

BMJ Open Mortality during 6 years of follow-up in relation to visual impairment and eye disease: results from a population-based cohort study of people aged 50 years and above in Nakuru, Kenya

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ABSTRACT

Objective To estimate the association between (1) visual impairment (VI) and (2) eye disease and 6-year mortality risk within a cohort of elderly Kenyan people.

Design, setting and participants The baseline of the Nakuru Posterior Segment Eye Disease Study was formed from a population-based survey of 4318 participants aged ≥50 years, enrolled in 2007–2008. Ophthalmic and anthropometric examinations were undertaken on all participants at baseline, and a questionnaire was administered, including medical and ophthalmic history. Participants were retraced in 2013–2014 for a second examination. Vital status was recorded for all participants through information from community members. Cumulative incidence of mortality, and its relationship with baseline VI and types of eye disease was estimated. Inverse probability weighting was used to adjust for non-participation.

Primary outcome measures Cumulative incidence of mortality in relation to VI level at baseline.

Results Of the baseline sample, 2170 (50%) were re-examined at follow-up and 407 (10%) were known to have died (adjusted risk of 11.9% over 6 years). Compared to those with normal vision (visual acuity (VA) ≥6/12, risk=9.7%), the 6-year mortality risk was higher among people with VI (<6/18 to ≥6/60; risk=28.3%; risk ratio (RR) 1.75, 95% CI 1.28 to 2.40) or severe VI (SVI)/blindness (<6/60; risk=34.9%; RR 1.98, 95% CI 1.04 to 3.80). These associations remained after adjustment for non-communicable disease (NCD) risk factors (mortality: RR 1.56, 95% CI 1.14 to 2.15; SVI/blind: RR 1.46, 95% CI 0.80 to 2.68). Mortality risk was also associated with presence of diabetic retinopathy at baseline (RR 3.18, 95% CI 1.98 to 5.09), cataract (RR 1.26, 95% CI 0.95 to 1.66) and presence of both cataract and VI (RR 1.57, 95% CI 1.24 to 1.98). Mortality risk was higher among people with age-related macular degeneration at baseline (with or without VI), compared with those without (RR 1.42, 95% CI 0.91 to 2.22 and RR 1.34, 95% CI 0.99 to 1.81, respectively).

Conclusions Visual acuity was related to 6-year mortality risk in this cohort of elderly Kenyan people, potentially

Strengths and limitations of this study

- The cohort comprised a representative population-based sample in an area of ethnic, socioeconomic and educational diversity.
- There was a comprehensive assessment of ophthalmic characteristics and risk factors at baseline and follow-up.
- Data on mortality were collected through informant report, rather than from death certificates.
- There was a high loss to follow-up in this study, raising the possibility of selection bias.

because both VI and mortality are related to ageing and risk factors for NCD.

INTRODUCTION

Visual impairment (VI) is common, affecting approximately 253 million people globally.¹ It can impact on different aspects of people's lives, including reducing quality of life, and increasing poverty and depression.^{2–5} There is growing evidence from Europe, North America, Asia and Australia that VI and specific eye conditions are linked to increased risk of mortality,^{6–17} but data are lacking for low-income and middle-income countries (LMICs), particularly from sub-Saharan Africa.

There are several potential pathways by which VI may be linked with mortality. Both VI and mortality are related to ageing, and so confounding or residual confounding may explain the reported associations. There are also common underlying risk factors for both VI and mortality, such as smoking, obesity and poverty. For instance, VI due to age-related macular degeneration (AMD) is more

common among smokers,¹⁸ and smoking increases risk of mortality. An underlying disease may also cause both VI and mortality, for instance, diabetic retinopathy (DR) is related to poor control of diabetes, which also causes increased mortality. People with VI may find it more difficult to seek healthcare, due to a range of barriers,¹⁹ thereby increasing their mortality risk. Changes in the eye may be a marker of ageing or accelerated ageing,²⁰ and thereby linked to mortality. Finally, VI could exacerbate frailty, depression and functional difficulties, all linked to increased mortality.^{3 21 22}

It is important to explore whether there is an association between VI and mortality and, if there is, to identify possible pathways for this link, in order to understand how to reduce the vulnerability of people with VI to increased morbidity and mortality. Furthermore, these data may be useful to advocate for scaling up of eye health services in LMICs. The objective of this study was to investigate the association between VI and 6-year mortality risk within the Nakuru Eye Disease Cohort Study, a cohort of elderly Kenyan people.

