

# Validating Retinal Fundus Image Analysis Algorithms: Issues and a Proposal

Emanuele Trucco,<sup>1</sup> Alfredo Ruggeri,<sup>2</sup> Thomas Karnowski,<sup>3</sup> Luca Giancardo,<sup>4</sup> Edward Chaum,<sup>5</sup> Jean Pierre Hubschman,<sup>6</sup> Bashir al-Diri,<sup>7</sup> Carol Y. Cheung,<sup>8</sup> Damon Wong,<sup>9</sup> Michael Abramoff,<sup>10-13</sup> Gilbert Lim,<sup>14</sup> Dinesh Kumar,<sup>15</sup> Philippe Burlina,<sup>16</sup> Neil M. Bressler,<sup>17</sup> Herbert F. Jelinek,<sup>18</sup> Fabrice Meriaudeau,<sup>19</sup> Gwénolé Quéllec,<sup>20</sup> Tom MacGillivray,<sup>21</sup> and Bal Dhillon<sup>22</sup>

<sup>1</sup>VAMPIRE project, School of Computing, University of Dundee, Dundee, United Kingdom

<sup>2</sup>Department of Information Engineering, University of Padova, Padova, Italy

<sup>3</sup>Oak Ridge National Laboratory, Oak Ridge, Tennessee

<sup>4</sup>Istituto Italiano di Tecnologia, Genoa, Italy

<sup>5</sup>University of Tennessee Health Science Center, Memphis, Tennessee

<sup>6</sup>Jules Stein Eye Institute, Los Angeles, California

<sup>7</sup>REVIEW Group, University of Lincoln, Lincoln, United Kingdom

<sup>8</sup>Singapore Eye Research Institute, Singapore

<sup>9</sup>Institute for Infocomm Research, A\*STAR, Singapore

<sup>10</sup>Department of Ophthalmology and Visual Sciences, University of Iowa, Iowa City, Iowa

<sup>11</sup>Department of Electrical and Computer Engineering, University of Iowa, Iowa City, Iowa

<sup>12</sup>Department of Biomedical Engineering, University of Iowa, Iowa City, Iowa

<sup>13</sup>Department of Veterans Affairs, Iowa City VA Medical Center, Iowa City, Iowa

<sup>14</sup>School of Computing, National University of Singapore, Singapore

<sup>15</sup>RMIT University, Melbourne, Australia

<sup>16</sup>Applied Physics Laboratory, Johns Hopkins University, Laurel, Maryland

<sup>17</sup>Wilmer Eye Institute, Johns Hopkins University, Baltimore, Maryland

<sup>18</sup>Charles Sturt University, Albury, Australia

<sup>19</sup>University of Burgundy, Le2i, Le Creusot, France

<sup>20</sup>INSERM, U1101, SFR ScInBioS, Brest, France

<sup>21</sup>VAMPIRE project, Clinical Research Imaging Centre, University of Edinburgh, Edinburgh, United Kingdom

<sup>22</sup>VAMPIRE project, NHS Princess Alexandra Eye Pavilion, University of Edinburgh, Edinburgh, United Kingdom

Correspondence: Emanuele Trucco, VAMPIRE/CVIP, School of Computing, University of Dundee, Dundee, UK; manueltrucco@computing.dundee.ac.uk.

Submitted: June 6, 2012

Accepted: December 2, 2012

Citation: Trucco E, Ruggeri A, Karnowski T, et al. Validating retinal fundus image analysis algorithms: issues and a proposal. *Invest Ophthalmol Vis Sci.* 2013;54:3546-3559. DOI:10.1167/iops.12-10347.

This paper concerns the validation of automatic retinal image analysis (ARIA) algorithms. For reasons of space and consistency, we concentrate on the validation of algorithms processing color fundus camera images, currently the largest section of the ARIA literature. We sketch the context (imaging instruments and target tasks) of ARIA validation, summarizing the main image analysis and validation techniques. We then present a list of recommendations focusing on the creation of large repositories of test data created by international consortia, easily accessible via moderated Web sites, including multicenter annotations by multiple experts, specific to clinical tasks, and capable of running submitted software automatically on the data stored, with clear and widely agreed-on performance criteria, to provide a fair comparison.

Keywords: validation, fundus image analysis, reference standards

This paper was born of discussions involving several international research groups on automatic retinal image analysis (ARIA; see Ref. 1 for a recent review) at IEEE Engineering in Medicine and Biology Conference (EMBC) 2011, Boston. There was unanimous recognition of two key facts.

1. Efforts in the community are shifting from generating algorithms to detect, localize, or measure retinal features and properties, validated with small sets of test data, to

generating measurements of clinical and public health significance for clinicians, eye care providers, and biomedical scientists and researchers, requiring larger and “real-life” sets of test data.

2. The current methods for validating ARIA algorithms are neither uniform nor widely agreed upon. Issues include how to deal with the variability of expert annotations; the availability of public, large, structured “real-life” datasets for testing; and the accepted definition of reference (gold) standards in different applicative contexts.

It was felt that the discussion was sufficiently important to seek the opinion of further groups, and to publish the result of the discussion. This paper is that result.

Purely for reasons of space and consistency, we concentrate on the validation of algorithms processing fundus camera images, currently the largest section of the ARIA literature.

ARIA algorithms are currently used for the following main purposes.

1. *Screening/monitoring*, for example of diabetic retinopathy (DR), glaucoma, or age-related macular degeneration. The goal is to identify images showing signs of a target condition in large sets (tens of thousands to millions). The images (patients) selected are referred for clinical attention. False negatives (missing patients with disease) must be minimized; limited amounts of false positives (false alarms) are acceptable, but should also be considered as a factor to avoid unnecessary use of resources or side effects of unnecessary treatments. It has been shown that appropriate screening of DR is cost-effective.<sup>2,3</sup> DR screening facilitates early detection of patients with mild stages of DR, and thus early intervention (e.g., by targeting a patient's blood glucose and blood pressure levels, or by laser treatment) and ultimately the prevention of vision loss (outcome of interest). ARIA screening promises to eliminate inefficiencies within the current DR screening workflow by providing a faster, more cost-effective and accurate disease diagnosis. It will also eventually improve economics of eye disease management and cost savings for patients, public health care providers, and the government, and improve general eye health.
2. *Computer-assisted* diagnosis and risk stratification, for example diagnosis of retinopathy of prematurity (ROP)/Plus disease given measurements of tortuosity and width not detected readily by clinical examination alone. The purpose is to detect the presence or likelihood of a disease from specific signs. ARIA performance must be demonstrated to be more precise than diagnosis in the absence of computer assistance or generate richer data improving a clinician's diagnosis. Unlike that with screening and monitoring, the outcome is not necessarily binary (refer/do not refer), and the diagnosis usually depends on a combination of factors beyond ARIA measurements (e.g., age, symptoms, clinical features).
3. (c) *Biomarkers* aimed to determine whether the occurrence of measurable features in retinal images is linked significantly (in a statistical sense) with specific outcomes or conditions that impact treatment decisions, prognosis, or diagnosis, for example retinal vessel width with lacunar stroke and coronary heart disease. ARIA features may also be useful for testing effects of new drugs and therapies (e.g., changes in retinal vascular parameters with a novel drug for hypertension) or for discovery of novel pathways in the natural history of diseases (e.g., microvascular disease pathways in heart attacks). Links to cognitive performance and gene expression have also been reported.<sup>4,5</sup>

