Association of reduced retinal arteriolar tortuosity with depression in older participants from the Northern Ireland Cohort for the Longitudinal Study of Ageing

O'Neill, R. A., Maxwell, A. P., Kee, F., Young, I., Hogg, R. E., Cruise, S., McGuinness, B., & McKay, G. J. (2021). Association of reduced retinal arteriolar tortuosity with depression in older participants from the Northern Ireland Cohort for the Longitudinal Study of Ageing. *BMC Geriatrics*, 21(1), 62. https://doi.org/10.1186/s12877-021-02009-z

**Published in:**
BMC Geriatrics

**Document Version:**
Publisher's PDF, also known as Version of record

**Queen's University Belfast - Research Portal:**
Link to publication record in Queen's University Belfast Research Portal

**Publisher rights**
Copyright 2021 the authors.
This is an open access article published under a Creative Commons Attribution License (https://creativecommons.org/licenses/by/4.0/), which permits unrestricted use, distribution and reproduction in any medium, provided the author and source are cited.

**General rights**
Copyright for the publications made accessible via the Queen's University Belfast Research Portal is retained by the author(s) and / or other copyright owners and it is a condition of accessing these publications that users recognise and abide by the legal requirements associated with these rights.

**Take down policy**
The Research Portal is Queen's institutional repository that provides access to Queen's research output. Every effort has been made to ensure that content in the Research Portal does not infringe any person's rights, or applicable UK laws. If you discover content in the Research Portal that you believe breaches copyright or violates any law, please contact openaccess@qub.ac.uk.

Download date: 06. Jan. 2022
Association of reduced retinal arteriolar tortuosity with depression in older participants from the Northern Ireland Cohort for the Longitudinal Study of Ageing

R. A. O’Neill, A. P. Maxwell, F. Kee, I. Young, R. E. Hogg, S. Cruise, B. McGuinness and G. J. McKay *

Abstract

Introduction: The retina shares similar anatomical and physiological features with the brain and subtle variations in retinal microvascular parameters (RMPs) may reflect similar vascular variation in the brain. The aim of this study was to assess associations between RMPs and measures of depression in the Northern Ireland Cohort for the Longitudinal Study of Ageing.

Methods: RMPs (arteriolar and venular caliber, fractal dimension and tortuosity) were measured from optic disc centred fundus images using semi-automated software. Depression was characterised by the Centre for Epidemiologic Studies Depression Scale (CES-D) in the absence of mild cognitive impairment or use of anti-depressive medications. Associations between depression and RMPs were assessed by regression analyses with adjustment for potential confounders.

Results: Data were available for 1376 participants of which 113 (8.2%) and 1263 (91.8%) were classified with and without depression. Participants had a mean age of 62.0 ± 8.4 yrs., 52% were female, and 8% were smokers. Individuals with depression had a higher CES-D score than those without (22.0 ± 6.2 versus 4.4 ± 3.9). Lower values of arteriolar tortuosity were significantly associated with depression, before and after adjustment for potential confounders (odds ratio = 0.79; 95% confidence intervals: 0.65, 0.96; \( P = 0.02 \)).

Conclusion: Decreased retinal arteriolar tortuosity, a measure of the complexity of the retinal microvasculature was associated with depression in older adults independent of potential confounding factors. Retinal measures may offer opportunistic assessment of microvascular health associated with outcomes of depression.

Keywords: Retinal microvascular parameters, Depression score, Centre for epidemiological studies- depression, CES-D, NICOLA

* Correspondence: g.j.mckay@qub.ac.uk
Centre for Public Health, Queens University Belfast, Belfast, Northern Ireland

© The Author(s). 2021 Open Access This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons licence, and indicate if changes were made. The images or other third party material in this article are included in the article’s Creative Commons licence, unless indicated otherwise in a credit line to the material. If material is not included in the article’s Creative Commons licence and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this licence, visit http://creativecommons.org/licenses/by/4.0/. The Creative Commons Public Domain Dedication waiver (http://creativecommons.org/publicdomain/zero/1.0/) applies to the data made available in this article, unless otherwise stated in a credit line to the data.
Introduction
Depression is a serious, common mental health condition that incorporates feelings of sadness or hopelessness necessitating clinical intervention. Depression prevalence increases with age and is associated with higher levels of morbidity, suicide, self-neglect and reduced physical, cognitive and social functioning [1–4]. As lifespans increase, identification of those at greater risk of age-related conditions, including depression, becomes increasingly important to ensure appropriate provision of treatment [5, 6].

The most rapidly expanding age demographic in Northern Ireland (NI) are those aged 50 years (yrs) and above. The NI population provides a unique opportunity for evaluation of depression in a setting historically shaped by conflict and an undertone of sectarian violence [7, 8]. ‘The Troubles’ was a period of conflict that spanned from 1969 to 1994 and resulted in increased psychological morbidity and mental health problems among individuals exposed to higher levels of conflict [9–11]. During this time, a five-fold increase in antidepressant prescriptions was reported in the NI population (1989–2000), especially among those aged > 55 yrs. who were three times more likely to be prescribed antidepressant medication than the 15–24 yrs. demographic [9], highlighting the increasing burden of depression among the local older population.