MATERIALS AND METHODS

The methodology of the Nakuru Eye Disease Cohort Study has been reported previously,²³ and is summarised here.

Baseline study population

The baseline population-based survey was conducted in 2007/2008. A total of 100 clusters each of 50 participants were selected with a probability proportional to the size of the population across Nakuru district. Households were selected within clusters using a modified compact segment sampling method.²⁴ Eligible individuals were those aged ≥ 50 years living in the household for at least 3 months in the previous year, and multiple subjects could be included per household.

Baseline ophthalmic and general examination

All participants were invited to undergo a comprehensive ophthalmic examination at a screening clinic.²³ The objectives of the survey and the examination process were explained to those eligible in the local dialect, in the presence of a witness. A subject was examined only after informed written (or thumbprint) consent was obtained.

All participants underwent Logarithm of the Minimum Angle of Resolution (logMAR) visual acuity (VA) testing on each eye separately and corrected VA (by refraction or pinhole) when less than 20/40 Snellen equivalent. Participants had two non-stereoscopic, digital, 45° fundus photographs (one disc and one macula centred) taken per eye by an ophthalmic clinical officer. Digital images were graded for the presence of AMD and DR at an approved grading centre (Moorfields Eye Hospital

Table 1 Baseline characteristics of the Nakuru Eye Disease Cohort Study

Baseline characteristics		Deceased n=409	Followed-up or known to be alive n=3032	P value*	Unknown status n=973	P value†
Age in years, mean (SD)		71.6 (12.8)	62.4 (9.6)	<0.001	63.2 (10.6)	0.60
Sex, % (n)	Female	42 (173)	53 (1612)	<0.001	53 (516)	0.59
	Male	58 (236)	47 (1420)		47 (457)	
Tribe, % (n)	Kikuyu	69 (283)	62 (1881)	0.02	61 (596)	0.005
	Kalenjin	23 (93)	24 (736)		19 (186)	
Education, % (n)‡	None	6 (26)	9 (283)	<0.001	12 (114)	0.21
	Primary	44 (179)	32 (952)		33 (323)	
	Secondary	43 (174)	48 (1448)		44 (427)	
	Higher	7 (29)	11 (327)		11 (105)	
Residence, % (n)	Rural	74 (303)	71 (2167)	0.43	51 (498)	<0.001
	Urban	26 (106)	29 (865)		49 (475)	
SES quartile, % (n)‡	Lower	33 (136)	24 (709)	0.002	26 (247)	0.17
	Middle lower	24 (96)	26 (776)		23 (219)	
	Middle upper	24 (97)	26 (777)		23 (218)	
	Upper	19 (79)	24 (731)		29 (281)	

*P value describes the strength of evidence that each variable is associated with mortality, among those where we know the mortality status (Null hypothesis is that the odds of death are equal in each category of the variable).

†P value describes the strength of evidence that each variable is associated with mortality status being missing (ie, comparison of known vs unknown mortality status). (Null hypothesis is that odds of knowing the mortality status of an individual at follow-up are equal in each group).

‡There were 27 missing values for education, and 48 missing values for SES.

SES, socioeconomic status.

