14)	(global)
High school	25 (11.96)	39 (9.31)	23 (11.00)	
University	102 (48.80)	113 (26.97)	69 (33.01)	
Diabetes (n (%)) (N=763)	23 (12.17)	53 (13.80)	21 (11.05)	0.63
Hypertension (n (%)) (N=837)	161 (77.03)	318 (75.89)	152 (72.73)	0.56
Asthma (n (%)) (N=828)	21 (10.14)	36 (8.67)	14 (6.80)	0.47
Oral corticosteroid use (n (%)) (N=837)	3 (1.44)	2 (0.48)	4 (1.91)	0.22
CFH Y402 H (n (%)) (N=766)				
TT (low AMD risk)	79 (40.93)	188 (49.21)	86 (45.03)	0.05
TC (intermediate AMD risk)	91 (47.15)	143 (37.43)	90 (47.12)	(global)
CC (high AMD risk)	23 (11.92)	51 (13.35)	15 (7.85)	
ARMS2 A69S (n (%)) (N=703)				
GG (low AMD risk)	118 (66.67)	232 (66.48)	112 (63.28)	0.91
GT (intermediate AMD risk)	53 (29.94)	103 (29.51)	59 (33.33)	(global)
TT (high AMD risk)	6 (3.39)	14 (4.01)	6 (3.39)	

Table 3. Lifestyle and nutritional characteristics of participants according to lifetime ambient total UVR exposure of the Alienor Study (Bordeaux, France, 2006-2008) (N=837)

	Ambient total UVR, kJ/cm ²			p-value
	≤ 39.649 (N=209)]39.649- 40.173[(N=419)	≥40.173 (N=209)	
Smoking (n (%)) (N=828)				
Non smoker	111 (54.15)	296 (71.15)	138 (66.67)	0.0005
< 20 pack-years	47 (22.93)	68 (16.35)	31 (14.98)	
≥ 20 pack-years	47 (22.93)	52 (12.50)	38 (18.36)	
Physical activity (n (%)) (N=837)				
Low	29 (13.88)	83 (19.81)	31 (14.83)	0.20
Medium	112 (53.59)	226 (53.94)	108 (51.67)	
High	47 (22.49)	76 (18.14)	43 (20.57)	
No answer	21 (10.05)	34 (8.11)	27 (12.92)	
Body mass index, kg/m ² (mean (sd)) (N=829)	25.90 (4.00)	26.17 (3.97)	26.61 (3.65)	0.17
Fasting plasma measurements, mmol/l (mean (sd)) :				
Glucose (N=759)	5.16 (1.03)	5.13 (1.12)	5.13 (1.22)	0.96
Total cholesterol (N=787)	5.83 (0.91)	5.83 (1.03)	5.72 (0.92)	0.37
HDL- cholesterol (N=786)	1.60 (0.40)	1.61 (0.39)	1.57 (0.39)	0.66
LDL- cholesterol (N=785)	3.66 (0.80)	3.67 (0.91)	3.61 (0.79)	0.73
Triglycerides (N=786)	1.25 (0.61)	1.23 (0.60)	1.18 (0.48)	0.37
Dietary intake (mean (sd)):				
Total energy Kcal/day (N=797)	1835.7 (574.1)	1678.2 (513.4)	1704.5 (535.3)	0.003
Vitamin C, mg/day (N=797)	89.62 (56.98)	81.95 (60.96)	92.10 (82.66)	0.15
Vitamin E, mg/day (N=797)	6.72 (4.72)	6.30 (4.51)	6.70 (4.43)	0.45
Lutein and zeaxanthin, mg/day (N=781)	0.60 (0.92)	0.55 (0.91)	0.71 (0.90)	0.14
Omega 3 fatty acids, mg/day (N=797)	0.48 (1.20)	0.43 (1.00)	0.41 (1.16)	0.79

Table 4. Associations of cataract extraction with lifetime ambient UVR exposure in the Alienor Study (Bordeaux, France, 2006-2008) (Odds-ratios (OR) and 95 % confidence intervals (CI))

	Cataract surgery				Odds-ratio*	P
	No		Yes			
	(N=846 eyes)		(N=542 eyes)			
	N	%	N	%		
Total UV (kJ/cm ²)						
≤39.649 (n=350 eyes)	230	65.71	120	34.29	0.75 (0.51-1.12)	0.16
]39.649-40.173[(n=696 eyes)	430	61.78	266	38.22	1.00 (ref)	
≥40.173 (n=342 eyes)	186	54.39	156	45.61	1.53 (1.04-2.26)	0.03
UVA (kJ/cm ²)						
≤38.617 (n=350 eyes)	230	65.71	120	34.29	0.75 (0.51-1.12)	0.16
]38.617-39.131[(n=696 eyes)	430	61.78	266	38.22	1.00 (ref)	
≥39.131 (n=342 eyes)	186	54.39	156	45.61	1.53 (1.04-2.26)	0.03
UVB (kJ/cm ²)						
≤1.028 (n=350 eyes)	230	65.71	120	34.29	0.75 (0.51-1.10)	0.14
]1.028 -1.042[(n=696 eyes)	430	61.78	266	38.22	1.00 (ref)	
≥1.042 (n=342 eyes)	186	54.39	156	45.61	1.53 (1.04-2.26)	0.03