The retinal microvasculature shares similar anatomical and physiological features with other end organs including the brain, heart and kidneys, and subtle variations in retinal microvascular parameters (RMPs) may reflect similar variation within the cerebral, renal and coronary circulation [12–14]. Recent advances in retinal fundus imaging provide a unique and non-invasive assessment of the microvasculature currently not possible elsewhere in the body [15–18], enabling opportunistic evaluation of vascular disease and the wider systemic circulation [13–18]. Variation in RMPs have been previously reported in association with cognitive impairment, depression and cerebrovascular disease [19–22] although the nature of associations between RMPs and depression have been inconsistent [23–27]. A ‘vascular depression’ hypothesis has previously implicated cerebrovascular disease in the predisposition, precipitation, and perpetuation of geriatric depressive syndromes [28]. As such, the aim of this study was to assess associations between RMPs and measures of depression in the Northern Ireland Cohort for the Longitudinal Study of Ageing (NICOLA).

Methods
Participants
NICOLA is a longitudinal cohort study consisting of 8468 adults aged 50 yrs. and over, located in Northern Ireland (those resident in care homes or other residential institutions at baseline were excluded from the study) [7]. The study included a computer-aided personal interview (CAPI), a self-completion questionnaire (https://static-content.springer.com/esm/art%3A10.1186%2Fs12882-020-02031-0/MediaObjects/12882_2020_2031_MOESM2_ESM.pdf) and approximately 45% completed a health assessment. The CAPI was extensive in scope and included assessment of demographic, social and health-related factors, and was conducted at individual home appointment between December 2013 and March 2016. Measures of cardiovascular, physical, cognitive and visual function were determined and biological samples collected, including visual health with retinal fundus photography. Written informed consent was obtained prior to participation following ethical approval from the School of Medicine, Dentistry and Biomedical Sciences Ethics Committee, Queen’s University Belfast (SREC 12/23) and in accordance with the Helsinki Declaration.

Measurement and classification of depression and co-variates
Participants undertook the Centre for Epidemiologic Studies Depression Scale questionnaire (CES-D) which consisted of 20 questions scored from 0 to 3, to assess depressive symptoms with summative scores ranging between 0 and 60 [29]. Cognitive function was assessed using the 30 point Montreal Cognitive Assessment (MoCA) in addition to questions on subjective cognitive decline (SCD) and difficulties associated with basic activities of daily living (ADL), such as dressing, walking, bathing or showering, eating, getting in or out of bed, and using the toilet. Use of anti-depressive medications was defined as use of psycholeptic drugs (Anatomical Therapeutic Chemical [ATC] classification N05) and/or psychoanaleptic drugs (ATC classification N06). Mean arterial blood pressure (MABP) was an average of two individual systolic (SBP) and diastolic blood pressure (DBP) measurements (2/3 DBP + 1/3 SBP). Diabetes was characterised by HbA1c ≥48 mmol/mol, use of diabetic medications or self-reported diabetes. Cardiovascular disease (CVD) was characterised by self-reported history of angina, heart attack, congestive heart failure or stroke. Participant height was measured to the nearest centimetre using a seca 240 wall mounted measuring rod and weight was measured in kilograms using seca electronic floor scales. High and low-density lipoprotein (HDL and LDL) was measured from individual participant blood samples. Educational attainment was dichotomised as primary schooling and below or secondary level and above (including university education). Smoking status was classified as current smokers versus non-smokers. Alcohol consumption was categorised as: non-drinker, light drinker (0–7 units per week), moderate drinker (7–14 units per week).
and heavy drinker (>14 units per week). Exclusion criteria included those individuals who failed to complete the CES-D questionnaire, were classified with mild cognitive impairment (MCI; defined as MoCA test score ≤26 and SCD, in the absence of depression or problems with ADL activities), were taking anti-depressive medication or had retinal images or insufficient quality (Supplementary Figure 1). Depression was defined as a CES-D score ≥16 in the absence of MCI or anti-depressive medication use.

**Measurement of retinal microvascular parameters**

Retinal photography was performed through the dilated pupil using a Canon CX-1 Digital Fundus Camera (Canon USA, Melville, NY, USA), following dilatation from a single drop of 1% tropicamide. RMPs (central retinal arteriolar equivalent [CRAE], central retinal venular equivalents [CRVE], arteriolar to venular ratio [AVR], fractal dimension and tortuosity) were measured from optic disc centred fundus images and analysed using the semi-automated software Vessel Assessment and Measurement Platform for Images of the Retina (VAMPIRE; VAMPIRE group, University of Dundee, Dundee, Scotland, Version 3.1, Fig. 1), by a trained grader blinded to participant data [30, 31]. Analysis was undertaken on left eye images except when unavailable or of insufficient quality, in which case the right eye image was used. A paired samples t-test was used to compare a sub-sample of left and right eye measurements from 75 participants. Intraclass correlation coefficients (ICCs) were calculated to assess intergrader reliability with mean values of 0.87 (CRAE) and 0.91 (CRVE).

