Association between optical coherence tomography based retinal microvasculature characteristics and myocardial infarction in young men

Robert Kromer1, Eike Tigges2, Nargis Rashed1, Inga Pein2, Maren Klemm1 & Stefan Blankenberg2

Incident myocardial infarction (MI) is a leading cause of adult mortality in the United States. However, because MI has a relatively low incidence in the young population, little information exists on the disease in younger adults. Because the retina has the unique quality that its vasculature is readily and noninvasively visible, the retina is frequently studied to evaluate correlations between vessels and cardiovascular diseases. In the current study, we evaluated the retinal microvasculature of patients who had experienced an MI before 50 years of age (n = 53 subjects) and age- and sex-matched patients who had not experienced an MI (n = 53 patients). We used circular optical coherence tomography (OCT) scans to image peripapillary venules and arterioles. The diameter of each vessel was measured and the respective arterial-venous ratio (AVR) was calculated. We did not detect any significant differences between MI and control subjects in retinal vessel calibre or AVR.

Mortality from coronary heart disease (CHD) has significantly declined in the developed world over the past several decades. However, it remains the leading cause of adult mortality in the United States1. The CHD classification describes disease from subclinical atherosclerosis to acute coronary events, including myocardial infarction (MI). Even though CHD and its subsequent events primarily occur in patients over the age of 45 years, an age threshold has not been defined2. In fact, sudden cardiac death (predominantly due to CHD) is the most common cause of sudden death in adults under 40 years of age3. Therefore, CHD and its sequelae constitute an important health issue. This is especially true in younger patients who have a 15-year mortality rate of 30% (for patients < 40 years of age)4 and whose active lifestyles are at risk. Unfortunately, the mortality rate significantly increased to 45% for subjects who had experienced a previous MI4. These findings highlight the need to develop a risk stratification, which could lead to earlier diagnoses in high risk patients. However, CHD in the younger adult population is less common than CHD in older populations. As a result, comparatively little information exists on the condition in younger populations.

The retina has the unique characteristic that its vasculature can be directly visualized and its microcirculation can be noninvasively imaged in a matter of seconds. Retinal microvasculature analyses can quantify parameters related to vessel structure and function1. Additionally, blood vessels in the eye and heart are exposed to similar intrinsic- and extrinsic-influencing factors5. Moreover, the central retinal artery and its direct branches, generally evaluated with ophthalmoscopy, are true arteries because they have three vessel wall layers (intima, media, and adventitia). They also closely resemble small arteries in other organs, including the heart and brain6. Thus, changes in retinal vasculature characteristics may be early warning signs of ocular and/or systemic disease. In support of this theory, atherosclerotic changes in retinal arteries, characterized by thickening of the vessel wall, lipid deposition in the intima, and calcification of larger arteries, is strongly associated with atherosclerotic...
Our results indicate, with a power of 99%, that men who experienced an MI before the age of 50 years did not have changes in the retinal microvasculature, as examined with vessel calibre and AVR. However, coronary microvascular dysfunction is not likely an isolated process, but part of a systemic one\cite{16,17}. This theory is supported by the finding that retinal arteriolar narrowing may be an early indicator of microvascular damage caused by aging, hypertension, and/or other processes and that it reflects intimal thickening and medial hyperplasia, hyalinization, and sclerosis\cite{18}. Because similar pathological processes occur in the coronary arterioles of patients with hypertension\cite{19,20}, changes that occur in retinal arterioles may offer insight into changes that occur in the systemic microcirculation. Additionally, AVR, an index of retinal arteriolar narrowing, has been shown to be correlated with long-term average blood pressure\cite{21} and to be a marker of inflammation and endothelial dysfunction\cite{22}.
Furthermore, retinal vascular architecture changes could offer insight into subclinical CHD. Previous studies have shown that a smaller retinal arteriolar caliber is associated with left ventricular concentric remodeling, identified using cardiac MRI\textsuperscript{11,23}. This association remained significant even after adjusting for traditional risk factors. Our study only examined male subjects and our results may have been different if women had also been included in our study cohort. The Atherosclerosis Risk in Communities Study included 9648 patients and provided evidence that a smaller AVR was linked to an increased risk of incident CHD and MI in women, but not in men\textsuperscript{24}. Comparable gender differences in the association between AVR and CHD were also observed in the Blue Mountains Eye Study, Beaver Dam Eye Study, and Cardiovascular Health Study\textsuperscript{25–27}. A recent meta-analysis by McGeechan et al.\textsuperscript{28} (n > 22,000 subjects) concluded that both wider venules and narrower arterioles were associated with an increased risk of CHD in women, but not in men. It is important to note that all of the above-mentioned studies only included participants who were middle-aged or older. Therefore, it is likely that microvascular dysfunction plays a more integral role in the pathogenesis of CHD in women than in men\textsuperscript{29}. These gender differences may result from the influence of hormones or other substances that inhibit atherosclerosis development. Furthermore, these findings suggest that microvascular processes are the main contributors to CHD development\textsuperscript{30}, as supported by the finding that revascularization procedures have better outcomes in men than in women\textsuperscript{31}.