In addition, three further areas would benefit from reliable ARIA systems:

4. *Longitudinal* studies, whereby ARIA provides a means to study quantitatively the evolution and characterization of a disease to assist treatment planning or gauge patient response to a treatment.
5. *Computer-aided* or image-guided surgery, an emerging application of ARIA algorithms,<sup>6-9</sup> for example, vitreo-retinal microsurgery, for which ARIA allows registration

of intraoperative imagery with preoperative imagery for image-guided surgical interventions.<sup>10</sup>

6. *Telehealth*. ARIA disease screening and monitoring could play a very important role here, for example in less developed countries where the incidence of diabetes is rising and screening made difficult by the lack of resources and clinicians.<sup>11</sup> A World Health Organization consultation report reviewing principles for the care of diabetic retinopathy noted that retinal imaging systems and image analysis methods perform at least as well as human providers. Computer-assisted telehealth programs can therefore become a scalable method for providing expert care. The cost-effectiveness of telehealth screening of DR is accepted as an alternative to eye specialist exams in the United States, and is applied on a societal scale under various national health care programs, for example in the United Kingdom.<sup>12,13</sup>

This paper combines input from 14 international research groups on the validation of ARIA algorithms. We first define "validation," hence the scope of our discussion, and sketch the main techniques reported to date. We then give some compact background on the images and tasks for which ARIA algorithms are developed. We then discuss the issues making validation of ARIA algorithms difficult, and conclude with recommendations. We include in Appendix A a list of public datasets currently available for testing ARIA algorithms.

## VALIDATION IN RETINAL IMAGE ANALYSIS

### Validation: A Definition

For our purposes, *validation* indicates *the process of showing that an algorithm performs correctly by comparing its output with a reference standard*. In ARIA, target performance levels are normally represented by a reference ("gold") standard defined by expert performance, for example, regions traced manually around landmarks or lesions, image quality level, or scores attached to DR screening images.

Validating ARIA algorithms implies therefore (1) selecting a data (image) sample representative for the specific validation purposes, (2) collecting reference standard annotations on the sample images, (3) running algorithms on the sample images, and (4) comparing the output with the reference standard by performing statistical analysis to assess the agreement of the algorithm's output and reference standard, for example sensitivity, specificity, positive and negative predictive value, and area under ROC (receiver operating characteristic) curve.

### Techniques

We identify four main types of validation processes in the ARIA literature, each involving its own reference standards. From the most general (defined in terms of clinical concepts) to the most detailed (defined in terms of image elements), these are as follows:

1. *Outcome* oriented, for example disease/no disease;
2. *Disease* grading, for example severity of DR, ROP Plus or Pre-plus;
3. *Feature* grading, for example tortuosity level of vessels or eye vasculature, width of retinal vessels;
4. *Image/pixel/measurement* oriented, for example locating microaneurysms, measuring area or perimeter of target regions, locating vessel bifurcations.

The key validation task is to *assess quantitatively the agreement of automatic and manual measurements*. How exactly to declare agreement or disagreement between sets of data is the object of discussion in the literature. Techniques vary with the type of the validation process (see above), but in general they are drawn from statistics. The ones reported most frequently in ARIA papers include graphs (e.g., scattergrams, Bland-Altman plots), integral indexes like correlation coefficients (e.g., Pearson), and statistical tests. ROC curves and associated coefficients (e.g., specificity, sensitivity, area under the curve), imported from signal processing, are frequently used to quantify the accuracy of detection and classification and are advocated strongly by some authors.<sup>14</sup>

Two major issues in the generation of reference standard data are the *variability of expert judgment* and the *need for generating annotations directly comparable to the algorithm's output*. The former is addressed by having several experts annotate the same dataset; however, it is not ultimately clear how to obtain a single reference value from multiple ones (e.g., by averaging of values, discussion and consensus, interrater reliability metrics such as ACI or Kappa, or just keeping histograms/distributions of multiple values). This is because some annotation tasks are not part of normal clinical practice and clinicians are not used to them, or do not see their relevance (e.g., tracing accurate contours around lesions).

A related question is where to set the “outcome” for validation. In a screening program, a refer/no refer decision with an associated uncertainty level seems the obvious choice; other cases are not so clear. A related point is that ARIA algorithms often consist of a pipeline of modules; and while testing the outcome of the algorithm is the main goal, it may be interesting and useful to test each individual module.

For these reasons some authors have begun to explore alternative validation paradigms. One is estimating *simultaneously* the quality of ARIA results and reference standards summarizing annotations from multiple experts, for example STAPLE for image segmentation<sup>15,16</sup> and other tasks.<sup>17</sup> Imperfect reference standards are the motivation behind weak learning methods in pattern recognition and machine learning<sup>18,19</sup>; these methods are currently used only rarely in ARIA<sup>20</sup> but might provide very useful modeling tools (see also subsection “Variation of Expert Judgment”). An interesting viewpoint is offered by Quellec et al.,<sup>21</sup> who found, in brief, that a disagreement on DR severity between the algorithm and one expert would predict disagreement between the expert and a more experienced one. The ability to model expert disagreement would be a powerful tool for validation.

## IMAGES AND TASKS

We discuss the characteristics of digital images and of the instruments generating them, as well as the clinical tasks for which images are ultimately created and analyzed. Both play a substantial role in validation. For reasons of space, we omit other imaging modalities like optical coherence tomography (OCT) and fluorescein angiography (FA) angiography.

### Instruments and Images

The majority of ARIA systems reported to date consider single color images from digital fundus cameras, although acquiring two images per eye for both eyes (posterior pole, optic disc [OD] centered) is increasingly common in screening programs. Images are acquired with or without dilating the patient's pupil through eye drops, respectively *mydriatic* and *nonmydriatic*. In the latter case, cameras require a high-power flash to allow

enough light to enter the pupil and be reflected back by the retina. Concurrent illumination and imaging, performed through the same optical path, is the main engineering challenge faced by fundus cameras.