**Statistical analysis**

Analyses were performed using Statistical Package for Social Sciences (Version 24.0. Armonk, NY: IBM Corp). Population characteristics were described using frequencies and percentages for categorical variables or mean and standard deviation (SD) for continuous variables. RMPs were converted into standardised Z-scores, before inclusion in regression models (a standard deviation [SD] increase from the mean). Independent samples t-tests and chi-squared tests were used to compare the distribution of demographic factors and clinical variables between participants according to depression classification. Logistic regression was used to evaluate associations between RMPs and depression status as a binary trait. Minimally adjusted models included age and sex (model 1); with additional adjustment for BMI (model 2) and educational attainment, alcohol consumption, smoking status, CVD, MABP, triglycerides, diabetes, body mass index (BMI) high and low-density lipoprotein (HDL and LDL) levels (model 3). In a sensitivity analysis,
participants with diabetes were excluded. $P < 0.05$ was considered statistically significant.

**Results**

Data were available for 1376 participants of which 113 (8.2%) and 1263 (91.8%) participants were classified with and without depression (Table 1). Participants had a mean age of 62.0 ± 8.4 yrs. and 52% were female, with 8% categorised as smokers (Table 1). The mean CES-D score for all participants was 5.9 ± 6.4. MABP was 98.2 ± 12.6 mmHg, and 23 and 6% of participants were characterised with diabetes and history of CVD respectively. As expected, individuals with depression had a higher CES-D test score than those without (22.0 ± 6.2 versus 4.4 ± 3.9, respectively). Those classified as free from depression had a higher mean age (62.2 ± 8.4 yrs. versus 59.3 ± 8.1 yrs), a lower percentage were female (52% versus 59%) and they had a lower mean BMI (28.3 ± 4.7 kg/m² versus 29.6 ± 5.4 kg/m²) compared to those with depression. Participants with depression were more likely to smoke (12% versus 8%), and a higher proportion had diabetes (29% versus 23%).

Left and right eye CRAE and CRVE comparisons from 75 participants were not significantly different ($P_{\text{CRAE}} = 0.08; P_{\text{CRVE}} = 0.89$). Only mean arteriolar tortuosity was significantly lower in those with depression in unadjusted comparisons (0.085 versus 0.113; $P = 0.03$; Table 2). Decreased retinal arteriolar tortuosity was associated with depression before and after adjustment for potential confounding variables across all regression models (unadjusted: odds ratio [OR] = 0.81; 95% confidence interval [CI]: 0.67, 0.98; $P = 0.03$; minimally adjusted: OR = 0.81; 95% CI: 0.67, 0.98; $P = 0.03$; and fully adjusted: OR = 0.79; 95% CI: 0.65, 0.96; $P = 0.02$; Table 2 and Table 3). No additional significant associations between RMPs and depression were detected (Table 3). In a sensitivity analysis that excluded the 317 participants with diabetes, the effect size of the

| Table 1 | Participant summary characteristics |
|---------|-----------------------------------|
| **Participant characteristics** | **All (n = 1376)** | **No Depression (n = 1263)** | **Depression (n = 113)** | **P-Value** |
| Mean age (years, SD) | 62.0 ± 8.4 | 62.2 ± 8.4 | 59.3 ± 8.1 | $< 0.01$ |
| Female, n (%) | 717 (52.1) | 650 (51.5) | 67 (59.3) | 0.11 |
| Smoking status, yes n (%) | 115 (8.4) | 101 (8.0) | 14 (12.4) | 0.11 |
| Alcohol consumption, non-drinker, n (%) | 280 (20.3) | 259 (20.5) | 21 (18.6) | 0.26 |
| Education, secondary and above, n (%) | 1228 (89.2) | 1124 (89.0) | 104 (92.0) | 0.32 |
| Diabetes, yes n (%) | 317 (23.0) | 284 (22.5) | 33 (29.2) | 0.10 |
| Mean BMI (kg/m², SD) | 28.4 ± 4.8 | 28.3 ± 4.7 | 29.6 ± 5.4 | $< 0.01$ |
| Mean arterial blood pressure (mm Hg, SD) | 98.2 ± 12.6 | 98.3 ± 12.6 | 96.8 ± 12.8 | 0.24 |
| Cardiovascular disease, no n (%) | 1292 (93.9) | 1187 (94.0) | 105 (92.9) | 0.65 |
| Mean triglyceride (mmol/L, SD) | 1.6 ± 0.9 | 1.6 ± 0.9 | 1.6 ± 0.8 | 0.50 |
| Mean HDL cholesterol (mmol/L, SD) | 1.6 ± 0.5 | 1.6 ± 0.5 | 1.6 ± 0.4 | 0.88 |
| Mean LDL cholesterol (mmol/L, SD) | 3.5 ± 1.1 | 3.4 ± 1.1 | 3.5 ± 1.1 | 0.56 |
| Mean CES-D score (SD) | 5.9 ± 6.4 | 4.4 ± 3.9 | 22.0 ± 6.2 | $< 0.01$ |