**Figure 1.** Correlation plots showing the influence of age on examined retinal microvasculature parameters in healthy control subjects (A–C) and in those who experienced a myocardial infarction (MI) at a young age (<50 years, D–F). The central retinal arterial equivalent (CRAE-6, A and D), central retinal venous equivalent (CRVE-6, B and E), and arteriovenous ratio (AVR, C and F) were examined. Age was not significantly correlated with any retinal microvasculature parameter examined (all p > 0.05). All data shown were collected from the right eye of study subjects.
A prior study showed that middle-aged and older men with a history of MI do not have an altered retinal microvasculature\textsuperscript{24}. Our study differed in that it only included patients who experienced an MI before the age of 50. Our study adds to the literature because little is known about baseline clinical characteristics and CHD risk factors in the young MI population\textsuperscript{32}. Prior studies that examined younger CHD or MI patients used an age cut-off of 40 or 45 years to distinguish between young and “nonyoung” patients\textsuperscript{33–38}. These investigations

**Table 2.** Retinal microvasculature parameters in healthy control subjects and in those who experienced a myocardial infarction (MI) at a young age (<50 years). All parameters were measured in both eyes. Measured values were compared between groups using two-tailed paired t-tests. Central retinal arterial equivalent (CRAE-6); Central retinal venous equivalent (CRVE-6); Arterial-venous ratio (AVR).

**Figure 2.** The arteriovenous ratio (AVR) in the right eye and left eye for healthy subjects and for those who experienced a myocardial infarction (MI) at a young age (<50 years). The AVR was not significantly different between groups in either the right eye (mean of differences = 0.002; \( p = 0.85 \)) or the left eye (mean of differences = −0.005; \( p = 0.56 \)). Statistical significance was tested using a paired t-test. Vertical columns represent 25–75% percentile with indicated mean and error bars represent 10–90% percentile.
found that young patients were more likely to be male, to be a smoker, and to have a family history of CHD. In agreement, young men with hypertension are significantly less likely to develop CHD than women in the same age group. In summary, epidemiological data, clinical findings, and the current results support the hypothesis that microvascular changes play little to no role in the development of CHD and MI in young men. Klein et al. showed in a study involving 8772 adults that the pathogenesis of retinal arteriolar changes differs. They demonstrated that retinal arteriolar changes like focal narrowing, generalized narrowing, changed AVR or arteriovenous nicking seem to be related to hypertension, inflammatory factors and endothelial dysfunction. However, clinical and subclinical atherosclerosis was seen independently to these retinal arteriolar changes. They suggested that atherosclerosis, as a disease of larger arteries, provides an independent prediction for ischemic diseases of the heart and other organs. In our study, we did not find any retinal arteriolar changes in terms of altered AVR in men who experienced an MI before the age of 50 years. Therefore, early-onset atherosclerosis might play a major role in the development of CHD and MI in this population rather than hypertension, inflammatory factors or endothelial dysfunction.