Table 2 The 6-year weighted mortality risk by level of visual impairment (VI) among the Nakuru Eye Disease Cohort Study Participants, stratified by age and gender

Weighted using inverse probability-weights	Overall		Male		Female		<60 years		≥60 years	
	N	Risk per 1000/6 years (95% CI)	N	Risk per 1000/6 years (95% CI)	N	Risk per 1000/6 years (95% CI)	N	Risk per 1000/6 years (95% CI)	N	Risk per 1000/6 years (95% CI)
Visual acuity at baseline (better eye presenting)										
All individuals	409/3441	119 (106 134)	236/1656	143 (123 165)	173/1785	98 (84 113)	86/1503	56 (4570)	323/1938	169 (151 188)
Normal (≥6/12)	280/2901	97 (84 111)	162/1378	118 (98 142)	118/1523	77 (6493)	78/1420	54 (4268)	202/1481	138 (120 159)
Near normal (<6/12 to ≥6/18)	27/170	158 (107 226)	14/84	168 (104 258)	13/86	148 (85 245)	3/27	106 (32 301)	24/143	168 (114 241)
VI (<6/18 to ≥6/60)	77/275	283 (236 336)	45/142	316 (252 388)	32/133	249 (178 338)	3/31	99 (31 271)	74/244	307 (252 368)
Severe VI (<6/60 to ≥3/60)	4/16	260 (86 566)	2/10	187 (34 603)	2/6	379 (51 875)	2/2	-	2/14	136 (27 475)
Blind (<3/60)	19/50	385 (245 548)	13/30	436 (247 646)	6/20	310 (133 570)	0/6	-	19/44	438 (279 612)
Any VI (<6/18)	100/341	297 (248 351)	60/182	328 (268 395)	40/159	262 (191 347)	5/39	137(57 297)	95/302	318 (262 380)

Reading Centre) by a senior grader, with adjudication by a clinician for confirmed cases and 5% of randomly selected images to ensure quality control. The presence of cataract was recorded by the ophthalmologist (WM) on slit-lamp examination after pupil dilation.

Detailed interviews were undertaken in the local language on demographic details, information on risk factors, socioeconomic status (SES) and full medical history.

A nurse performed and recorded measures of participants: height (Leicester Height Measure); weight (Seca 761) and three measures of blood pressure (Omron Digital Automatic Blood Pressure Monitor Model HEM907), each 10 min apart. Capillary blood was taken from all participants for random blood glucose (Accutrend GC system).

Assessment of vital status at follow-up

Follow-up was conducted from January 2013 to March 2014. A meeting was held approximately 1 week before the follow-up examination clinic for a given cluster. A list of study participants was given to the chief and a local village guide was recruited to assist location of the study participants. The village guide was someone who knew and was well known by the community (or the village chief him/herself). The advance team visited homes of baseline participants on the day prior to the examination clinic and confirmed their identity using the national identity cards and invited them to attend the examination clinic the following day. All identified participants were also asked to help locate baseline participants that had not been found.

Each local field guide was asked to classify the baseline study participant for that cluster as ‘available’, ‘died’, ‘moved away’ or ‘unknown’. A participant was defined to have died if this was verified by at least two people from among the village chief, local guide or available study participant. Those who were known to have moved away were contacted when possible to either arrange follow-up at a more suitable location for the participant or to identify if they were alive or had died in the follow-up period. Any participant for whom nobody could identify as being alive or having moved away was recorded as ‘unknown’.

Definitions and statistical analyses

All participants who had complete examinations at baseline were considered ‘at-risk’ for mortality during follow-up. Follow-up status at 6 years was categorised as: (1) deceased (confirmed dead, as described above); (2) alive (ie, re-examined at follow-up, retraced but refused or unavailable at follow-up, or moved away but known to be alive) (3) unknown (ie, not retraced at follow-up, death not verified as described above or moved away but vital status unknown).