Values are n (%) for categorical variables and mean ± SD for continuous variables. $P$ values were calculated by independent samples $t$ and chi squared tests. $P < 0.05$ is considered statistically significant. Abbreviations: BMI body mass index, HDL high-density lipoprotein, LDL low-density lipoprotein, SD standard deviation, CES-D The Centre for Epidemiologic Studies Depression Scale

| Table 2 | Summary of participant retinal microvascular parameters |
|---------|-------------------------------------------------------|
| **Retinal microvascular parameters** | **All (n = 1376)** | **No Depression (n = 1263)** | **Depression (n = 113)** | **P-Value** |
| Mean CRAE (PX, SD) | 29.613 ± 2.186 | 29.636 ± 2.190 | 29.354 ± 2.136 | 0.19 |
| Mean CRVE (PX, SD) | 40.849 ± 3.314 | 40.857 ± 3.298 | 40.757 ± 3.501 | 0.76 |
| Mean AVR (SD) | 0.728 ± 0.062 | 0.728 ± 0.061 | 0.724 ± 0.066 | 0.47 |
| Mean fractal dimension arteriolar (SD) | 1.557 ± 0.052 | 1.557 ± 0.053 | 1.563 ± 0.046 | 0.18 |
| Mean fractal dimension venular (SD) | 1.539 ± 0.051 | 1.539 ± 0.051 | 1.540 ± 0.047 | 0.89 |
| Mean tortuosity arteriolar (SD) | 0.111 ± 0.149 | 0.113 ± 0.153 | 0.085 ± 0.090 | 0.03 |
| Mean tortuosity venular (SD) | 0.068 ± 0.109 | 0.069 ± 0.111 | 0.063 ± 0.075 | 0.38 |

Values are n (%) for categorical variables and mean ± SD for continuous variables. $P$ values were calculated by independent samples $t$ tests. Abbreviations: CRAE central retinal arteriolar equivalent, CRVE central retinal venular equivalent, AVR retinal arteriolar/venular ratio, SD standard deviation, PX pixels. $P < 0.05$ was considered statistically significant.
association observed between arteriolar tortuosity and depression was slightly attenuated and no longer significant given the reduced sample size (Supplementary Table 2).

Discussion
We detected significantly lower levels of retinal arteriolar tortuosity, a complexity measure of the twisting and turning of the retinal microvasculature [32], associated with depression independent of potential confounders in older adults. Depression is a condition that affects a significant number of older people, many with underlying chronic illness or cognitive impairment, which leads to disruption of daily life and increased levels of morbidity and mortality. Age and disease related processes, including arteriosclerosis and inflammatory, endocrine, and immune changes, have been associated with the disease processes that characterises depression, especially in those of advanced age [33].

Alexopoulos and colleagues proposed a ‘vascular depression’ hypothesis implicating cerebrovascular disease in the predisposition, precipitation, and perpetuation of geriatric depressive syndromes [28], which provided a rationale for the investigation of associations between RMPs and depression [34]. Previous studies have reported associations between severe and transitory depression and vascular endothelial dysfunction [35–38] but to our knowledge, the current study is the first to report associations between decreased arteriolar tortuosity and later-life depression. Endothelial cells also play a key role in regulating retinal microvascular blood flow and angiogenesis and our findings may implicate microvascular endothelial dysfunction and reduced retinal arteriolar tortuosity and in older people with depression independent of potential confounding factors. Indeed, a recent systematic-review and meta-analyses reported associations between peripheral and cerebral forms of microvascular dysfunction with an increased odds of incident late-life depression supporting the hypothesis that microvascular dysfunction is causally linked to depression and a potential target for the prevention and treatment of symptoms [39]. The meta-analysis by van Agtmaal and colleagues evaluated several outcome measures including retinal vessel calibre [23, 40], detecting no association with depression, in line with the findings from our study.