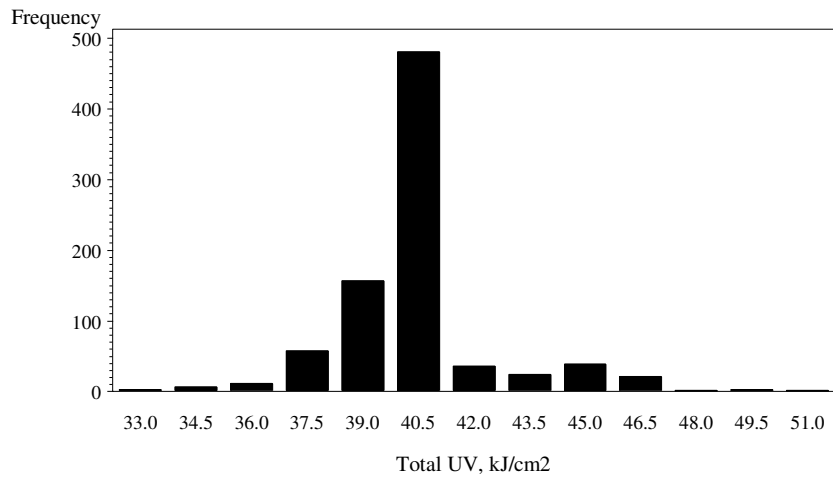
* estimated using multivariate Generalized Estimating Equations (GEE) logistic regression models, adjusted for age, gender, educational level, smoking, diabetes, oral corticosteroid, asthma, total energy intake and CFH Y402H.

Table 5. Associations of AMD with lifetime ambient UVR exposure in the Alienor Study (Bordeaux, France, 2006-2008) (Odds-ratios (OR) and 95 % confidence intervals (CI))

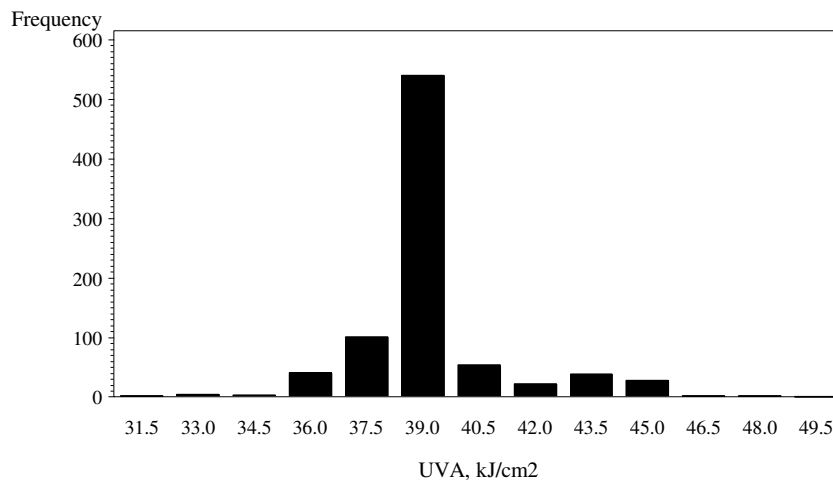
	Early AMD	P	Late AMD	P
	OR (CI)*		OR (CI)*	
	(N=238 eyes)		(N=49 eyes)	
Total UV (kJ/cm²)				
≤39.649 (n=300 eyes)	1.69 (1.06-2.69)	0.03	1.03 (0.33-3.26)	0.95
]39.649-40,173[(n=559 eyes)	1.00 (ref)		1.00 (ref)	
≥40.173 (n=295 eyes)	1.59 (1.04-2.44)	0.03	1.11 (0.36-3.36)	0.86
UVA (kJ/cm²)				
≤38.617 (n=300 eyes)	1.69 (1.06-2.69)	0.03	1.03 (0.33-3.26)	0.95
]38.617-39.131[(n=559 eyes)	1.00 (ref)		1.00 (ref)	
≥39.131 (n=295 eyes)	1.59 (1.04-2.44)	0.03	1.11 (0.36-3.36)	0.86
UVB (kJ/cm²)				
≤1.028 (n=302 eyes)	1.66 (1.04-2.64)	0.03	1.03 (0.33-3.25)	0.96
]1.028 -1.042[(n=557 eyes)	1.00 (ref)		1.00 (ref)	
≥1.042 (n=295 eyes)	1.58 (1.03-2.42)	0.04	1.11 (0.36-3.36)	0.86

* estimated using multivariate Generalized Estimating Equations (GEE) logistic regression models, age, gender, educational level, smoking, CFH Y402H and ARMS2 A69S polymorphisms, cataract extraction, dietary intake of total energy and of omega3 fatty acids.

Total UV



UVA



UVB

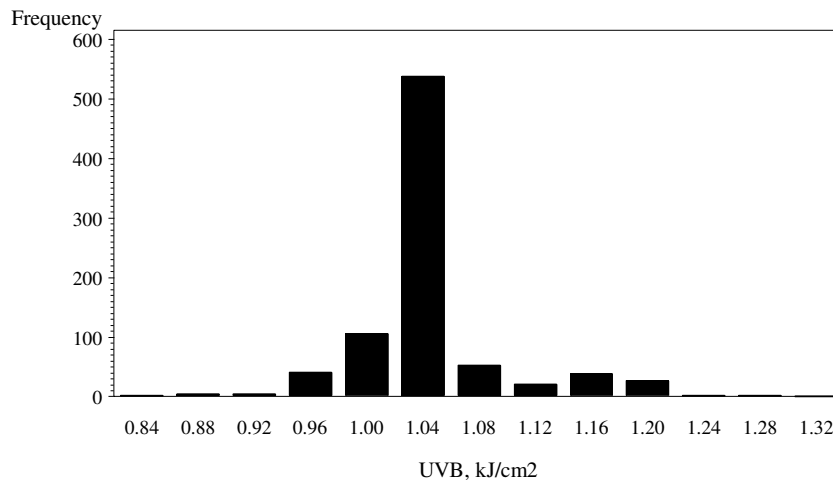


Figure 1. Distribution of lifetime ambient total UV, UVA and UVB radiations exposure in participants of the Alienor Study (Bordeaux, 2006-2008)