The main manufacturers of fundus cameras are currently Zeiss (Wetzlar, Germany), CenterVue (Padova, Italy), Topcon (Oakland, NJ), Nidek (Gamagori, Japan), Canon (Tokyo, Japan), Kowa (Nagoya, Japan). Depending on the clinical application, various optical filters are available, the most common being green filters for red-free photography and barrier filters for fluorescein angiography. A typical camera has a field of view (FOV) of 30° to 50° with  $\times 2.5$  magnification. Some modifications are possible through zoom or auxiliary lenses, from 15°, which provides  $\times 5$  magnification, to 140° with a wide-angle lens, which reduces the image by half. The actual image resolution, that is, how many millimeters are captured by a pixel, depends on various factors, mainly the properties of the optics and the resolution of the complementary metal-oxide-semiconductor/charge-coupled device (CMOS/CCD) image sensor employed; approximately  $3000 \times 3000$  pixels is now common. Higher-resolution sensors could be used, but the optics pose a limit to the resolution achievable, that is, to the size of the smallest distance that can be imaged in focus. As the optical quality of the eye prevents resolving features smaller than 20  $\mu\text{m}$ , very high resolutions without adaptive optics may not be useful. We notice that the diffusion of current imaging equipment and technology is somewhat limited by infrastructure requirements; most existing are bulky, are expensive (approximately \$25,000), and require special skills to operate. User-friendly, cost-effective handheld retinal cameras have been developed to tackle these issues, but work in this area is still limited.<sup>19,22</sup>

### Image Quality

Image quality depends on acquisition procedures, operators and their training, blur, occlusions (e.g., cataract, eyelashes), widespread lesions, artifacts introduced by the instrument, and conditions; for instance, fundus images of infants for ROP assessment tend to be lower quality than those of adults. Quality considerations are essential for proper validation, as image quality is at the basis of exclusion criteria applied in screening programs.

As quality influences the performance of ARIA systems, it seems advisable to divide a set of test images into quality classes, for example good, acceptable, and unusable. In the interest of applicability, quality classes should be defined by practitioners, using national standards for specific tasks whenever present. The UK diabetes screening program, for instance, defines three image quality grades (inadequate, minimum, achievable).<sup>23</sup>

Automated systems for assessing retinal image quality exist,<sup>24–29</sup> but quality is not always considered in the wider ARIA literature with respect to preparing datasets. Capturing quality definitions applied by experts in an ARIA algorithm is difficult, as clinicians learn from examples and practice; images considered viable for clinical analysis may not always produce good results with ARIA systems. ARIA quality estimation has often relied on heuristic measures (e.g., contrast level of vessels in specific retinal regions) and the use of approximate classes for quality classifications such as “adequate” or “inadequate.” Such labels are used in supervised learning systems, combined with pattern recognition and image-processing methods based on image features like sharpness of vessel regions, quantity of the blood vessel, and color characteristics like the shape of the color histogram.



## Tasks

To complete the background picture for our discussion of validation, we identify the main ARIA tasks addressed in the literature and the main measures used for validation. We do not aim to review ARIA techniques for each task; for these the reader is referred to the recent review by Abramoff et al.<sup>1</sup>

### Anatomical Landmarks: Location and Measurement.

In fundus images, the main landmarks are the OD, the macular region, and the vasculature. Most detection methods reported combine anatomical knowledge (relative locations of retinal landmarks, vasculature geometry) with image features (brightness levels, vessel density, orientation and thickness) to locate OD and macula, thus identifying a reference coordinate system for the location of retinal lesions.

Reference standard sets must specify therefore OD and macula locations. Measures used to compare reference standard annotations and ARIA estimates are normally the distance between annotated and estimated OD centers, or integral measures of contour agreement based on area (e.g., the Dice coefficient), or point-to-point distances along the contours.

The target clinical task determines the accuracy of the “location of the anatomical landmark.” For example, locating OD (with approximate size) and macula centers is generally sufficient to establish a retinal coordinate system; estimating the ellipticity of the OD requires accurate contour detection including parapapillary atrophy segmentation.<sup>30</sup>

### Vasculature Segmentation and Measurement.

The retinal vasculature is an important indicator of various diseases; its changes underlie the development of other signs such as retinal lesions. Indicators of disease in the vasculature include width and tortuosity changes, venous beading, focal arterial narrowing and neovascularization. Measurements can be local (e.g., width, branching angles) and global (e.g., fractal dimension of whole network).<sup>31,32</sup> Subtle vessel changes may occur during the early stages of disease development, consequent to changes to blood flow dynamics; and associations of these changes have been found with age or with risk of stroke.<sup>31,33</sup> There is therefore substantial interest in automatically segmenting the vasculature and measuring its properties. This requires, typically, locating the vessels, establishing overall branching patterns and connectivity, and computing target measurements.

The majority of datasets used for validating vasculature-related ARIA algorithms (see Appendix A) concentrate on the first step. They consist of images with corresponding reference standard in the form of vascular masks (binary pixel images) generated by clinicians using a software drawing tool. Two well-known examples are DRIVE and STARE.

These datasets suffer from two limitations. First, there is currently no absolute, objective definition of the location of the edge of a retinal blood vessel. The observed vessel in a standard retinal image corresponds to the blood column within the vessel. However, as the column depth reduces toward the vessel edge, the intensity drops off, blurring the edge's appearance. Second, generating vessel masks is one of the most labor-intensive annotation tasks. The anti-aliasing effect along vessel edges makes it difficult to determine exactly whether an individual pixel belongs properly to a vessel.

Both DRIVE and STARE use multiple observers, who provide sometimes significantly different reference standards. As algorithms approach human levels of performance, it becomes difficult to use such variable standards to assess performance. On the other hand, the datasets provide an easily accessible reference standard. The REVIEW dataset uses a different approach. A limited number of vessels have their edges marked using a contour tool with subpixel accuracy.

This allows more accurate assessment of algorithms for vessel width determination, but does not provide sufficient detail to analyze the overall segmentation performance.

Work addressing vascular connectivity exists, and some standard definitions of angles have been suggested.<sup>34</sup> However, there appears to be relatively little published work on vessel branching angles, and we know no publicly available reference standard dataset addressing this aspect.<sup>35,36</sup>

### Diabetic Retinopathy and Diabetes-Related Retinal Lesions.

DR has attracted a large part of ARIA research, and we dedicate a longer background section to it. Prevalence is expected to grow exponentially,<sup>37,38</sup> affecting 300 million people worldwide by 2025.<sup>39</sup> DR, a specific microvascular complication of diabetes, is a major cause of vision loss in people aged 25 to 60. Of the 246 million people with diabetes, approximately a third have DR, and a third of these have vision-threatening retinopathy—the majority caused by diabetic macular edema (DME).<sup>40</sup> DR imposes a huge economic burden on patients, health care systems, and society, estimated at US\$500 million annually in the United States alone.

This background poses a demand for reliable automated early-screening procedures. The challenge for ARIA is to find cost-effective techniques with sufficient sensitivity and specificity to reliably identify those at risk of vision loss. Clinically, the primary validation method of interest is outcome oriented: how well the presence of disease can be detected (refer/no refer), and how sensitive the systems can be made (near-zero false negatives) with a manageable level of specificity.

