

University of Dundee

Towards Standardization of Retinal Vascular Measurements

Mookiah, Muthu Rama Krishnan; McGrory, Sarah; Hogg, Stephen; Price, Jackie; Forster, Rachel; MacGillivray, Thomas J.

Published in:

Computational Pathology and Ophthalmic Medical Image Analysis - First International Workshop, COMPAY 2018, and 5th International Workshop, OMIA 2018, Held in Conjunction with MICCAI 2018, Proceedings

DOI:

[10.1007/978-3-030-00949-6_35](https://doi.org/10.1007/978-3-030-00949-6_35)

Publication date:

2018

Document Version

Peer reviewed version

[Link to publication in Discovery Research Portal](#)

Citation for published version (APA):

Mookiah, M. R. K., McGrory, S., Hogg, S., Price, J., Forster, R., MacGillivray, T. J., & Trucco, E. (2018). Towards Standardization of Retinal Vascular Measurements: On the Effect of Image Centering. In Z. Taylor, H. Bogunovic, D. Snead, M. K. Garvin, X. J. Chen, F. Ciompi, Y. Xu, L. Maier-Hein, M. Veta, E. Trucco, D. Stoyanov, N. Rajpoot, J. van der Laak, A. Martel, & S. McKenna (Eds.), *Computational Pathology and Ophthalmic Medical Image Analysis - First International Workshop, COMPAY 2018, and 5th International Workshop, OMIA 2018, Held in Conjunction with MICCAI 2018, Proceedings* (pp. 294-302). (Lecture Notes in Computer Science (including subseries Lecture Notes in Artificial Intelligence and Lecture Notes in Bioinformatics); Vol. 11039 LNCS). Springer Verlag. https://doi.org/10.1007/978-3-030-00949-6_35

General rights

Copyright and moral rights for the publications made accessible in Discovery Research Portal are retained by the authors and/or other copyright owners and it is a condition of accessing publications that users recognise and abide by the legal requirements associated with these rights.

- Users may download and print one copy of any publication from Discovery Research Portal for the purpose of private study or research.
- You may not further distribute the material or use it for any profit-making activity or commercial gain.
- You may freely distribute the URL identifying the publication in the public portal.

Take down policy

If you believe that this document breaches copyright please contact us providing details, and we will remove access to the work immediately and investigate your claim.

Towards standardization of retinal vascular measurements: on the effect of image centering

Muthu Rama Krishnan Mookiah¹, Sarah McGrory², Stephen Hogg¹, Jackie Price⁴, Rachel Forster⁴, Thomas J. MacGillivray³, and Emanuele Trucco¹

¹ VAMPIRE project, CVIP, Computing (SSE), University of Dundee, UK.

² Centre for Cognitive Ageing and Cognitive Epidemiology, University of Edinburgh, UK.

³ VAMPIRE project, Centre for Clinical Brain Sciences, University of Edinburgh, UK.

⁴ Usher Institute for Population Health Sciences and Informatics, University of Edinburgh, UK.

{m.r.k.mookiah,e.trucco}@dundee.ac.uk

Abstract. Within the general framework of consistent and reproducible morphometric measurements of the retinal vasculature in fundus images, we present a quantitative pilot study of the changes in measurements commonly used in retinal biomarker studies (e.g. caliber-related, tortuosity and fractal dimension of the vascular network) induced by centering fundus image acquisition on either the optic disc or on the macula. To our best knowledge, no such study has been reported so far. Analyzing 149 parameters computed from 80 retinal images (20 subjects, right and left eye, optic-disc and macula centered), we find strong variations and limited concordance in images of the two types. Although analysis of larger cohorts is obviously necessary, our results strengthen the need for a structured investigation into the uncertainty of retinal vasculature measurements, ideally in the framework of an international debate on standardization.

Keywords: Biomarkers, Retina, Microvasculature, VAMPIRE, Uncertainty.