Previous research demonstrated a significant correlation between AVR and age, which was not replicated in our study. This might be attributed to the low patient number compared to epidemiological studies and the young age of the included participants. The study excluded patients with both high alcohol intake and body mass index $>30$ kg/m$^2$. These might have effects on the microvasculature and could lead to an altered AVR.

Our study had several limitations. First, patients were not examined immediately after the MI had occurred. In some cases, subjects had experienced the MI decades earlier. This may have influenced our results because arteriolar narrowing may have been alleviated with antihypertensive therapy. However, antihypertensive therapy does not affect retinal venules. Second, additional data regarding cardiovascular risk factors were not obtained. Third, our sample size adequately powered the epidemiological and population studies. Our study also had several strengths. This is the first study to evaluate the retinal vasculature in subjects who had experienced an MI at a young age. Additionally, SD-OCT is an objective, accurate, and established method for evaluating the retinal vessels. Lastly, only images that met the OSCAR-IB consensus criteria for retinal OCT quality were included in data analyses.

In summary, we demonstrated that men who had experienced an MI before 50 years of age do not have associated changes in the retinal microvasculature. Further studies are needed to identify factors that can feasibly be used as screening tools to determine which patients have a high-risk for MI. This information can then be used to support clinical decision making.

Methods

The study protocol was reviewed and approved by the Ethics Committee of the Hamburg Medical Council. All study conduct adhered to the tenets of the Declaration of Helsinki (7th revision, 64th Meeting, Fortaleza, Brazil) and followed Good Clinical Practices (GCPs). Written informed consent was obtained from each participant before study evaluations were performed.

Study subjects. Subjects who were already participating in the University Heart Center “MI Young Study” at the University Medical Centre Hamburg-Eppendorf were prospectively recruited for participation in the current study. The ongoing “MI Young Study” began in 2015 and is evaluating the risk profile of young (<50 years of age) MI patients using routinely-performed cardiac diagnostics, echocardiography, and genetic profiling. Age- and sex-matched healthy control subjects were prospectively recruited from the Department of Ophthalmology at the University Medical Center Hamburg-Eppendorf. Subjects were selected based on careful review of medical histories and records, with particular attention given to retinal and cardiovascular disease.

Subjects in the MI Young Study who met all of the following ophthalmic criteria were considered for study participation: (i) best-corrected logMAR visual acuity of 0.3 or better, (ii) spherical refractive error within $\pm 5.0$ dioptres (D), and (iii) cylindrical refractive error within $\pm 2.0$ D. Subjects were excluded if any of the following were present: (i) intense alcohol abuse (more than 40 g of alcohol daily), (ii) body mass index $>30$ kg/m$^2$, (iii) intraocular pressure $\geq 21$ mmHg, (iv) anterior ischemic optic neuropathy, or (v) congenital optic nerve abnormality. All participants in both the MI and control groups had a physiological or medication controlled systolic blood pressure within normal limits (within the range of 100–130 millimetres mercury (mmHg) systolic and 60–85 mmHg diastolic). Healthy control subjects with cardiovascular diseases, including angina, myocardial infarction, stroke, heart failure, hypertensive heart disease, cardiomyopathy, heart arrhythmia, congenital heart disease, valvular heart disease, carditis, aortic aneurysms, peripheral artery disease, thromboembolic disease and venous thrombosis, were excluded from the study.

Study evaluations. Study participants underwent a thorough ophthalmic examination, including: (i) slit lamp-assisted biomicroscopy (anterior and posterior segment thru non-dilated pupils) to identify any ocular pathology, (ii) non-contact tonometry (Nidek Tonometer NT-530) to identify undetected glaucoma, and (iii) SD-OCT imaging to collect study measurements (SPECTRALIS; Heidelberg Engineering, Heidelberg, Germany).