Some researchers<sup>2,41–44</sup> have reported systems with performance deemed acceptable for clinical deployment, but reports of large-scale studies remain rare and may not hold true for particular datasets with unique idiosyncrasies. Lesions targeted by ARIA systems include microaneurysms, cotton wool spots, soft exudates, and small hemorrhages for nonproliferative retinopathy; and ischemic areas in the retina, loss of vessels, and vessel proliferation for proliferative retinopathy.<sup>45–48</sup>

Table 1 summarizes ARIA work on DR detection, in particular microaneurysms and exudates, over the past 20 years. Many algorithms have been designed for the detection of DR in various types of retina images (color fundus, angiogram, red-free). The majority of these algorithms are validated on modest numbers of retina images annotated by experts, usually not available publicly. Performance is evaluated in terms of sensitivity and specificity at either the lesion, region, or image level.

In general, results depend on methodology and dataset, stressing the need for large, internationally agreed-upon datasets for validation. For instance, the winning team of the Retinopathy Online Challenge (provided in the public domain at <http://roc.healthcare.uiowa.edu/>, Niemeijer et al.<sup>49</sup>), led by Quéllec, achieved only 60% sensitivity with eight false positives per image; but earlier work by Niemeijer (2005)<sup>50</sup> reported 100% sensitivity and 87% specificity on image-level screening, even though the per lesion sensitivity was only approximately 30%.

**Glaucoma.** Glaucoma is a disease of the optic nerve, resulting in a gradual and progressive loss of vision. The main indicator in ARIA is the cup-to-disc ratio (CDR), that is, the ratio of the size of the optic cup to that of the OD. Various imaging techniques are used in relation to glaucoma. Currently, fundus photography remains the only modality in which the characteristic colors of the retina and pathologies are preserved. Stereo imaging approaches exist in which a depth map of the OD region is computed from two retinal photographs acquired from displaced viewpoints.<sup>71,72</sup> Tomographic imaging of the retina in three dimensions has been made possible by confocal laser scanning, used in the Heidelberg retinal tomograph, and OCT, which exploits interferometry to achieve tomographic micrometer-resolution imaging of the retinal layers.

TABLE 1. ARIA Literature on DR at a Glance

Reference No.	DR Task	Method	Public Dataset?	No. of images	Image Type	Validation Level	Sensitivity	Specificity
2	MA, dot HM	Pixel clustering, kNN	No	16,670	Color fundus	Image	47.7%	90%
51	MA	Radon transform, wavelet preprocessing	Yes	100	Color fundus	Lesion	50%	>10 false positive per image
49	MA	Wavelet transform	Yes	100	Color fundus	Lesion	60%	8 false positive per image
52	MA	Double-ring filter	Yes	100	Color fundus	Lesion	65%	27 false positive per image
53	MA	Wavelet, genetic algorithm	No	1115	Color fundus + angiogram	Lesion	90.24%	89.75%
54	MA	Diameter closing, feature extraction, classification	No	115	Color fundus	Lesion	88.50%	2.13 false positive per image
55	MA	Edge inference	No	49	Color fundus	Lesion	68%	>40 false positive per image
48	Exudates	Machine learning	No	430	Color fundus	Lesion	95%	88%
43	MA, dot HM	Combination of methods	No	15,473	Color fundus	Image	97.90%	67.40%
56	MA	Watershed contrast normalization	No	1677	Color fundus	Image	85.40%	83.10%
57	MA	Wavelet template matching	No	995	Green channel fundus	Lesion	87.90%	96.20%
58	MA	Generalized eigenvectors	No	70	Color fundus	Region	93%	NA
59	MA	2-D adaptive filtering, region growing	No	11	Angiogram	Region	90.72%	82.35%
50	MA, dot HM	Pixel classification	No	240	Color fundus	Region	100%	87%
60	MA, dot HM	RetinaLyze (proprietary)	No	400	Color fundus	Image	96.70%	71.40%
61	Drusen	Histogram-adaptive local thresholding	No	23	Color fundus	Region	98.80%	99.31%
62	MA, HM	Morphological, region growing, neural network	No	142	Color fundus	Region	77.50%	88.70%
63	Exudates	Dynamic clustering with domain knowledge	No	543	Color fundus	Image	100%	74%
64	MA	Morphological, region growing	No	46	Color fundus	Image	90%	80%
65	Exudates	Statistical classification with local window-based verification	No	200	Color fundus	Image	100%	70%
66	MA	Manual rule-based classifier	No	3885	Red-free	Image	85%	76%
46	MA, HM, exudates, cotton wool	Estimate background intensity, extract candidate regions for classification	No	268	Color fundus	Lesion	94%	69%
67	MA, dot HM	Pattern recognition	No	400	35-mm color slides	Lesion	NA	NA
68	MA	Manual rule-based classifier	No	88	Angiogram	Lesion	82%	84%
69	HM, exudates	Neural network	No	480	Red-free	Region	73.80%	73.80%
70	MA	Matched filter	No	6	Angiogram	Lesion	45%	>150 false positive per image

MA, microaneurysm; HM, haemorrhage; kNN, k-nearest neighbor.

Table 2 summarizes glaucoma-related ARIA reports (although numerous papers have described individual optic disc and cup detection, only papers leading to a CDR or glaucoma detection outcome have been included). ARGALI is a recent ARIA system for glaucoma assessment.<sup>73-75</sup> It uses active contour methods based on level sets to segment the cup and the disc and calculate the CDR. A similar approach was adopted by Joshi et al.<sup>76</sup> Some authors have reported stereo techniques recovering depth information.<sup>71,72,77,78</sup> An alternative approach is the use of machine learning to assign a predictive score for the risk of glaucoma directly to images.<sup>79</sup> AGLAIA (Automatic GLaucoma Diagnosis and Its Genetic Association Study through Medical Image InformAtics)<sup>67,80,81</sup> computes 13 image cues for glaucoma assessment, and aims to integrate clinical and genome data in a holistic glaucoma analysis.

No publicly available datasets for validation are known to us, although some may be available on request from the authors under specific agreements, for example ORIGA-light.<sup>80-82</sup>

**Retinopathy of Prematurity.** Retinopathy of prematurity (ROP) is a disease involving abnormal development of retinal vasculature in premature infants, which can lead to retinal detachment and visual loss. The main indicators of ROP severity are the Plus disease, the stage when treatment is required, and the Pre-plus disease, a predictor of sight-threatening Plus disease development. Plus and Pre-plus diseases can be diagnosed by recognizing their specific signs, namely abnormal vascular dilation and tortuosity. The number of infants requiring ROP examinations has recently increased, thanks to improved survival of very low-birth-weight infants.