1 Introduction and motivation

The eye is the only organ allowing direct, non-invasive and inexpensive observation of a rich portion of the human microvasculature. Ophthalmoscopic instruments include nowadays fundus cameras, optical coherence tomography (OCT), scanning laser ophthalmoscopes, ultra-widefield angiography, autofluorescence and OCT-angiography. Fundus camera imaging remains the most common modality, given its use in many decades of clinical practice and research. A number of software packages have been developed to quantitate the morphometry of the retinal vasculature efficiently in large numbers of images, e.g. IVAN [22], SIVA [23], QUARTZ [24] and VAMPIRE [25]. Coupled with the increasing availability of cross-linked clinical data repositories, the above has enabled a plethora of studies on retinal vascular biomarkers for a variety of conditions, among others diabetes, stroke, dementia, and cardiovascular disease. Morphometric vascular parameters commonly adopted include the central retinal arterio-venular equivalents (CRAE, CRVE) and their ratio, the arterio-venous ratio (AVR),

all of which summarize measures of vessel calibers around the optic disc (OD); measures of tortuosity; bifurcation coefficients; and the fractal dimension (FD), assessing the complexity of the vascular network. Details of such measurements can be found in many publications; see e.g. MacGillivray *et al.* [1] for an introduction.

A crucial assumption of biomarker studies is that the *retinal vascular measurements are accurate, consistent and reliable*. Accuracy, consistency and reliability depend, in turn, on a considerable number of factors [2,3], not all of which easily controllable. The effects of several of these factors have been reported in a few studies (Section 2). In this paper, we focus on a specific and little investigated factor, *the centering of fundus images*. Centering is commonly of two types: on the macula, named Type I, or on the OD, named Type II (Figure 1). Clinical protocols require one or both, and for one eye or both (left and right), depending on the pathology of interest. Crucially for our discussion, *a standard $\sim 30^\circ$ - 45° field-of-view (FOV) image centered on the macula does not capture vessels in nasal quadrants* (Figure 1) *which would be visible in an OD-centered image*. Many studies on retinal vascular biomarkers have drawn on existing clinical repositories of retinal images, but it is not always specified whether images of different types have been analyzed separately. To our best knowledge, no reports exist on the quantitative difference in retinal fundus measurements of the same eye induced by different centering. This is what we present in this paper, contributing to the body of studies summarized in the next section.

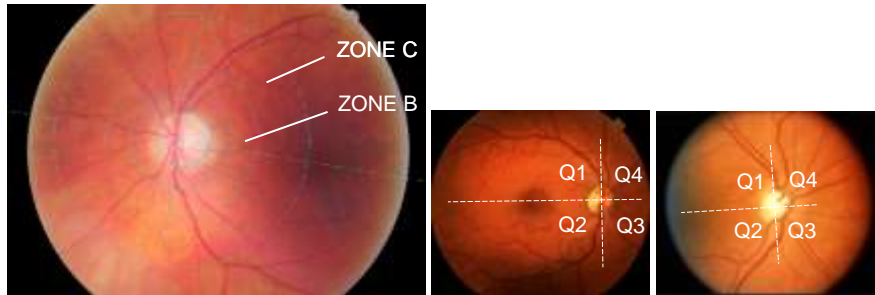


Figure 1. Illustration of quadrants and retinal coordinates centered on the OD and circular zones used to compute retinal measurements in a left eye (left). The x axis goes through the estimated centers of OD and macula. Examples of Type I (center) and Type II (right) images of right eyes; Type I (macula-centered) images miss parts of the vasculature in nasal quadrants (Q3, Q4).

2 Related work

The consistency and repeatability of retinal vascular measurements in fundus images, on which we focus (other imaging modalities have been considered, e.g. OCT [5]), has been investigated sparsely. The examples below, far from a systematic review, highlight concerns raised in the last decade. Notice that there is ample variability in the cohorts, numbers and statistical methods used in these studies themselves.

Chandler *et al.* [4] studied the variations of CRAE and CRVE, measured with IVAN in 30 fundus camera images (3 to 5 Mpixels from various cameras) of 3 subjects, applying systematically increasing blur. They found significant caliber broadening in

blurred images, and almost twice as much on average for CRAE (20 μm) as for CRVE (10 μm). Lim *et al.* [6] analyzed the effect of variations of axial length and myopic refractive errors on the retinal vasculature in an Asian diabetic population ($n=2,882$, Singapore Malay Eye Study), finding narrower retinal arterioles and venules, less tortuous arterioles, and increased branching coefficients in both arterioles and venules depending on axial length and refractive errors.

Knudtson *et al.* [7] analyzed the effect of the pulse cycle on width estimates in fundus images of 30 subjects. The retinal vessel diameter for one large arteriole/venule and one small arteriole/venule were measured by two trained graders. Results showed that the width of large retinal venules were less variable compared to that of arterioles. A related, more recent study was reported by Hao *et al.* [13].