Optical coherence tomography. The SD-OCT system noninvasively obtains high-resolution, microscopic, cross-sectional images of the retina. The device combines conventional OCT technology with cSLO technology. A super-luminescent diode emits a light beam with a wavelength of 870 nm. The SD-OCT has a scan rate of up to 40,000 A-scans per second, a depth resolution of 7 µm, and a transverse resolution of 14 µm. The cSLO utilizes a laser to illuminate the retina and produce real-time, en face images. These fundus images are then used by the SD-OCT system as an anatomical reference. Before obtaining SD-OCT scans, the scan circle was centred on the optic disc. The automatic real-time averaging mode, resulting in the achievement of even higher quality. In this study, high-quality SD-OCT scans with at least 50 frame averaging were used to provide images with low
noise. Three high-resolution SD-OCT peripapillary scans with 3.5 mm in diameter and linked cSLO images were acquired from each subject. All SD-OCT images were obtained by a single examiner to minimize variability. Additionally, all included scans satisfied the consensus criteria for retinal OCT quality assessments (OSCAR-IB) to improve scan comparability and ensure that only the highest quality OCT images were used. More specifically, any image that did not meet all of the following quality criteria were excluded from analyses: (i) in focus and visible fundus before and during image acquisition, (ii) absence of scan and algorithm failures, and (iii) a continuous scanned layer. A representative peripapillary SD-OCT scan with a linked cSLO image is shown in Fig. 3.

Manual classification of retinal vessels in cSLO images. Retinal vessels were manually classified as arterioles or venules because automated classification methods do not yield adequate results, even when colour fundus photographs are used. Determining vessel classification is more complicated on cSLO images than on colour fundus images because cSLO images are monochromatic. Around the optic disc, only vessels that are close to each other can be reliably categorized as arteries or veins because a direct comparison is needed to reliably distinguish between vessel types. The following comparisons were used to categorize vessel: (i) the central light reflex is wider in arteries and smaller in veins, (ii) arteries are brighter than veins and veins are darker and deeper than arteries, and (iii) arteries are 30% thinner than their neighbouring veins.

Calculation of the arterial-venous ratio. Vessel diameters were indirectly measured on peripapillary SD-OCT scans using vessel-associated shadowing, as previously described. Vessel diameters were measured indirectly, and their shadow width on the SD-OCT scan was determined, assuming that retinal vessel blood flow attenuates the SD-OCT signal underneath the vessel. The central retinal equivalent was computed from these measurements using formulas established by Knudtson et al., which optimises the popular approach reported by Parr and Hubbard. One limitation of the Parr-Hubbard formula is that the number of vessels measured significantly impacts the overall AVR calculation. Knudtson et al. based their formula on the six largest retinal arterioles and venules to overcome this. The following equations were used to calculate CRAE-6 for arterioles and CRVE-6 for venules:

Figure 3. Peripapillary spectral domain optical coherence tomography (SD-OCT) scan (B) and the linked confocal scanning laser ophthalmoscope (cSLO) fundus image (A) obtained from a subject who had not experienced a myocardial infarction at a young age (<50 years). The peripapillary OCT scanning circle was 3.5 mm in diameter and is shown in the cSLO image (red circle, A). The manually identified vessel borders are shown on the SD-OCT scan (red lines, B).
Arterioles: $\hat{W} = 0.88 \omega_1^2 + w_2^{1/2}$

Venules: $\hat{W} = 0.95 \omega_1^2 + w_2^{1/2}$

where $w_1$, $w_2$, and $\hat{W}$ are, respectively, the widths of the narrower branch, the wider branch, and the estimate of parent trunk arteriole or venule. The AVR is then calculated as the quotient of CRAE-6 and CRVE-6 using the same iterative procedure of combining the largest and smallest vessel in each pairing.