An SES score was developed based on information collected on job, housing conditions and ownership of material goods and livestock, based on previous work in the same population.²⁵ Hypertension was defined on the

Table 3 Association between visual acuity (VA) category and 6-year mortality risk

VA category	Age/sex-adjusted RR	Age/sex, SES* adjusted RR	Age/sex, NCD risk† factor adjusted RR	Fully adjusted ‡ RR
Normal ($\geq 6/12$)	Reference	Reference	Reference	Reference
Near normal ($< 6/12$ to $\geq 6/18$)	0.92 (0.57–1.50)	0.84 (0.51–1.39)	0.87 (0.51–1.48)	0.82 (0.48–1.41)
VI ($< 6/18$ to $\geq 6/60$)	1.75 (1.28–2.40)	1.77 (1.30–2.40)	1.56 (1.13–2.16)	1.56 (1.14–2.15)
SVI/blind ($< 6/60$)	1.98 (1.04–3.80)	1.95 (1.01–3.76)	1.51 (0.82–2.77)	1.46 (0.80–2.68)
P value	0.004	0.003	0.04	0.03

*SES=SES quartile, location, ethnic group, education.

†NCD risk factor=smoking, alcohol, diabetes, hypertension, BMI.

‡Age=sex, plus all SES and NCD risk factors.

BMI, body mass index; NCD, non-communicable disease; RR, risk ratio; SES, socioeconomic status; SVI, severe visual impairment; VI, visual impairment.

basis of the average of the second and third reading, with cut-offs used of systolic blood pressure ≥ 140 mm Hg and/or diastolic blood pressure ≥ 90 mm Hg and/or self-reported hypertension medication. Diabetes was defined as (1) self-reported in the history, or (2) random glucose of ≥ 11.0 mmol/L.

Statistical analysis was performed using STATA V.14 (Stata). All analyses accounted for the cluster survey design using Taylor linearised variance estimation to calculate standard errors. Pearson χ^2 tests corrected for the survey design were used to calculate p values to assess differences between participants whose mortality status is known and those where mortality status is unknown, that is, lost to follow-up (LTFU).

An inverse probability-weighting (IPW) model²⁶ was developed, in order to allow estimation of mortality risk while accounting for those LTFU. Multivariable logistic regression was used to identify independent baseline covariates associated with LTFU. Covariates for which there was the evidence of univariable association with the outcome ($p < 0.10$ across all categories of the variable) were kept in a multivariable model (age, sex, rural/urban and mother tongue). From this final model, the probability of being followed up was estimated, based on the presence or absence of each of these baseline covariates. The inverse of this probability formed the weighting to be applied to account for those LTFU.

The final step was to remove those individuals LTFU from the cohort, so that all subsequent analysis would be performed on only those with complete outcome records, with IPW applied to account for those LTFU. A sensitivity analyses for this approach involved repeating the analyses without applying IPW (ie, standard unweighted complete case analysis), and assessing the impact on the results.

Six-year mortality risk was calculated by dividing the number of deaths identified at follow-up by the number of people at risk at baseline. The 95% CIs were estimated assuming a Poisson distribution of events. This was done for the population overall, and stratified by each covariate.

Age/sex-adjusted risk ratios (RRs) for each covariate in relation to mortality were estimated using a Poisson

regression model with robust error variance to allow for the clustered design and including IPW. Mortality status was the binary outcome and the distribution was assumed to be Poisson. These analyses were adjusted for the clustered design, as well as the use of IPW, by setting the clusters as the primary sampling unit and weighting using the inverse probability of being followed up. The model was further adjusted using a set of four SES variables only (SES quartile, location, ethnic group, education), then a set of five non-communicable disease (NCD) risk factors only (smoking, alcohol, diabetes, hypertension, body mass index (BMI)) before finally adjusting for all nine SES and NCD variables.

Patient and public involvement

Patients and the public were not directly involvement in the development of the research question and outcome measures, or the design or conduct of study. There are no plans to disseminate results directly to the study participants.

RESULTS

At baseline, 4414 participants were examined. The follow-up assessment was conducted, on average, 5.6 years (SD 0.6) after the baseline, expressed for simplicity as 6 years (meaning that there were 6 years between the baseline and follow-up wave, rather than that each participant was followed up on average for 6 years as the time of LTFU or death was not known for individuals). Of the baseline participants, 3032 were known to be alive at follow-up (69% 2170 re-examined at the follow-up plus 862 known to be alive but not re-examined), 409 (9%) were known to have died and 973 had unknown vital status (22%).