Technological advances in non-invasive retinal image acquisition and analysis have enabled more accurate quantification of microvascular health [17]. A small number of studies investigating associations between RMPs and depression have reported inconsistent findings [23–27]. Furthermore, these studies were largely limited to the assessment of retinal vessel calibre only and were unable to consider other RMPs such as fractal dimension or tortuosity, coefficients that reflect the status of microvascular health. Sun and colleagues reported no significant associations between retinal microvascular calibre and depression similarly characterised using the CES-D questionnaire in 2420 individuals ≥65 yrs. old in the population-based Cardiovascular Health Study [26]. The Rotterdam population-based study also characterised depression using either the CES-D questionnaire or the Hospital Anxiety and Depression Scale, and reported no significant associations between retinal microvascular calibre and late-life depression in 3605 participants ≥55 yrs. [23]. Kim and colleagues identified retinal arteriolar narrowing in individuals characterised with depression using the CES-D questionnaire in a cross-sectional study of 1744 older adults (mean age of 78 yrs) from the Cardiovascular Health Study [27]. Other forms of depression and age categories have also been considered. Li and colleagues measured symptoms of antenatal anxiety and depression using the State-Trait Anxiety Inventory and Edinburgh Postnatal Depression Scale reporting associations between wider retinal arterioles and depression in 952 pregnant women [25]. Wider

### Table 3 Logistic regression analysis of retinal microvascular parameters and depression

| Retinal parameter | OR 95% CI P-Value | OR 95% CI P-Value | OR 95% CI P-Value |
|-------------------|------------------|------------------|------------------|
| CRAE (PX)         | 0.88 0.72, 1.08 0.21 | 0.89 0.73, 1.09 0.27 | 0.86 0.70, 1.06 0.17 |
| CRVE (PX)         | 0.99 0.82, 1.20 0.92 | 0.98 0.81, 1.19 0.85 | 0.97 0.79, 1.18 0.75 |
| AVR               | 0.92 0.75, 1.12 0.40 | 0.94 0.77, 1.15 0.52 | 0.92 0.75, 1.13 0.45 |
| Fractal dimension arteriolar | 1.12 0.91, 1.37 0.28 | 1.13 0.92, 1.38 0.24 | 1.12 0.91, 1.37 0.30 |
| Fractal dimension venular | 1.00 0.83, 1.20 0.97 | 1.00 0.82, 1.20 0.96 | 0.99 0.82, 1.19 0.89 |
| abTortuosity arteriolar | 0.81 0.67, 0.98 0.03 | 0.80 0.66, 0.97 0.02 | 0.79 0.65, 0.96 0.02 |
| abTortuosity venular | 0.93 0.76, 1.13 0.45 | 0.90 0.74, 1.11 0.33 | 0.91 0.74, 1.12 0.36 |

Abbreviations: CRAE central retinal arteriolar equivalent, CRVE central retinal venular equivalent, AVR retinal arteriolar/venular ratio, CI confidence interval, OR odds ratio, PX pixels. aRMPs were transformed into standardised Z-scores (based on a SD increase) before inclusion in regression models. bTortuosity values were skewed and therefore log-transformed before inclusion in regression models. Model 1 was adjusted for age (yrs) and sex; model 2 was adjusted for model 1 covariates plus BMI; model 3 was adjusted for model 2 covariates plus alcohol consumption, smoking status, educational attainment, history of cardiovascular disease, triglycerides, diabetes, mean arterial blood pressure, body mass index, high and low-density lipoprotein levels.
Retinal arterioles were also reported by Meier and colleagues in 865 adolescents and young adults from the Brisbane Longitudinal Twin Study and the Twin Eye Study with symptoms of anxiety characterised by the Somatic and Psychological Health Report questionnaire, with the authors suggesting this may implicate microvascular variation as an early mechanistic factor in depression disease aetiology [24], in contrast to arteriolar narrowing described by Kim and colleagues in older adults [24, 27].

Our study had several limitations. Firstly, the study population consisted of Caucasian participants, aged > 50 yrs. and may represent the ‘worried-well’, limiting the generalisability of our results. Secondly, although NICOLA is a longitudinal study, only baseline data were available for this cross-sectional analysis precluding assumptions regarding causality. Thirdly, although adjustment for a large number of potential confounders was made, the possibility of residual confounding remains. Additionally, despite adjusting for cardiometabolic, atherosclerotic and diabetic risk factors, we cannot fully discount their potential influence on our findings, given previously reported associations [41–45].

Despite the limitations, our study had several strengths. Similar to previous investigations of RMPs and depression [23–27], optic disc centred fundus images provided a more accurate quantification of RMPs compared to macula centred images, which generally limit assessment to the temporal retinal arcades. Although our analysis was cross-sectional in design, it provides a rationale for further evaluation of RMPs in future waves of NICOLA data. If clinical value is associated with identifying those individuals at increased risk of depression by using RMPs, earlier identification of older individuals at increased risk of adverse vascular events associated with depression may be possible. In this study, retinal images from the left eye were analysed except when unavailable or of insufficient quality, in which case the right eye image was used. This was unlikely to have limited the study conclusions, as comparable investigations have previously reported high correlations between RMPs from right and left eye comparisons [46–48], similar to the comparisons undertaken in a subset of NICOLA participants.