Motivated by conflicting findings in the literature of the association of diseases with the FD, Huang *et al.* [8] performed a stability analysis of three FD measures (box, information and correlation coefficient) against different vessel segmentations from human annotators, automatic segmentation methods, threshold values, and regions of interest. Using 20 images from DRIVE [9], the authors observed substantial variations, leading them to recommend the use of vessel probability maps directly in biomarkers studies [10]. FD stability was also studied by Wainwright *et al.* [11] against variations of image quality, color, and format in a set of 30 images from the Blue Mountains Eye Study (9.6 Mb, 3,888x2,595 pixels), processed with IRIS-Fractal software. Simulated degradations resulted in significant variations of the FD coefficient.

Yip *et al.* [21] compared vessel width measurements (CRAE, CRVE) from two semi-automatic software applications, SIVA and IVAN, with 200 fundus camera images from the Singapore Chinese Eye Study. They found only moderate associations (ICC ~ 0.5 ; for ICC see Section 3, Analysis) and discordant associations with body mass index and arterial blood pressure. The same authors report similar results in [22] including the RA application in the comparison. Recently, McGrory *et al.* [3] reported a similar comparison of SIVA and VAMPIRE with 655 images of participants in the Lothian Birth Cohort 1936 studies. They found ICC values indicating poor to limited agreement for all retinal parameters (0.159–0.410), but consistent associations with systemic variables relating to blood pressure, as well as significant differences in the magnitude of association between retinal and systemic variables for 7 of 77 comparisons. We omit some reports of small-scales comparison between software applications for reasons of space.

Other authors analyzed morphometric parameter variations induced by further factors (see [3] and references therein) including image resolution, operators and fundus cameras make and models.

In summary, several, independent authors have measured considerable variability of retinal vascular measurements. Understanding and reducing such variability seems crucial as statistical associations within biomarker studies rely on accurate and consistent measurements. To our best knowledge, we contribute the first quantitative pilot study on the effect of fundus image centering, an important part of any imaging protocol.

3 Materials and Methods

Data set. 4 fundus-camera images of each of 20 subjects (2 per eye, macula and OD-centered, 80 images in total) were sourced from the Edinburgh Type 2 Diabetes Study (ET2DS), a population-based cohort study designed to investigate potentially modifiable risk factors for cognitive decrements in type 2 diabetes [12]. Images were acquired with a TOPCON TRC-50FX digital fundus camera at 35° FOV after pupil dilation using 1% tropicamide. Ethical approval for the ET2DS was granted by the Lothian Research Ethics Committee, and written informed consent was obtained from all participants; see Prince *et al.* [26] for details on the recruitment protocol. The images did not present diabetic lesions upsetting the detection and quantification of the vasculature, hence were considered a suitable sample for our purposes.

Retinal measurements. All images were measured by a trained operator (author [1]) with VAMPIRE 3.1 (Universities of Dundee and Edinburgh), obtained from its authors [1,3] following a standard protocol¹. For each image, VAMPIRE computes 151 measurements (Section 1) and their basic statistics (mean, median, standard deviation, max, min). Measurements are computed by vessel type (arteriole or venule), by region (zone, whole image, quadrant) and vessel (path, generation). We considered the 149 measures describing vessel morphology: 39 widths and functions thereof (e.g. CRAE, CRVE, AVR, basic statistics, width gradients, different width estimation algorithms by artery and vein, average ratio length-diameter at branching points), 104 tortuosity measurements, and 6 FD coefficients (3 per vessel network type, arterial or venous).

Analysis. Two-way mixed model intra-class coefficients (ICC) were computed to evaluate the extent of correspondence between two measurements (e.g., right eye versus left eye, or OD centered versus macula centered) of the same parameter (e.g., CRAE). The ICC quantifies this agreement, combining a measure of correlation with a test of the difference in means correcting for systematic bias and agreement based on chance alone. ICCs are thought to be more appropriate for assessing whether two variations in measuring a quantitative parameter provide similar results than Pearson's r , which measures the extent to which two variables are linearly dependent [14]. Method-comparison studies have demonstrated that a perfect linear relationship does not necessarily reflect good or even moderate agreement as measured by ICC [15,16]. ICC results are usually interpreted using 0.00–.49 = poor, 0.50–0.74 = moderate, and 0.75–1.00 = excellent [16]. Single-measure coefficients and 95% confidence intervals (CI) as well as correlations (raw, uncorrected Pearson's r s) were also computed. Tortuosity measurements were log-transformed to improve their distributions, which were positively skewed, as done elsewhere [3]. ICCs and Pearson's correlation were used to examine agreement between macula- and OD-centered images (right and left eye separately). In addition, we also analyzed measurement symmetry between right and left eye (macula and OD-centered images separately).