Table 1 provides the baseline characteristics of participants who had died during the follow-up, those were re-examined at follow-up and those who were LTFU. In comparison to those who had died, those who were re-examined were younger, more likely to be female, Kalenjin speakers, and had higher SES, while those of unknown

status were more likely to be of 'other' tribes and urban residence.

Table 2 shows the weighted 6-year mortality risk by level of VI. Overall, the 6-year mortality risk was 11.9% over 6 years. Risk increased with worsening levels of VI, from 9.7% (95% CI 8.4% to 11.1%) among those with normal vision to 38.5% (95% CI 24.5% to 54.8%) among those who were blind. This pattern was observed in both males and females, but was less clear in people aged <60 years given the low mortality in this group and consequent small numbers. In each subgroup, the lowest risk of mortality was among people with normal vision. Mortality risk among people with VI was higher for males than for females, and among those ≥ 60 years vs <60 years. Estimates changed little after weighting for LTFU (online supplementary table for unweighted estimates).

Compared with those with normal vision (VA >6/12, risk=9.7%), the mortality risk was significantly higher among people with VI (VA <6/18 to $\geq 6/60$; risk=28.3%; RR 1.75, 95% CI 1.28 to 2.40) or severe VI (SVI)/blindness (VA <6/60: risk=34.9%; RR 1.98, 95% CI 1.04 to 3.80) (table 3). There was a weakening of the association after adjustment for NCD risk factors or full adjustment for both SES and NCD risk factors, although the overall trends between worsening vision and increased 6-year mortality risk remained evident (VI: RR 1.56, 95% CI 1.14 to 2.15; SVI/blind: RR 1.46, 95% CI 0.80 to 2.68).

People with any VI had a higher mortality risk than those without VI (29.7% vs 9.7%; RR 1.54, 95% CI 1.22 to 1.93), and this association remained after adjustment for SES and NCD risk factors (RR 1.37, 95% CI 1.10 to 1.71) (table 4). Other risk factors associated with 6-year mortality risk after comprehensive adjustment included increasing age (oldest vs youngest age group: RR 4.68, 95% CI 3.55 to 6.18) and diabetes (RR 2.34, 95% CI 1.81 to 3.03). Being underweight was associated with an increased 6-year mortality risk (underweight vs normal: RR 1.60, 95% CI 1.24 to 2.07).

Risk of mortality was analysed by prevalence of specific eye diseases at baseline (table 5). The presence of cataract (or aphakia/pseudophakia) and any VI (ie, VA <6/18 in better eye) was associated with higher mortality risk (RR 1.57, 95% CI 1.24 to 1.98), whereas cataract alone (or aphakia/pseudophakia) was not (RR 1.26, 95% CI 0.95 to 1.66). Mortality risk was higher among people with AMD at baseline (with or without VI), compared with those without, although these associations were not statistically significant (RR 1.42, 95% CI 0.91 to 2.22 and RR 1.34, 95% CI 0.99 to 1.81, respectively). DR was associated with a threefold increased mortality risk (RR 3.18, 95% CI 1.98 to 5.09). The number of people with DR and any VI was too small to make meaningful inferences.

DISCUSSION

VI was associated with increased mortality risk during 6 years of follow-up in a cohort of elderly Kenyan people. The risk of mortality increased with worsening vision. This

association was reduced after adjustment for the presence of NCD risk factors, and to a lesser extent for SES indicators. Among eye conditions, DR was most strongly associated with mortality risk, although the number affected was small. Cataract with VI was also associated with elevated mortality, as were AMD and cataract without visual loss at baseline (although these estimates lacked precision).

Previous studies have also shown a positive relationship between VI and mortality, with evidence available from the USA,⁶⁻⁹ UK,¹⁰ Australia,¹¹ Japan,¹³ Singapore,¹⁴ 15 China¹⁶ and India.¹⁷ Others have failed to find evidence for this association, including in India,²⁷ Iceland²⁸ and Taiwan.²⁹ Data from LMICs are sparse, in particular for sub-Saharan Africa, and so comparison of our study findings to those from similar settings is not possible.