TABLE 2. ARIA Literature on Glaucoma at a Glance

Reference No.	Method	Public Dataset?	No. of Images	Image Type	Validation Level	Results
83	Intra-image learning	ORIGA, <sup>84</sup> on request	650	Nonstereo fundus	CDR	Mean CDR error: 0.081
85	Statistical model based	ORIGA, <sup>84</sup> on request	650	Nonstereo fundus	CDR	Mean CDR error: 0.100
86	Depth-discontinuity model	No	138	Stereo fundus	CDR	Mean CDR error: 0.09
87	Sliding window, regression	ORIGA, <sup>84</sup> on request	650	Nonstereo fundus	CDR	Mean CDR error: 0.091
88	AGLAI framework	No	291	Nonstereo fundus	Glaucoma	AUC: 0.73
89	Active contour model, r-bends	No	138	Nonstereo fundus	CDR	Mean CDR error: 0.09
90	Higher-order spectra, texture features	No	60	Nonstereo fundus	Glaucoma	Detection: 61%
91	Active contour, depth reconstruction	No	80	Stereo fundus	CDR, glaucoma	Mean CDR error: 0.110, AUC: 0.90
92	Appearance-based analysis	No	575	Nonstereo fundus	Glaucoma	AUC: 0.88
93	Intensity profiling	No	50	Nonstereo fundus	CDR, Glaucoma	Mean CDR error: 0.14, AUC: 0.87
94	Regional information	No	170	Nonstereo fundus	CDR	Mean CDR error: 0.100
95	Level set	No	104	Nonstereo fundus	CDR	Mean CDR error: 0.089
96	Thresholding, 3-D reconstruction	No	80	Stereo fundus	Glaucoma	AUC: 0.83
97	Hybrid wavelet edges, kinking	No	27	Nonstereo fundus	CDR	Mean CDR error: 0.093
98	ARGALI	No	23	Nonstereo fundus	CDR	Correlation with ground truth: 0.89
99	Deformable model	No	25	Stereo fundus	CDR	Correlation with ground truth: 0.71
100	Pixel features, kNN	No	58	Stereo fundus	CDR	Correlation with ground truth: 0.93
101	Discriminatory analysis	No	NA	Nonstereo fundus	NA	Not provided

AUC, area under the ROC curve.

Therefore, computer-assisted solutions that can either increase the productivity of ophthalmologists' screening or allow trained paramedical personnel to carry out part of the screening themselves will be of significant clinical benefit.

Many ROP-related systems for computer-aided diagnosis have been reported recently, and Table 3 attempts a summary. Retinal Image multiScale Analysis (RISA) provides a semiautomatic tool for the labeling of the skeleton trees, followed by an automatic procedure to measure vessel width and tortuosity and from these derive Plus or Pre-plus diagnosis.<sup>102</sup> The Computer-Aided Image Analysis of the Retina (CAIAR) system semiautomatically identifies the retinal vessels, with provision for manual pixel editing if any vessel is inappropriately represented, and then automatically measures width and tortuosity of each identified vessel.<sup>103</sup> ROPtool semiautomatically traces retinal blood vessels; its reliability in measuring tortuosity and dilation of vessels was assessed in two distinct studies.<sup>104,106</sup> VesselMap (Imedos, Jena, Germany) is a commercial semiautomatic software program developed to analyze vessels in an adult retinal image. It performs the tracking of the

main retinal vessels, providing information only about the vessel diameter, and was used also in ROP images.<sup>105</sup>

No public annotated datasets on ROP seem to be available to date for ARIA system training and validation.

**Age-Related Macular Degeneration.** Age-related macular degeneration (AMD) is a condition of great interest because of its relevance and prevalence. It has, however, attracted less ARIA research than DR, and we do not discuss it in this paper. The reader is referred to several examples of studies.<sup>1,18,61</sup>

**VALIDATION ISSUES AND CHALLENGES**

We now discuss the factors introducing uncertainties in the reference standard (section "Validation in Retinal Image Analysis"). Some are shared with other areas of medical image analysis. Indeed, the very definition of the reference standard varies with a number of factors, and may not reflect the true state of a disease. All this results in assessment variations. As it is unreasonable to pursue an accuracy higher than that of the

TABLE 3. ARIA Literature on ROP at a Glance

Reference No.	DR Task	Method	Public Dataset?	No. of Images	Image Type	Validation Level	Sensitivity	Specificity
102	ROP diagnosis	Semiautomated	No	20	Color fundus	Image	50%-100%	46%-93%
103	Vessel tortuosities and width measurement	Semiautomated	No	10	Color fundus	Image feature	Ground truth correlation: 0.49-0.67 (tortuosities), 0.42 (width)	
104	ROP diagnosis	Semiautomated	No	185	Color fundus	Image	97%	94%
105	Vessel width measurement	Semiautomated	No	20	Color fundus	Image feature	Ground truth correlation: 0.80	
106	Vessel width measurement	Semiautomated	No	30	Color fundus	Image feature	NA	NA

reference standard used, it seems essential to characterize quantitatively the variations of reference standards (see “Techniques” subsection). Much work is still needed to achieve this goal, given the number and nature of the uncertainty sources involved.

Several practical challenges also exist. For instance, it is extremely time-consuming to collect sufficiently large, carefully constructed and annotated reference standard datasets. Time issues are exacerbated when it is necessary to annotate multiple images per eye, for instance with fluorescein angiography sequences or longitudinal studies. Another issue is that the point in the diagnostic process at which “outcome” should be set is not always clear, as factors beyond retinal measurements influence diagnosis. In automatic DR screening, for example, at least two possible end points exist that impact the four types of validation listed in “Techniques”: identifying the presence of DR (refer/no refer decision) and identifying specific lesions from which conclusions are then drawn by the specialist.<sup>107</sup>

### Variation of Expert Judgment

In engineering, reference standards for testing are normally objective measurements from instruments more accurate than the one being tested (see, e.g., camera calibration procedures); in medical image analysis, reference standards are instead built from statistical or explicit consensus among experts. Such judgments vary, in general, with experts (interobserver variations), and, to a lesser extent, over repeated judgments by the same expert (intraobserver variations).

These variations depend at least on experience, task, background, image quality (e.g., this often makes it difficult, even for the expert practitioner, to decide without ambiguity the presence of a druse), and interpretation of the annotation task, although a detailed protocol should minimize variations (see subsection “Annotation Protocols” below). Recent relevant work has been reported by Quellec et al. and Abramoff<sup>2,20</sup> on the maximum meaningful performance achievable with automatic binary decision systems, given the characteristics of the reference standard obtained from clinicians. The study focused on the ROC area under the curve as the evaluation index. The authors ran tests with two ARIA systems for DR detection, 500 images, and the reference standard obtained by three experts. They concluded, interestingly, that meaningful performance measured against a single expert could not be improved (on the dataset used), whereas it could be improved significantly when compared against a committee of experts.

Hubschman et al.<sup>108</sup> quantified the interobserver difference in an ischemia grading task with branch retinal vein occlusion. A single ultra-wide-field fluorescein angiogram image was segmented by four retina specialists into regions belonging to four different levels of retinal perfusion (normally perfused, partially perfused, nonperfused, insufficient quality). The cardiac cycle can also have an influence on vessel caliber measurements (see subsection “Physiological Short-Term Changes in Time”).