Furthermore, the well-characterised population-based study design added validity to our findings. Finally, the CES-D questionnaire to classify depression is well accepted with previously reported sensitivity and specificity values of 0.87 and 0.70, respectively [49] and the exclusion of participants using anti-depressive medications or with MCI improved the characterisation of depression, given those with depressive symptoms are more likely to score poorly on tests of cognitive function [50].

In conclusion, we report decreased retinal arteriolar tortuosity in association with depression in an older population independent of potential confounding factors. These retinal measures may provide non-invasive assessment of microvascular complications associated with depression.

Supplementary Information
The online version contains supplementary material available at https://doi.org/10.1186/s12877-021-02009-z.

Additional file 1.

Abbreviations
NICOLA: The Northern Ireland Cohort for the Longitudinal study of Ageing; BMI: Body mass index; HDL: High-density lipoprotein; LDL: Low-density lipoprotein; CRAE: Central retinal arteriolar equivalent; CRVE: Central retinal venular equivalent; AVR: Retinal arteriolar/venular ratio; SD: Standard deviation; PX: Pixels; SBP: Systolic blood pressure; DBP: Diastolic blood pressure; CI: Confidence intervals; β: beta value; VAMPER: Vessel Assessment and Measurement Platform for Images of the Retina; MABP: Mean arteriolar blood pressure; RMPs: Retinal microvascular parameters; Yrs: Years; CVD: Cardiovascular disease; CAP: Computer-aided personal interview; NI: Northern Ireland; CES-D: The Centre for Epidemiologic Studies Depression Scale; MCI: Mild cognitive impairment

Acknowledgments
We are grateful to all the participants of the NICOLA Study, and the whole NICOLA team, which includes nursing staff, research scientists, clerical staff, computer and laboratory technicians, managers and receptionists. The Atlantic Philanthropies, the Economic and Social Research Council, the UKCRC Centre of Excellence for Public Health Northern Ireland, the Centre for Ageing Research and Development in Ireland, the Office of the First Minister and Deputy First Minister, the Health and Social Care Research and Development Division of the Public Health Agency, the Wellcome Trust/Wolfson Foundation and Queen’s University Belfast provide core financial support for NICOLA. The authors alone are responsible for the interpretation of the data and any views or opinions presented are solely those of the authors and do not necessarily represent those of the NICOLA study team. Rachael O’Neill was supported by a PhD studentship from the Department for Economy, Northern Ireland. This work was supported by the following funders who provide core financial support for the NICOLA Study: the Atlantic Philanthropies, the Economic and Social Research Council, the UKCRC Centre of Excellence for Public Health Northern Ireland; the Centre for Ageing Research and Development in Ireland, the Office of the First Minister and Deputy First Minister, the Health and Social Care Research and Development Division of the Public Health Agency; the Wellcome Trust/Wolfson Foundation and Queen’s University Belfast.

Authors’ contributions
RON undertook retinal measurements, data analysis and was a major contributor to the writing of the manuscript. GMK and APM proposed the research hypothesis and were major contributors to the writing of the manuscript. FK and IY conceived and designed the NICOLA. BMG acquired and interpreted NICOLA health assessment data. RH acquired and interpreted NICOLA ophthalmic data. SC acquired the NICOLA CAPI questionnaire data. All authors revised the manuscript critically for important intellectual content; and agreed to be accountable for all aspects of the work in ensuring that questions related to the accuracy or integrity of any part of the work were investigated and resolved. Finally all authors read and approved the final manuscript before submission.

Funding
Rachael O’Neill was supported by a PhD studentship from the Department for Economy, Northern Ireland.

This work was supported by the following funders who provide core financial support for the NICOLA Study: the Atlantic Philanthropies; the Economic and Social Research Council; the UKCRC Centre of Excellence for
Public Health Northern Ireland; the Centre for Ageing Research and Development in Ireland; the Office of the First Minister and Deputy First Minister; the Health and Social Care Research and Development Division of the Public Health Agency; the Wellcome Trust/Wolfson Foundation; and Queen’s University Belfast.

Availability of data and materials
The data that support the findings of this study are available from the Northern Ireland Cohort for the Longitudinal Study of Ageing but restrictions apply to the availability of this data, which was used under license for the current study, and so is not publicly available. Data may however be available from the corresponding authors upon reasonable request and provided there is permission from NICOLA.

Ethics approval and consent to participate
Written informed consent was obtained from participants prior to taking part following ethical approval from the Queen’s University Belfast School Research Ethics Committee in compliance with the Declaration of Helsinki (SREC 12/23).

Consent for publication
Not applicable.

Competing interests
The authors declare that they have no competing interests.