### Statistical analyses.

Eyes from each participant were analysed separately (phenotype). Data are presented as mean ± standard deviation. All statistical analyses were performed using a commercially available software package (Prism 7 for Mac OSX; GraphPad Software, Version 7.0a, GraphPad Software, Inc., La Jolla, USA). Two-tailed, paired parametric t-tests were used to test the statistical significance of differences between continuous variables. Correlation analyses were performed using Pearson correlation calculations because data sampled from the populations followed an approximate Gaussian distribution. Statistical significance was defined as $P < 0.05$.

A two-tailed power analysis was also performed with a significance level (alpha) of 0.05. Because Schuster et al found an AVR difference of 0.09 between healthy patients and those with hypertensive retinopathy, the power analysis was performed for an AVR difference of 0.09. A post-test power analysis was also performed on the smallest detected difference to verify that observed differences were not insignificant because of relatively small sample size.

### References

1. Go, A. S. et al. Heart disease and stroke statistics–2014 update: a report from the American Heart Association. *Circulation* **129**, e28–e292 (2014).
2. Rubin, J. B. & Borden, W. B. Coronary heart disease in young adults. *Curr Atheroscler Rep* **14**, 140–149 (2012).
3. Arzamendi, D. et al. Increase in sudden death from coronary artery disease in young adults. *Am. Heart J.* **161**, 574–580 (2011).
4. Cole, J. H., Miller, J. I., Sperling, L. S. & Weintraub, W. S. Long-term follow-up of coronary artery disease presenting in young adults. *J. Am. Coll. Cardiol.* **41**, 521–528 (2003).
5. Liew, G., Wang, J. J., Mitchell, P. S. & Wong, T. Y. Retinal vascular imaging: a new tool in microvascular disease research. *Circ Cardiovasc Imaging* **1**, 156–161 (2008).
6. Flammer, J. et al. The eye and the heart. *European Heart Journal* **34**, 1270–1278 (2013).
7. Hogan, M. J. & Feeney, L. The ultrastructure of the retinal blood vessels. *I. The large vessels*. *J. Ultrastruct. Res.* **39**, 10–28 (1963).
8. Tedeschi-Reiner, E., Strozzi, M., Skoric, B. & Reiner, Z. Relation of Atherosclerotic Changes in Retinal Arteries to the Extent of Coronary Artery Disease. *The American Journal of Cardiology* **96**, 1107–1109 (2005).
9. Wong, T. Y. et al. Relation of retinopathy to coronary artery calcification: the multi-ethnic study of atherosclerosis. *Am. J. Epidemiol.* **167**, 51–58 (2008).
10. Klein, R., Klein, B. E., Moss, S. E. & Wang, Q. Hypertension and retinopathy, arteriolar narrowing, and arteriovenous nicking in a population. *Arch. Ophthalmol.* **112**, 92–98 (1994).
11. Wong, T. Y. et al. The prevalence and risk factors of retinal microvascular abnormalities in older persons: The Cardiovascular Health Study. *Ophthalmology* **110**, 658–666 (2003).
12. Leung, H. et al. Relationships between age, blood pressure, and retinal vessel diameters in an older population. *Invest. Ophthalmol. Vis. Sci.* **44**, 2900–2904 (2003).
13. Lowell, J. et al. Measurement of retinal vessel widths from fundus images based on 2-D modeling. *IEEE Trans Med Imaging* **23**, 1196–1204 (2004).
14. Wojtkowski, M., Bajraszewski, T., Targowski, P. & Kowalczyk, A. Real-time in vivo imaging by high-speed spectral optical coherence tomography. *Opt Lett* **28**, 1745–1747 (2003).
15. Cerone, R. et al. Ultra-high-resolution high-speed retinal imaging using spectral-domain optical coherence tomography. *Opt Express* **12**, 2435–2447 (2004).