Intuitively, uncertainty can be reduced by increasing the number of experts and the number of annotations per expert. There is, however, little consensus on how to proceed statistically when such data are available, and expert numbers are usually small (two to five in most ARIA papers). Relevant techniques have been mentioned in “Techniques.”

### Annotation Protocols

Procedures used to take photographs represent another source of variability. For example, if the eye is not positioned in the

same location, the vessels may be captured at slightly different angles, resulting possibly in different measurements.

As noted in “Techniques,” annotating specific image elements, like circling regions or tracing vessels on a computer screen, is a task that doctors do not normally perform in clinical practice. To maximize annotation accuracy as well as relevance to translation, it seems desirable to align validation tasks with those that doctors specialize in. Relevant research has been reported recently by Quellec et al.,<sup>20</sup> who designed graphical user interfaces (GUIs) to automatically detect elements that catch the attention of clinicians in their daily clinical practice. This avoids requesting clinicians to explicitly annotate anatomical structures, a task they have not been trained for. Usage logs (zoom, magnifying glass, and the like) are used to train weakly supervised lesion detectors and validate their outputs.

Datasets used for high-profile clinical studies, such as AREDS (see Appendix A), were annotated at national grading centers using established protocols. As the intent was to evaluate the efficacy of drugs (for AREDS, by the National Institutes of Health/U.S. Department of Agriculture), it should be believed that the grading protocol was highly refined and scrutinized, and should be trusted to a higher extent than those of datasets developed at individual institutions for testing a specific algorithm. However, datasets not considering ARIA validation might not provide ARIA-relevant data, for example pixel-level delineations allowing ROC analysis.

### Physiological Short-Term Changes in Time

Taking photographs at random instants in the pulse cycle may result in unrecognized variations in the measurements of retinal vessel diameters, however technically sophisticated, both among subjects and over time in the same individual. A few studies have investigated this in detail,<sup>109-113</sup> and some conclusions appear to be conflicting. It has been reported that the maximum variation at different points in the pulse cycle ranged from 4.3% to 4.8% for major retinal venules and from 3.1% to 3.9% for major retinal arterioles. Venular diameter was smallest in early systole, increasing to a maximum level in early diastole and decreasing thenceforth. The arteriole diameter peaked slightly earlier. In another investigation of 10 volunteers, it was shown that a summary measure of the retinal venule diameters and the arteriolar diameters change at different points in the cardiac cycle. Across the cardiac cycle, the central retinal venular equivalent (CRVE) changed by 3.1% and the central retinal arteriolar equivalent (CRAE) changed by 4.3%. However, recent work by Kumar et al.<sup>113</sup> has shown that there is no significant change in the average of the width of six large vessels in the region often chosen for arteriole-to-venule ratio (AVR) estimation. As quantifying vascular changes is a primary ARIA concern,<sup>112</sup> there is a need for more extensive studies and test sets with repeated images of the same eye over time. Gating the eye fundus camera with the electrocardiogram has also been proposed. Moret et al.<sup>114</sup> have measured changes to the vessel shape and size, also evidenced from the use of dynamic vessel analyzers (DVA) for disease diagnosis.<sup>115</sup> DVA can provide temporal resolutions above 20 frames/s but comes with limits on spatial resolution and methodology, and the instrumentation is suitable only for highly specialized research facilities.

### Different Imaging Instruments

Because of the nature of retinal imaging, there is usually a high level of customization for each modality. Algorithms used for the segmentation of the optic nerve head in a confocal scanning technique may not be applicable to retinal images



without some level of customization. Even without the same class of machines, the variation of instruments can have a large effect on algorithm processes. In retinal fundus imaging, resolution variations can have a large effect on algorithm performance, and resolution requirements depend on the task at hand. For example, a lower resolution may be acceptable for optic nerve head segmentation, but not for estimating the tortuosity of retinal neovascularization. The selection of FOV and imaging field can also affect performance, particularly when one is relying on or assuming the visibility of retinal landmarks (OD, macula, arcades). The color calibration model used in the camera CCDs for digital capture can also affect algorithm performance, possibly causing images of the same retina taken by different devices to appear significantly different.

### Image Quality

Image quality was discussed earlier. We reiterate that image quality depends on, among other factors, instrument characteristics, acquisition procedure, and target conditions, and that capturing quality definitions applied by experts for implementation in ARIA systems is difficult; in general, images considered viable for clinical analysis may not produce good results with ARIA systems.

### Datasets

The key observation is that different datasets may lead to somewhat inconsistent performance assessments, as preparation protocols may differ. For instance, the best-known public retinal datasets are probably DRIVE<sup>116</sup> and STARE.<sup>117</sup> Soares et al.<sup>47</sup> compared binary vessel masks from both datasets with masks obtained from their algorithm. Accuracy differences were noted between the two datasets, associated with different segmentation methods and differences in the extent and detail of the manual tracing provided by the experts. However, other groups have reported diverse methods of vessel segmentation and identification of proliferative retinopathy,<sup>48</sup> and a comparison of these methods has not shown large differences in accuracy with images from the STARE dataset.<sup>118</sup>

### Task, Previous Knowledge, Condition

Expert judgment and annotations can be different for the same image and target measure depending on the task a clinician has in mind. For instance, when asked to determine the width of a retinal vessel from a fundus images, a doctor thinking surgically might try to keep at a distance from the vessel, hence overestimating width. If the task does affect annotations, it seems advisable to group reference annotations also by clinical task and to specify annotation protocols accordingly. At the moment, some public repositories group images by conditions, for example MESSIDOR for DR (see Appendix A).

### Patient Characterization: Metadata

The growing volume of electronic images and datasets potentially available to researchers makes it challenging to mine the rich information embedded in the data. This obviously includes image properties, but also *contextual information*, that is, clinical metadata that may be relevant for the disease incidence and for organizing consistent validation datasets. Contextual patient characterization data include ethnicity, age, sex, medical data (hypertension, diabetes, heart diseases), lifestyle factors (e.g., smoking), systemic data (body mass index, cholesterol level, blood

pressure, and so on), ocular data (refractive error, lens opacity, and so on), comorbid diseases, and in general all data normally considered in clinical studies.

These factors are rarely discussed for data used in validating ARIA algorithms, in part because their effect on retinal images, and consequently on ARIA results, is the object of current international investigation. Content-based approaches may have particular relevance in utilizing contextual information together with image analysis to assign risk profiles to patients in a screening environment. Quellec et al.<sup>119,120</sup> have reported work on the inclusion of demographic and biological data in an image-based DR severity scale, concluding that this inclusion leads to significant improvement in classification performance. A larger clinical study aimed at confirming these findings is currently being performed on a dataset of 25,702 examination records from the Ophdiat screening network (provided in the public domain at <http://reseau-ophdiat.aphp.fr/index.html>).<sup>3</sup>

### Human in the Loop

A final consideration is that some ARIA algorithms are semiautomatic and involve deliberately human intervention, a paradigm known in robotics as “human in the loop.” This complicates the objective evaluation of performance, as it seems to require decoupling human and automatic contributions. Specialized techniques apparently have not yet been considered in the ARIA literature.