On the basis of our findings, and those in the wider literature, consideration can be given to the potential pathway for the association between VI and mortality. There was clear evidence for confounding by age, as both VI and mortality are independently related to older age. Consequently, imperfect adjustment for age may have allowed for residual confounding as a partial explanation for the association. There was little evidence for confounding by SES, although in this setting high SES was associated with greater prevalence of NCD risk factors,³⁰ and a somewhat reduced mortality risk. The presence of NCD risk factors may also act as confounders of the association of VI on mortality, since the association was attenuated after adjustment for these indicators, as found in other studies.¹⁰ Significant associations persisted, however, between VI and mortality after comprehensive multivariable adjustment in this study, as occurred in previous studies,¹²⁻¹⁴ 17 31 suggesting that residual confounding or direct effects of VI on mortality may be operational.

Exploring the relationship between different eye conditions and mortality may help to clarify whether independent biological pathways exist. DR is known to be associated with increased mortality,¹⁵ 32 as was also shown in this study. This link is unsurprising given the well-known relationship between uncontrolled diabetes with both DR and mortality. However, the relatively small number of people with DR in this population means that this link cannot be the sole driver of the VI-mortality association. Our study, as well as others, has shown cataract to be associated with increased mortality,⁶ 12 17 33 34 though this association is not always demonstrated.¹⁵ 35 Some studies have suggested that this relationship varies by cataract type.⁸ 34 36-38 It is hypothesised that the association between cataract and mortality arises as lens opacification (cataract) is an indicator of accelerated ageing.²⁰ 39 The evidence for a link between AMD and mortality is more complex; some studies show that late AMD is associated with mortality, but not early AMD.⁶ 40-42 Others found no association between AMD and mortality,¹⁵ 33 35 or only among women.⁴³

There are other potential pathways between VI and mortality not explored in this study. For example, NCD risk factors may be mediators of the effect of VI on

Table 4 Multivariable analysis of baseline covariables and 6-year mortality risk in the Nakuru Eye Disease Cohort Study

	No at risk	Deaths	Risk per 1000/6 years (95% CI)	Age/sex-adjusted risk ratio	Age/sex/socioeconomic status (SES) adjusted risk ratio	Age/sex/SES- non-communicable disease (NCD) risk factor adjusted risk ratio
Any visual impairment (<6/18)						
No	3071	307	100 (87 to 115)	Reference	Reference	Reference
Yes	341	100	297 (248 to 351)	1.54 (1.22 to 1.93)	1.55 (1.24 to 1.94)	1.37 (1.10 to 1.71)
Gender						
Male	1656	236	143 (123 to 165)	Reference	Reference	Reference
Female	1785	173	98 (84 to 113)	0.74 (0.63 to 0.87)	0.68 (0.56 to 0.83)	0.82 (0.63 to 1.06)
Age						
50–59	1503	86	56 (45 to 70)	Reference	Reference	Reference
60–69	1036	96	94 (77 to 115)	1.64 (1.24 to 2.17)	1.58 (1.19 to 2.09)	1.40 (1.07 to 1.83)
70–79	571	107	191 (161 to 224)	3.27 (2.55 to 4.20)	3.15 (2.39 to 4.15)	2.74 (2.09 to 3.60)
80+	331	120	363 (311 to 419)	6.36 (4.85 to 8.33)	5.76 (4.39 to 7.57)	4.68 (3.55 to 6.18)
SES risk factors						
Location						
Rural	2470	303	123 (108 to 140)	Reference	Reference	Reference
Urban	971	106	110 (83 to 145)	1.14 (0.90 to 1.45)	1.15 (0.90 to 1.48)	1.18 (0.91 to 1.53)
SES quartile						
Lower	845	136	164 (137 to 197)	Reference	Reference	Reference
Lower middle	872	96	110 (91 to 134)	0.75 (0.60 to 0.93)	0.72 (0.58 to 0.91)	0.75 (0.59 to 0.95)
Upper middle	874	97	110 (89 to 134)	0.86 (0.66 to 1.11)	0.82 (0.63 to 1.06)	0.84 (0.65 to 1.09)
Upper	810	79	99 (78 to 126)	0.89 (0.69 to 1.14)	0.78 (0.58 to 1.05)	0.76 (0.56 to 1.03)
Ethnic group						
Kikuyu	2164	283	131 (116 to 148)	Reference	Reference	Reference
Kalenjin	829	93	113 (91 to 139)	0.83 (0.66 to 1.03)	0.78 (0.62 to 0.99)	0.81 (0.63 to 1.04)
Other	448	33	76 (48 to 119)	0.84 (0.56 to 1.26)	0.82 (0.55 to 1.21)	0.81 (0.55 to 1.20)
Education						
No education	309	26	86 (58 to 125)	Reference	Reference	Reference
Primary	1131	179	161 (138 to 188)	0.95 (0.65 to 1.41)	0.96 (0.63 to 1.46)	0.97 (0.63 to 1.50)
Secondary	1622	174	107 (91 to 125)	0.85 (0.59 to 1.22)	0.84 (0.57, 1.24)	0.87 (0.59 to 1.28)
College/Uni	356	29	81 (52 to 124)	0.94 (0.57 to 1.55)	0.95 (0.57 to 1.55)	1.01 (0.62 to 1.64)
Risk factors for NCD						
Smoking						