¹ VAMPIRE operator training consists of a 3-day module followed by periodic intra- and inter-operator repeatability and consistency checks on sets of images analyzed by certified, experienced operators.

4 Results

Full-result tables are reported in the supplementary material² and summarized here. Following a well-established protocol (based on [17] and developments), VAMPIRE requires a minimum number of vessels visible in Zone B and C. These are however only partially visible in macula-centered images (e.g. Figure 1, center), as nasal quadrants are minimally or not at all visible, leading to higher rejection rates than in OD-centered ones (not enough vessels), or to an analysis based on fewer vessels in fewer quadrants. 5 macula-centered images of the right eye had major AVR segments missing in Q2, and one image in Q1. Similarly, 3 macula-centered images of the left eye had AVR segments missing in Q2. All four quadrants were visible in all OD-centered images (right and left eye).

In the right eye, 5 **width-related measurements** (of 39) showed at least moderate correlation, association and significance (defined for our purposes as $r > 0.5$, ICC > 0.6 , $p < 0.1$) between OD- and macula-centered image, including CRAE, CRVE, arterial average ratio length-diameter in Zone C and the width gradient of the main artery in Q2 (LDR). Between 13 and 20 images supported these computations for OD- and macula centered images. In the left eye, only 3 width-related measures satisfied our conditions: CRAE (but not CRVE), AVR (not found in the right eye), and the venular (not arterial) LDR, with 12 to 20 images supporting the computation. Only the CRAE and arterial LDR satisfied our conditions in both eyes. For additional illustration, Bland-Altman graphs of two measures (right eyes) are shown in Figure 2.

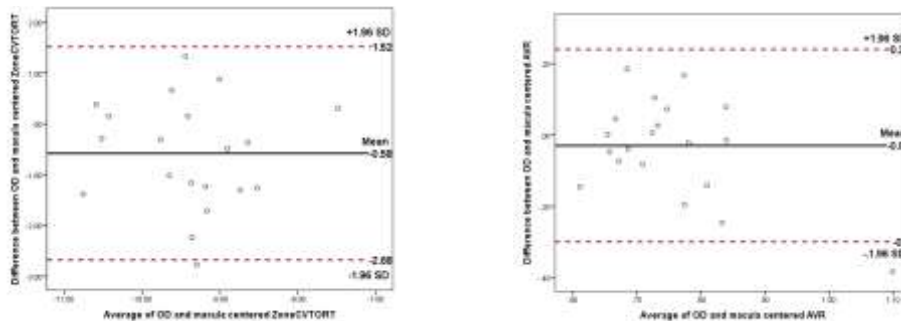


Figure 2. Bland-Altman plots visualizing, for illustration, the association of venular tortuosity in Zone C (left) and AVR (right) between OD- and macula-centered images (right eyes).

Tortuosity measures with at least moderate correlation and ICC (defined as above) between OD- and macula-centered images were only 17 (of 104) in the right and 20 in the left eyes. Of these, only 10 satisfied our conditions in both eyes: 8 arterial and 2 venular tortuosity measures, including 7 taken in Q1 and mean arterial tortuosity in Zone C. Of the 6 **FD measures** (3 for arteries, 3 for veins), only 2 of 20 images of the right eye supported full computation, leading to excellent but obviously not significant

²The Supplementary Material for this article can be found online at:
<http://vampire.computing.dundee.ac.uk/upload/files/Mookiah-et-al-OMIA-2018-Supplementary-Material/>

correlation and association; but in the left eyes, all 6 measures could be computed on the full set (20 images). Good and significant correlation ($r \sim 0.7$, $p < 0.01$) and moderate association (ICC ~ 0.7) was found for arterial measures only.

5 Discussion

Number of images and vessels measured. The absence or large occlusion of the nasal quadrants in macula-centered images implies higher image rejection rates (not enough vessels) or smaller number of vessels contributing to measurements compared to OD-centered images. Results show that *values for the same eye vary in the two cases*. What can we say of this variation (in our sample) is summarized below.

Width-related parameters. Given our results (Section 4), even imposing minimal requirements on r , ICC and significance, the effect of considering different sets of vessels for CRAE/CRVE and AVR calculations is considerable. This is supported by results reported by Heitmar *et al.* [18] in a related analysis.

Tortuosity. Again discrepancies between Type I and II images seem strong. In our sample, only 10 tortuosity measures satisfied our requirements simultaneously in both eyes. We notice that tortuosity values tend to be very small numbers, hence the numerical stability [19] of calculations involving them must be considered carefully.