16. Sax, F. & Cannon, R. S., Hanson, C. & Epstein, S. Impaired forearm vasodilator reserve in patients with microvascular angina. Evidence of a generalized disorder of vascular function? *JAMA* **28**, 1745–1747 (2003).
17. Lekakis, J. P., et al. Peripheral vascular endothelial dysfunction in patients with angina pectoris and normal coronary arteriograms. *Am. Coll. Cardiol.* **31**, 541–546 (1998).
18. Tso, M. O. & Jampol, L. M. Pathophysiology of hypertensive retinopathy. *Ophthalmology* **89**, 1132–1145 (1982).
19. Tanaka, M. et al. Quantitative analysis of narrowings of intramyocardial small arteries in normal hearts, hypertensive hearts, and hearts with hypertrophic cardiomyopathy. *Circulation* **75**, 1130–1139 (1987).
20. Moss, M., Yaron, R., Gotsman, M. S. & Hasin, Y. Histologic evidence for small-vessel coronary artery disease in patients with angina pectoris and patent large coronary arteries. *Circulation* **74**, 964–972 (1986).
21. Sharrett, A. R. et al. Retinal arteriolar diameters and elevated blood pressure: the Atherosclerosis Risk in Communities Study. *Am. J. Epidemiol.* **150**, 263–270 (1999).
22. Klein, R. et al. Are retinal arteriolar abnormalities related to atherosclerosis?: The Atherosclerosis Risk in Communities Study. *Arterioscler Thromb Vasc Biol* **20**, 1644–1650 (2000).
23. Cheung, N. et al. Retinal arteriolar narrowing and left ventricular remodeling: the multi-ethnic study of atherosclerosis. *J. Am. Coll. Cardiol.* **50**, 48–55 (2007).
24. Wong, T. Y. et al. Retinal arteriolar narrowing and risk of coronary heart disease in men and women. The Atherosclerosis Risk in Communities Study. *JAMA* **287**, 1153–1159 (2002).
25. Wang, J. et al. Retinal vascular calibre and the risk of coronary heart disease-related death. *Heart* **92**, 1583–1587 (2006).
26. Wong, T. Y. et al. Quantitative retinal venular calibre and risk of cardiovascular disease in older persons: the cardiovascular health study. *Arch. Intern. Med.* **166**, 2388–2394 (2006).
27. Wang, J. J. et al. Retinal vessel diameter and cardiovascular mortality: pooled data analysis from two older populations. *European Heart Journal* **28**, 1984–1992 (2007).
28. McGeechan, K. et al. Meta-analysis of retinal vessel calibre and risk for coronary heart disease. *Ann Intern Med* **151**, 404–413 (2009).
29. Bugiardini, R. & Bailey Merz, C. N. Angina with ‘normal’ coronary arteries: a changing philosophy. *JAMA* **293**, 477–484 (2005).
30. Wexler, L. F. Studies of acute coronary syndromes in women–lessons for everyone. *N. Engl. J. Med.* **341**, 275–276 (1999).
31. Tan, Y. C. et al. Gender differences in outcomes in patients with acute coronary syndrome in the current era: A review. *Eur Heart J Acute Cardiovasc Care* **5**, 51–60 (2016).
32. Cho, K. I. et al. Gender differences in risk factors and clinical outcomes in young patients with acute myocardial infarction. J Epidemiol Community Health 70, 1057–1064 (2016).
33. Doughty, M. et al. Acute myocardial infarction in the young—The University of Michigan experience. Am. Heart J. 143, 56–62 (2002).
34. Pineda, J. et al. Premature myocardial infarction: clinical profile and angiographic findings. Int. J. Cardiol. 126, 127–129 (2008).
35. Ergelen, M. et al. Comparison of outcomes in young versus nonyoung patients with ST-elevation myocardial infarction treated by primary angioplasty. Coron. Artery Dis. 21, 72–77 (2010).
36. Chua, S. K. et al. Acute ST-elevation myocardial infarction in young patients: 15 years of experience in a single center. Clin Cardiol 33, 140–148 (2010).
37. Bangalore, S. et al. Age and gender differences in quality of care and outcomes for patients with ST-segment elevation myocardial infarction. The American Journal of Medicine 125, 1000–1009 (2012).