## DISCUSSION AND RECOMMENDATIONS

The ideal way to identify effective ARIA algorithms is to compare solutions proposed in the literature on an equal footing and with datasets recognized as meaningful by a representative cross section of the clinical community. This comparison is best achieved by the creation of common, accessible, and representative datasets including automated tools to run submitted algorithms on the data provided. Other areas of image-processing research have produced such repositories, for instance stereo and multiple-image analysis.<sup>121</sup> An ARIA move in this direction is the recent Diabetic Retinopathy Online Challenge.<sup>30</sup> Encouragingly, a variety of public datasets have appeared in recent years (see Appendix A). Most are still limited collections of images generated by individual sites, or contain limited annotations, or both. There is currently no coordinated consensus in the community on how to structure such datasets or what information to include, and this paper is meant to provoke thoughts toward that end.

So far, neither ARIA datasets nor epidemiological studies based on large populations have systematically taken into account the above issues. Furthermore, as images are often acquired independent of outcomes and participant characteristics, random variability may tend to underestimate the true associations found, for example between retinal vascular calibers and cardiovascular diseases. Future studies need to overcome these sources of variability before retinal features could be used as a more precise biomarker.

ARIA repositories of test data (not including alternative, promising validation paradigms like indirect methods) should, ideally,

- *Be created collaboratively* by consortia of international groups in order to achieve large data volumes and multiple annotators, to reduce opinion bias, to guarantee international visibility and credibility, and to ultimately generate useful results for clinicians.

- *Be easily accessible*, ideally via Web sites from which data could be downloaded following a suitable registration proce-



dure including legal disclaimers, no-redistribution clauses, acknowledgments needed for use, and so forth.

- *Be regularly maintained*, ideally by a consortium of international groups monitoring distribution and guaranteeing maintenance of data and annotations.

- *Be large in size*, and wherever possible dimensioned statistically to maximize power; tentatively, the minimum order of magnitude should be thousands of images. Large data repositories should be standardized, patient-friendly imaging protocols allowing large populations to be imaged effectively; for example, single 45° fovea-centered fields, or two 45° fields centered on the fovea and OD, respectively.<sup>122</sup>

- *Include metadata*, that is, non-image data characterizing imaging instruments, patients, and disease (subsection “Patient Characteristics: Metadata”).

- *Include automated tools for running software on the data*, as done on the Middlebury stereo site,<sup>121</sup> in which executable code is loaded and run on the site, and performance assessed in terms of predefined measures that are displayed in tabular form.

- *Organized by outcome*, which depends on the task at hand; for example, refer/do not refer in screening tasks, levels of tortuosity or width measurement in feature-oriented tasks. An image set could be used for multiple outcomes by providing annotations for some or all the points listed under “Techniques.”

- *Include image annotations*, providing the standard reference for comparison for the outcome stated (see previous point), preferably provided by as many clinicians as possible (ideally from different sites to eliminate possible opinion bias) to estimate interobserver variability, including arbitrated annotations; each expert should ideally annotate the dataset multiple times to estimate intraobserver variability (see also last paragraph of “Techniques” section for techniques managing variations in expert judgment). Uncertainty levels declared or agreed on by the annotators should also be considered. A surrogate of direct measurement yielding “true” gold standard could be the development of phantoms of the eye, including the retinal vasculature and pumps to simulate the blood flow, which could enable the calibration of ARIA vessel measurement methods. But given the current impossibility, with ARIA tasks, of obtaining ground truth measured by independent, highly accurate instruments, the only way to estimate “how good” an algorithm is to compare its output with expert judgment.

Creating such repositories for ARIA algorithms poses important challenges. The most obvious one is the sheer complexity of the task, as presented above. Acquiring images, generating the necessary annotations, and preparing the data for public use take time and impose significant costs. Governance and ethical issues, which vary internationally, may complicate further the release of clinical data for public research. A further point is the effort required for administration and maintenance, considering for instance that the obsolescence of imaging instruments limits the life of a dataset. It is arguable that retinal scans around 600 × 600 pixels are increasingly obsolete given the availability of much higher-resolution instruments.

Ultimately, outcome-oriented measures are required. It would be of little applicative interest to develop ever more accurate ARIA techniques if they could not be used to improve clinical outcomes. Outcome measures should be considered in a public health context, taking into account health economics, risks, and the impact of changes to services using automated algorithms. However, during the development of algorithms, it is helpful to compare the isolated performance of modules that will eventually become components in a larger system so that the most effective methods can be identified. This requires the

provision of comparative datasets with reference standard measurements of features that may ultimately prove diagnostic. These features must be selected based on expert clinical advice.

### Acknowledgments

Supported by the National Institutes of Health Grants R01 EY018853 and R01 EY019112 (MA), and the Department of Veterans Affairs I01 CX000119 (MA).

The authors thank the additional authors listed in Appendix B.

Disclosure: **E. Trucco**, None; **A. Ruggeri**, None; **T. Karnowski**, None; **L. Giancardo**, None; **E. Chaum**, None; **J.P. Hubschman**, None; **B. al-Diri**, None; **C.Y. Cheung**, None; **D. Wong**, None; **M. Abramoff**, None; **G. Lim**, None; **D. Kumar**, None; **P. Burlina**, None; **N.M. Bressler**, None; **H.F. Jelinek**, None; **F. Meriaudeau**, None; **G. Quellec**, None; **T. MacGillivray**, None; **B. Dhillon**, None