Fractal dimension. There was a marked discrepancy between OD- and macula centered images, with venular measures missing altogether in the right eye due to the exclusion of too many macula-centered images. Again this suggests that omitting substantial parts of the nasal quadrants induces substantial changes on FD measures compared to OD-centered images. This supports related findings and concerns by Huang *et al.* [8] on the stability of the FD of the retinal vasculature.

Symmetry. The right-left symmetry of morphometric measurements of the vascular network remains an object of study [20]. Our pilot strengthens the hypothesis that good symmetry levels must not be taken for granted. For instance, CRAE, CRVE, and AVR were poorly and not significantly correlated in macula-centered measurements of either eye; but in the OD-centered images, CRAE and CRVE showed strong correlation (r btw. 0.837 and 0.859, $p < 0.001$) and excellent agreement (ICC value range, 0.837 and 0.857, 95% CI); and good, significant correlation ($r=0.673$, $p < 0.01$) was obtained for AVR. Similar discrepancies were found for tortuosity and FD (details omitted for conciseness).

6 Conclusions

To our best knowledge, we have reported the first pilot study on the quantitative changes in retinal measurements commonly used in retinal biomarkers studies induced by centering fundus image acquisition on the OD or on the macula. Our results suggest that different centering induces substantial differences. The important risk is that this could lead, potentially, to fragile statistical conclusions in biomarker studies. Such studies should, ideally, consider *both* centering types and discuss the differences in associations for Type I and Type II images *separately*.

The main limit of our pilot is the modest number of images and subjects (80 images, $n=20$), larger however than those in published reports on related topics (Section 2). We notice that the question itself of what statistical analysis methods are resilient to what levels of uncertainty and errors requires attention. A second limit is the use of only two, if commonly used, statistics (r and ICC). We plan to extend our analysis to larger samples from independent populations to better understand the effects of centering on morphometric vascular measurements in the retina.

Ultimately, the many aspects of a protocol for reliable biomarker studies, of which centering is only one, require in our view an international collaborative standardization effort, which we strongly auspicate.

Acknowledgements. This work was partially supported by EPSRC grant EP/M005976/1 “Multi-modal retinal biomarkers for vascular dementia”. MRK Mookiah is funded by a National Institute for Health Research Global Health Research (NIHR) award (Dundee-Chennai Unit on Diabetes Outcome Research). This paper presents independent research funded by the NIHR. The views expressed are those of the authors and not necessarily those of the NHS, the NIHR or the Department of Health. The support of NHS Lothian R&D, Edinburgh Imaging and the Edinburgh Clinical Research Facility at the University of Edinburgh is gratefully acknowledged. ET2DS was supported by a grant from the UK Medical Research Council. We would like to thank participants of ET2DS and staff at the Wellcome Trust Clinical Research Facility and Princess Alexandra Eye Pavilion in Edinburgh.

References

1. MacGillivray T, Trucco E, Cameron J, Dhillon B, *et al.* Retinal imaging as a source of biomarkers for diagnosis, characterization and prognosis of chronic illness or long-term conditions. *British Journ Radiol* 87(1040), 2014.
2. E. Trucco, A. Ruggeri, T. Karnowski, L. Giancardo, *et al.* Validating Retinal Fundus Image Analysis Algorithms: Issues and a Proposal, *Investigative Ophthalmology & Visual Science*, vol. 54, pp. 3546-3559, 2013.
3. McGrory S, Taylor A, Pellegrini E, Ballerini L, *et al.* Towards standardization of quantitative retinal vascular parameters: comparison of SIVA and VAMPIRE measurements in the Lothian Birth Cohort 1936. *Transl Vis Sci and Techn* 7(2), 2018.
4. CS Chandler, S Gangaputra, LD Hubbard, NJ Ferrier, *et al.* Suboptimal Image Focus Broadens Retinal Vessel Caliber Measurement. *Inv Ophth Vis Sci* 52(12), 2011.
5. Shin JW, Shin YU, Uhm KB, Sung KR, *et al.* The Effect of Optic Disc Center Displacement on Retinal Nerve Fiber Layer Measurement Determined by Spectral Domain Optical Coherence Tomography. *PLoS ONE* 11(10): e0165538, 2016. doi:10.1371/journal.pone.0165538.
6. Lim LS, Cheung CY, Lin X, Mitchell P, *et al.* Influence of refractive error and axial length on retinal vessels geometric characteristics. *Inv Ophth Vis Sci* 52(2), 2011.
7. Knudtson, M. D., *et al.* Variation associated with measurement of retinal vessel diameters at different points in the pulse cycle. *British Journal of Ophthalmology* 88.1 (2004): 57-61.
8. Huang F, Zhang J, Bekkers EJ, Dashtbozorg B, *et al.* Stability analysis of fractal dimension in retinal vasculature. *Proc Int MICCAI Workshop on Ophthalmic Medical Image Analysis (OMIA)*, Munich, 2015.