38. Rathod, K. S. et al. Atypical risk factor profile and excellent long-term outcomes of young patients treated with primary percutaneous coronary intervention for ST-elevation myocardial infarction. Eur Heart J Acute Cardiovasc Care 5, 23–32 (2016).
39. Fox, C. S. et al. Increasing Cardiovascular Disease Burden Due to Diabetes Mellitus: The Framingham Heart Study. Circulation 115, 1544–1550 (2007).
40. Wong, T. Y. et al. Retinal microvascular abnormalities and 10-year cardiovascular mortality: a population-based case-control study. Ophthalmology 110, 933–940 (2003).
41. Piano, M. R. Alcohol’s Effects on the Cardiovascular System. Alcohol Res 38, 219–241 (2017).
42. Gong, W. et al. Effects of longitudinal body mass index variability on microvasculature over 5 years in adult Chinese. Obesity (Silver Spring) 24, 743–749 (2016).
43. Hughes, A. D. et al. Effect of antihypertensive treatment on retinal microvascular changes in hypertension. J. Hypertens. 26, 1703–1707 (2008).
44. Dahlöf, B., Stenkula, S. & Hansson, L. Hypertensive retinal vascular changes: relationship to left ventricular hypertrophy and arteriolar changes before and after treatment. Blood Press. 1, 35–44 (1992).
45. Tewarie, P. et al. The OSCAR-IB consensus criteria for retinal OCT quality assessment. PLoS ONE 7, e34823 (2012).
46. Li, H., Hsu, W., Lee, M. L. & Wang, H. A piecewise Gaussian model for profiling and differentiating retinal vessels. Image Processing 1, 1–1069–72 (2003).
47. Motte, J. et al. Vessel labeling in combined confocal scanning laser ophthalmoscopy and optical coherence tomography images: criteria for blood vessel discrimination. PLoS ONE 9, e102034 (2014).
48. Kondermann, C., Kondermann, D. & Yan, M. Blood vessel classification into arteries and veins in retinal images. Medical Imaging 6512, 651247–651247–9 (2007).
49. Schuster, A. K.-G., Fischer, J. E. & Vossmerbaeumer, U. Semi-automated retinal vessel analysis in nonmydriatic fundus photography. Acta Ophthalmologica 92, e42–9 (2014).
50. Knudtson, M. D. et al. Revised formulas for summarizing retinal vessel diameters. Curr. Eye Res. 27, 143–149 (2003).
51. Parr, J. C. & Spears, G. F. General caliber of the retinal arteries expressed as the equivalent width of the central retinal artery. Am. J. Ophthalmol. 77, 472–477 (1974).
52. Parr, J. C. & Spears, G. F. Mathematical relationships between the width of a retinal artery and the widths of its branches. Am. J. Ophthalmol. 77, 478–483 (1974).
53. Hubbard, L. D. et al. Methods for evaluation of retinal microvascular abnormalities associated with hypertension/sclerosis in the Atherosclerosis Risk in Communities Study. Ophthalmology 106, 2269–2280 (1999).

Author Contributions
R.K. and E.T. designed the study and its examinations. M.K. and S.B. revised the study protocol. R.K., N.R., and I.P. performed study examinations, compiled study data, and performed initial statistical analyses. R.K. and E.T. wrote the main manuscript text and prepared the figures. All authors reviewed the manuscript.

Additional Information
Competing Interests: The authors declare no competing interests.

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

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 license, and indicate if changes were made. The images or other third party material in this article are included in the article’s Creative Commons license, unless indicated otherwise in a credit line to the material. If material is not included in the article’s Creative Commons license 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 license, visit http://creativecommons.org/licenses/by/4.0/.

© The Author(s) 2018