Continued

Table 4 Continued

	No at risk	Deaths	Risk per 1000/6 years (95% CI)	Age/sex-adjusted risk ratio	Age/sex/socioeconomic status (SES) adjusted risk ratio	Age/sex/SES- non-communicable disease (NCD) risk factor adjusted risk ratio
Never	2396	256	107 (92 to 124)	Reference	Reference	Reference
Former	252	33	131 (93 to 180)	1.20 (0.85 to 1.69)	1.20 (0.85 to 1.71)	1.15 (0.79 to 1.66)
Current	775	120	156 (132 to 184)	1.19 (0.93 to 1.52)	1.17 (0.91 to 1.50)	1.19 (0.91 to 1.55)
Diabetes						
No	3202	354	111 (98 to 125)	Reference	Reference	Reference
Yes	216	54	248 (194 to 313)	2.16 (1.69 to 2.77)	2.20 (1.69 to 2.84)	2.34 (1.81 to 3.03)
Hypertension						
No	1670	179	108 (92 to 126)	Reference	Reference	Reference
Yes	1737	229	132 (113 to 153)	1.08 (0.90 to 1.29)	1.06 (0.88 to 1.28)	1.11 (0.92 to 1.34)
Alcohol						
Never	1335	117	87 (73 to 104)	Reference	Reference	Reference
Former	1520	221	147 (125 to 171)	1.18 (0.95 to 1.48)	1.18 (0.94 to 1.49)	1.15 (0.90 to 1.46)
Current	559	70	125 (100 to 157)	1.15 (0.86 to 1.53)	1.18 (0.87 to 1.60)	1.08 (0.77 to 1.51)
Body mass index						
Underweight	468	99	216 (169 to 271)	1.55 (1.20 to 2.00)	1.57 (1.22 to 2.02)	1.60 (1.24 to 2.07)
Normal	1697	199	117 (101 to 136)	Reference	Reference	Reference
Overweight	779	66	86 (66 to 111)	0.90 (0.69 to 1.18)	0.87 (0.66 to 1.14)	0.81 (0.62 to 1.05)
Obese	447	32	73 (52 to 100)	0.89 (0.63 to 1.25)	0.86 (0.59 to 1.24)	0.83 (0.57 to 1.20)

mortality (rather than confounders) for reasons such as lower ability to access NCD treatment, less exercise, poorer diets and so on. Consequently, the association between VI and mortality adjusted by NCD risk factors would be an underestimate of the total effect. There were also concerns about the accuracy of assessment of visual fields in this population. Consequently, it was not possible to determine the presence of glaucoma at baseline, although others have suggested a link between glaucoma and mortality.^{15 44} We also did not assess the impact of VI in accessing healthcare, although the Australian Blue mountain study showed that difficulties in walking explained some of the link between VI and mortality.¹¹ Only 18 people with cataract underwent cataract surgery during the follow-up period, so it was not possible to assess the impact on mortality.