{PAGE * MERGEFORMAT}

9. Staal, J., Abramoff, M., Niemeijer, M., *et al.* Ridge based vessel segmentation in color images of the retina. *IEEE Transactions on Medical Imaging* 23(4), 2004.
10. Grauslund, J., Green, A., Kawasaki, R., *et al.*: Retinal vascular fractals and microvascular and macrovascular complications in type 1 diabetes. *Ophthalmology* 117(7), 2010.
11. Wainwright A, Liew G, Burlutsky G, Rochtchina E, *et al.* Effect of image quality, color, and format on measurement of retinal vascular fractal dimension. *Inv Ophth Vis Sci* 51(11), 2011.
12. Ding J, Strachan MW, Reynolds RM, Frier BM *et al.* Diabetic retinopathy and cognitive decline in older people with type 2 diabetes: the Edinburgh Type 2 Diabetes Study. *Diabetes* 59(11), 2883-2889, 2010.
13. Hao HAO, Sasongko MB, Wong TY. Does Retinal Vascular Geometry Vary with Cardiac Cycle? Retinal Vessel Geometry during Cardiac Cycle. *Inv Ophth Vis Sci.* 2012; 53(9): 5799-5805.
14. Lee J, Koh D, Ong CN. Statistical evaluation of agreement between two methods for measuring a quantitative variable. *Comput Biol Med.* 1989; 19: 61–70.
15. Bédard M, Martin NJ, Krueger P, Brazil K. Assessing reproducibility of data obtained with instruments based on continuous measurements. *Exp Aging Res.* 2000; 26: 353–365.
16. Portney LG, Watkins, MP. *Foundations of Clinical Research: Applications to Practice.* Upper Saddle River, NJ: Pearson/Prentice Hall; 2009.
17. Knudtson MD, Lee KE, Hubbard LD, Wong TY, *et al.* Revised formulas for summarizing retinal vessel diameters. *Curr Eye Res.* 27(3), 2003.
18. Heitmar R, Kalitzeos AA, Panesar V. Comparison of Two Formulas Used to Calculate Summarized Retinal Vessel Calibers. { HYPERLINK "https://www.ncbi.nlm.nih.gov/pubmed/26390347" \o "Optometry and vision science : official publication of the American Academy of Optometry." } 92(11), 2015.
19. G Strang. *Linear algebra and applications.* 5th edition, Wesley-Cambridge Press, 2016.
20. J.R. Cameron, R.D. Megaw, A.J. Tatham, S. McGrory, *et al.* Lateral thinking: interocular symmetry and asymmetry in neurovascular patterning, in health and disease. *Progress in Retinal and Eye Research* 59, 2017.
21. WF Yip, CY Cheung, H Hamzah, C Han, *et al.* Are computer-assisted programs for measuring retinal vascular caliber interchangeable? *Inv Ophth Vis Sci.* 53, 2012.
22. Wong TY,, Islam FM,, Klein R,, *et al.* Retinal vascular caliber, cardiovascular risk factors and inflammation: the multi-ethnic study of atherosclerosis (MESA). *Inv Ophth Vis Sci.* 47, 2006.
23. Cheung CY,, Tay WT,, Mitchell P,, *et al.* Quantitative and qualitative retinal microvascular characteristics and blood pressure. *Journ Hypertens* 29, 2011.
24. Fraz MM, Welikala RA, Rudnicka AR, Owen CG, *et al.* QUARTZ: Quantitative Analysis of Retinal Vessel Topology and size. *Expert Systems with Applications* 42(20), 2011.
25. E. Trucco, A. Giachetti, L. Ballerini, D. Relan, *et al.* Morphometric measurements of the retinal vasculature in fundus images with VAMPIRE. In *Biomedical Image Understanding: Methods and Applications*, J. Lim, S. Ong, W. Xiong, Eds., John Wiley & Sons, 2015.
26. Prince JF, Reynolds RM, Mitchell RJ, *et al.* The Edinburgh Type 2 Diabetes Study: study protocol. *BMC Endocr Disord* 18(8), 2008.