Received: 14 October 2020 Accepted: 4 January 2021
Published online: 15 January 2021

References
1. Fiske A, Loebach-Wetherell J, Gatz M. Depression in older adults. Ann Rev Clin Psychol. 2009;5:363–89.
2. Gustafson Y, Nogueira V, O’Dwyer S, et al. Depression in old age in Austria, Ireland, Portugal and Sweden. Eur Geriatr Med. 2013;4(3):202–8.
3. Luppia M, Sikorski C, Luck T, et al. Age- and gender-specific prevalence of depression in latest-life - systematic review and meta-analysis. J Affect Disord. 2012;136(3):212–21.
4. Frankish H, Boyce N, Horton R. Mental health for all: a global goal. Lancet. 2013;382(9901):1493–4.
5. Harada CN, Natelson L, M. C., et al. Normal cognition and aging. Clin Geriatr Med. 2013;29(4):737–52.
6. Orlowe NV, Coletta C, Van Asten F, et al. Age-related changes of the retinal microvasculature. PLoS One. 2019;14(5):e0215916.
7. Mulholland C, Boyle C, Shannon C, et al. Exposure to “the troubles” in Northern Ireland influences the clinical presentation of schizophrenia. Schizop Res. 2008;108(1–2):278–82.
8. Kelly CB, Ansari T, Rafferty T, Stevenson M. Antidepressant prescribing and suicide rate in Northern Ireland. Eur Psychiatry. 2003;18(7):325–8.
9. O’Reilly D, Stevenson M. Mental health in Northern Ireland: have “the troubles” made it worse? J Epidemiol Community Health. 2003;57(7):488–92.
10. O’Neill S, Corry CV, Murphy S, et al. Characteristics of deaths by suicide in Northern Ireland from 2005 to 2011 and use of health services prior to death. J Affect Disord. 2014;168:466–71.
11. Hart NJ, Koronyo Y, Black KL, et al. Occular indicators of Alzheimer’s: exploring disease in the retina. Acta Neuropathol. 2016;132(6):767–87.
12. Kolb H. Simple Anatomy of the Retina. 2005;1 [Updated 2012 Jan 31]. In: Kolb H, Fernandez E, Nelson R, editors. Webvision: The Organization of the Retina and Visual System [Internet]. Salt Lake City (UT). University of Utah Health Sciences Center; 1995. Available from: https://www.ncbi.nlm.nih.gov/books/NBK1533/.
13. Muhammed LA. Localizing optical disc in retinal image automatically with entropy based algorithm. Int J Biomed Imaging. 2018;2016:1–7.
14. MacGillivray TJ, Trucco E, Cameron JR, et al. Retinal imaging as a source of biomarkers for diagnosis, characterization and prognosis of chronic illness or long-term conditions. Br J Radiol. 2014;87(1040):20120832.
15. MacGillivray TJ, Ong YT, Cheung CY, et al. Retinal vascular caliber measurements: clinical significance, current knowledge and future perspectives. Ophthalmol. 2013;120(3):325–36.
16. Ege BM, Heijlens OK, Larsen DV, et al. The relationship between age and colour content in fundus images. Acta Ophthalmol Scand. 2002;80(5):485–9.
17. Hanseen H, Minghetti A, Faude O, et al. Effects of different endurance exercise modalities on retinal vessel diameters in unipolar depression. Microvasc Res. 2018;120:111–6.
18. Cheung CYL, Ong S, Iram MK, et al. Retinal vascular fractal dimension is associated with cognitive dysfunction. J Stroke Cerebrovasc Dis. 2014;23(1):43–50.
19. McGrory S, Ballerini L, Okeley JA, et al. Retinal microvascular features and cognitive change in the Lothian-birth cohort 1936. Alzheimers Dement (Amst). 2019;11:500–9.
20. Williams MA, McGovan AJ, Cardwell CR, et al. Retinal microvascular network attenuation in Alzheimer’s disease. Alzheimers Dement (Amst). 2015;1(2):229–35.
21. Iram-MK, Lujendijk HI, Hoffman A, et al. Retinal vascular calibers and risk of late-life depression: the Rotterdam study. Am J Geriatr Psychiat. 2010;18(6):452–5.
22. Meier MH, Gillespie NA, Hansell NK, et al. Associations between depression and anxiety symptoms with retinal vessel caliber in adolescents and Young adults. Psychosom Med. 2014;76(9):732–8.
23. Li LJ, Iram MK, Broekman L, et al. Antenatal mental health and retinal vascular caliber in pregnant women. Trans Vis Sci Techn. 2013;2(2).
24. Sun A, Tikelis K, Klein R, et al. Are microvascular abnormalities in the retina associated with depression symptoms? The cardiovascular health study. Am J Geriatr Psychiat. 2007;15(4):335–43.
25. Kim DH, Newman AB, Hajjar I, et al. Retinal microvascular signs and functional loss in older persons: the cardiovascular health study. Stroke. 2011;42(6):1589–95.
26. Alexopoulos GS, Meyers BS, Young RC, et al. ‘Vascular depression’ hypothesis. Arch Gen Psychiatry. 1997;54(10):915–22.
27. Radloff LS. The CES-D Scale. A self-report depression scale for research in the general population. Appl Psychol Meas. 1977;1(3):385–401.
28. Flett AE, Doney AS, Hogg S, et al. A multimodal approach to cardiovascular risk stratification in patients with type 2 diabetes incorporating retinal, genomic and clinical features. Sci Rep. 2019;9(1):3591.
29. McKay GJ, Paterson EN, Maxwell AP, et al. Retinal microvascular parameters are not associated with reduced renal function in a study of individuals with type 2 diabetes. Sci Rep. 2018;8(1):3931.
30. Ramos L, Novo J, Rocco J, et al. Retinal vascular tortuosity assessment: inter-intra expert analysis and correlation with computational measurements. BMC Med Res Methodol. 2018;18:144.
31. Alexopoulos GS. Depression in the elderly. Lancet. 2005;365(9475):1961–70.
32. Taylor WD, Aizenstein HJ, Alexopuolos GS. The vascular depression hypothesis: mechanisms linking vascular disease with depression. Mol Psychiatry. 2013;18(9):963–74.
33. Cooper DC, Tomfohr LM, Milic MS, et al. Depressed mood and flow-mediated dilation: a systematic review and meta-analysis. Psychosom Med. 2011;73:365–9.
34. Shi H, Feng G, Wang Z, et al. Relationships between depressive symptoms and an endothelial function among outpatients of a general hospital in China. Med Sci Monit. 2015;21:1812–9.
35. Brudley AJM, Korszun A, Jones CJH, et al. Arterial endothelial function is impaired in treated depression. Heart. 2002;85:21–3.
36. Smith PJ, Blumenthal JA, Hindelitter AL, et al. Microvascular endothelial function and neurocognition among adults with major depressive disorder. Am J Geriatr Psychiat. 2018;26(10):1061–9.
37. van Agtmaal MJM, Houben AJHM, Pouwer F, et al. Association of microvascular dysfunction with late-life depression: a systematic review and meta-analysis. JAMA Psychiatry. 2017;74(7):729–39.
38. Nguyen TT, Wong TY, Islam FM, et al. Evidence of early retinal microvascular changes in patients with type 2 diabetes and depression. Psychosom Med. 2010;72(6):535–8.
39. Owen CG, Rudnicka AR, Weikala RA, et al. Retinal vasculometry associations with cardiometabolic risk factors in the European prospective investigation of cancer-Norfolk study. Ophthalmol. 2019;126(1):96–106.
40. Sabanayagam C, Lye WK, Klein R, et al. Retinal microvascular calibre and risk of diabetes mellitus: a systematic review and participant-level meta-analysis. Diabetologia. 2015;58:2476–85.
43. Sasongko MB, Wong TY, Donaghue KC, et al. Retinal arteriolar tortuosity is associated with retinopathy and early kidney dysfunction in type 1 diabetes. Am J Ophthalmol. 2012;153(1):176–83 e1.