There are several further limitations of the study, which need to be considered when interpreting the findings. There was a lack of data on date of death, and no verification from death certificates, as these are rarely available

in many African settings,⁴⁵ including Kenya. Cause of death could not be determined, and so we could not assess whether the relationship was stronger between VI and specific causes of mortality, notably cardiovascular and non-cancer causes, as demonstrated in previous studies,^{15 46–48} which would lend weight to a biological pathway for the association. The follow-up study was conducted after a period of postelection violence in the area. Consequently, there was a high LTFU in this study, raising the possibility of selection bias influencing the findings, although patterns changed little after weighting for LTFU. We did not adjust for the population sampling weights in our analysis, and so there could be concerns about the representativeness of the sample, although the selected sample had a similar demographic distribution to the general population.⁴⁹ Furthermore, the mortality rate may have been higher in this period, due to violence, and may have biased the association with VI if these deaths were disproportionately among people with VI, or among younger people (with lower prevalence of VI). Date of

Table 5 Risk of mortality during 6 years of follow-up by the presence of specific eye diseases at baseline

	No at risk	Deaths	Risk per 1000/6 years (95% CI)	Age/sex- adjusted risk ratio
Cataract present				
No	1921	142	72.5 (59.1 to 88.6)	Reference
Yes	1478	265	181.3 (161.6 to 202.8)	1.26 (0.95 to 1.66)
Cataract and visual impairment (VI) present (<6/18)				
No	3103	313	100.7 (87.5 to 115.5)	Reference
Yes	296	94	322 (267.3 to 381.8)	1.57 (1.24 to 1.98)
Age-related macular degeneration (AMD) present				
No	2270	225	99.2 (83.3 to 117.8)	Reference
Yes	319	57	183.4 (141.3 to 234.5)	1.34 (0.99 to 1.81)
AMD and VI present (<6/18)				
No	2529	265	105.5 (89.9 to 123.4)	Reference
Yes	60	17	282.7 (178.5 to 416.8)	1.42 (0.91 to 2.22)
Diabetic retinopathy (DR) present				
No	2513	264	105.7 (90.4 to 123.3)	Reference
Yes	55	18	318.4 (199.4 to 467.0)	3.18 (1.98 to 5.09)
DR and VI present (<6/18)				
No	2563	280	109.9 (94.1 to 128.0)	Reference
Yes	5	2	401.6 (37.5 to 920.4)	2.54 (0.57 to 11.36)

LTFU or death was not recorded, and so survival analysis was not possible. Another concern is that reports of local informants were used to categorise some people who had moved away as 'known to be alive', which may have created inaccuracies. The study may have been underpowered for some of the subgroup analyses, such as assessing the link between type of eye disease and mortality. We did not evaluate the association of different subtypes of AMD or cataract in relation to mortality because of small numbers. VI classification did not include loss of visual fields, and so the prevalence of functionally significant sight loss may have been underestimated. Self-reported diabetes was not confirmed (eg, from medical records). In terms of strengths, this was the first study of its kind in sub-Saharan Africa to assess the association between VI and mortality. The study participants comprised a representative population-based sample in an area of ethnic, SES and educational diversity. There was a comprehensive assessment of ophthalmic characteristics and risk factors at baseline, and every attempt was made to follow up all participants, and to record vital status.

In conclusion, VA was related to 6-year mortality risk in this cohort of elderly Kenyan people. The most likely explanation for the association is that both VI and

mortality are related to ageing and NCD risk factors. The implication is that continuity of care is needed, as people with VI require linkages to preventative and treatment services. Furthermore, we must advocate for the scale-up of eye care services in Kenya, as VI is linked to premature mortality.

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