44. Sasongko MB, Wong TY, Nguyen TT, et al. Retinal vessel tortuosity and its relation to traditional and novel vascular risk markers in persons with diabetes. Curr Eye Res. 2016;41(4):551–7.

45. Sieu N, Katon W, Lin EH, et al. Depression and incident diabetic retinopathy: a prospective cohort study. Gen Hosp Psychiatry. 2011;33(5):429–35.

46. Wong TY, Knudtson MD, Klein BE, et al. Computer-assisted measurement of retinal vessel diameters in the beaver dam eye study: methodology, correlation between eyes, and effect of refractive errors. Ophthalmol. 2004;111(6):1183–90.

47. Cheung N, Tikellis G, Saw SM, et al. Relationship of axial length and retinal vascular caliber in children. Am J Ophthalmol. 2007;144(5):658–62.

48. Leung H, Wang JJ, Rochtchina E, et al. Computer-assisted retinal vessel measurement in an older population: correlation between right and left eyes. Clin Exp Ophthalmol. 2003;31(4):326–30.

49. Vilagut G, Forero CG, Barnaglia G, et al. Screening for depression in the general population with the Centre for Epidemiologic Studies Depression (CES-D): a systematic review with meta-analysis. PLoS One. 2016;11(5):e0155431.

50. Yates JA, Clare L, Woods RT, et al. Subjective memory complaints, mood and MCI: a follow-up study. Aging Ment Health. 2017;21(3):313–21.

Publisher’s Note
Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

Ready to submit your research? Choose BMC and benefit from:
• fast, convenient online submission
• thorough peer review by experienced researchers in your field
• rapid publication on acceptance
• support for research data, including large and complex data types
• gold Open Access which fosters wider collaboration and increased citations
• maximum visibility for your research: over 100M website views per year

At BMC, research is always in progress.
Learn more biomedcentral.com/submissions