### References

1. Abramoff MD, Garvin M, Sonka M. Retinal image analysis: a review. *IEEE Rev Biomed Eng.* 2010;3:169–208.
2. Abramoff MD, Reinhardt JM, Russell SR, et al. Automated early detection of diabetic retinopathy. *Ophthalmology.* 2010;117:1147–1154.
3. Chabouis A, Berdugo M, Meas T, et al. Benefits of Ophdiat, a telemedical network to screen for diabetic retinopathy: a retrospective study in five reference hospital centres. *Diabetes Metab.* 2009;35:228–232.
4. Ding J, Patton N, Deary I, et al. Retinal vascular abnormalities and cognitive dysfunction: a systematic review. *Br J Ophthalmol.* 2008;92:1017–2005.
5. Patton N, Aslam T, McGillivray T, et al. Retinal image analysis: concepts, application and potential. *Progr Retin Eye Res.* 2006;25:99–127.
6. Mitchell B, Koo J, Iordachita M, et al. Development and application of a new steady-hand manipulator for retinal surgery. *IEEE Int Conf Robot Autom.* 2007:623–629.
7. Taylor R, Jensen P, Whitcomb L, et al. A steady-hand robotic system for microsurgical augmentation. *Int J Rob Res.* 1999; 18:1201–1210.
8. Balicki M, Han JH, Iordachita I, et al. Single fiber optical coherence tomography microsurgical instruments for computer and robot-assisted retinal surgery. *Med Image Comput Assist Interv.* 2009;12:108–115.
9. Camarillo DB, Krummel TM, Salisbury JK, et al. Robotic technology in surgery: past, present, and future. *Am J Surg.* 2004;188:2–15.
10. Fleming I, Balicki M, Koo J, et al. Cooperative robot assistant for retinal microsurgery. *Med Image Comput Assist Interv.* 2008;11:543–550.
11. Wild S, Sicree R, Roglic G, King H, Green A. Global prevalence of diabetes. *Diabetes Care.* 2004;27:1047–1053.
12. Lairson D, Pugh JA, Kapadia AS, Lorimor RJ, Jacobson J, Velez R. Cost-effectiveness of alternative methods for diabetic retinopathy screening. *Diabetes Care.* 1992;15:1369–1377.
13. Li R, Zhang P, Barker LE, Chowdhury FM, Zhang X. Cost-effectiveness of interventions to prevent and control diabetes mellitus: a systematic review. *Diabetes Care.* 2010;33:1872–1894.
14. Quellec G, Abramoff M. Estimating maximum measurable performance from automatic decision systems from the characteristics of the reference standard. *IEEE Trans Biomed Eng.* In press.

15. Commonwick O, Warfield, Simon D. A continuous STAPLE for scalar, vector, and tensor images: an application to DTI analysis. *IEEE Trans Med Imaging*. 2009;28:838-846.
16. Commonwick O, Warfield, Simon D. Estimation of inferential uncertainty in assessing expert segmentation performance from STAPLE. *IEEE Trans Med Imaging*. 2010;29:771-780.
17. Rutjes AW, Reitsma JB, Coomarasamy A, Khan KS, Bossuyt PM. Evaluation of diagnostic tests when there is no gold standard. A review of methods. *Health Technol Assess*. 2007;11:iii, ix-51.
18. Kankanahalli S, Burlina P, Wolfson Y, et al. Automated classification of severity of age-related macular degeneration from fundus photographs. *Invest Ophthalmol Vis Sci*. 2013;54:1789-1796.
19. Hammer DX, Ferguson RD, Ustun TE, Bigelow CE, Iftimia NV, Webb RH. Line-scanning laser ophthalmoscope. *J Biomed Opt*. 2006;11:041126.
20. Quellec G, Lamard M, Cazuguel G, et al. Weakly supervised classification of medical images. *Proc IEEE ISBI Int Symp on Biomed Imaging*. 2012:110-113.
21. Quellec G, Lamard M, Cazuguel G, et al. Automated assessment of diabetic retinopathy severity using content-based image retrieval in multimodal fundus photographs. *Invest Ophthalmol Vis Sci*. 2011;52:8342-8348.
22. Yogesan K, Constable IJ, Barry CJ, et al. Evaluation of a portable fundus camera for use in the teleophthalmologic diagnosis of glaucoma. *J Glaucoma*. 1999;8:297.
23. Hunter A, Lowell JA, Habib M, et al. An automated retinal image quality grading algorithm. In: *Proceedings from the IEEE EMBC International Symposium on Engineering in Medicine and Biology*. 2011:5955-5958.
24. Niemeijer M, Abramoff MD, van Ginneken B. Image structure clustering for image quality verification of color retina images in diabetic retinopathy screening. *Med Image Anal*. 2006;10:888-898.
25. Usher D, Himaga M, Dumskyj M. Automated assessment of digital fundus image quality using detected vessel area. In: *Proceedings of Medical Image Understanding and Analysis*. Liverpool: British Machine Vision Association; 2003:81-84.
26. Lalonde M, Gagnon L, Boucher M. Automatic visual quality assessment in optical fundus images. In: *Proceedings of Vision Interface*. Ottawa;2001:259-264.
27. Lee S, Wang Y. Automatic retinal image quality assessment and enhancement. In: *Proc SPIE: Image Processing*. 1999;1581-1590.
28. Davis H, Russell S, Barriga E, Abramoff M, Soliz P. Vision-based, real-time retinal image quality assessment. In: *Proceedings of 22nd IEEE International Symposium on Computer-Based Medical Systems*. 2009. Doi:978-1-4244-4878-4/09/.
29. Giancardo L, Abramoff M, Chaum E, et al. Elliptical local vessel density: a fast and robust quality metric for retinal images. In: *Proceedings from the International Conference of the IEEE EMBC Engineering in Medicine and Biology Society*. Vancouver; 2008:3534-3537.
30. Lu CK, Tang TB, Laude A, Deary IJ, Dhillon B, Murray AF. Quantification of parapapillary atrophy and optic disc. *Invest Ophthalmol Vis Sci*. 2011;52:4671-4677.
31. Kawasaki R, Che Azemin MZ, Kumar DK, et al. Fractal dimension of the retinal vasculature and risk of stroke: a nested case-control study. *Neurology*. 2011;76:1766-1767.
32. Azemin MZ, Kumar DK, Wong TY, Kawasaki R, Mitchell P, Wang JJ. Robust methodology for fractal analysis of the retinal vasculature. *IEEE Trans Med Imaging*. 2011;30:243-249.
33. Azemin MZ, Kumar DK, Wong TY, et al. Age-related rarefaction in the fractal dimension of retinal vessel. *Neurobiol Aging*. 2012;33:194.e1-194.e4.
34. Al-Diri B, Hunter A. Automated measurements of retinal bifurcations. In: *Medical Physics and Biomedical Engineering*. Munich; 2009.
35. Al-Diri B, Hunter A, Steel D, Habib M. Automated analysis of retinal vascular network connectivity. *Comput Med Imaging Graph*. 2010;34:462-470.
36. Al-Diri B, Hunter A, Steel D. An active contour model for segmenting and measuring retinal vessels. *IEEE Trans Med Imaging*. 2009;28:1488-1497.
37. Congdon NG, Friedman DS, Lietman T. Important causes of visual impairment in the world today. *JAMA*. 2003;290:2057-2060.
38. Fong DS, Aiello LP, Ferris FL, Klein R. Diabetic retinopathy. *Diabetes Care*. 2004;27:2540-2553.
39. World Health Organization. Diabetic retinopathy. Available at: <http://www.who.int/blindness/causes/priority/en/index6.html>. Accessed November 1, 2010.
40. Saaddine JB, Honeycutt AA, Narayan KMV, Zhang XZ, Klein R, Boyle JP. Projection of diabetic retinopathy and other major eye diseases among people with diabetes mellitus: United States, 2005-2050. *Arch Ophthalmol*. 2008;126:1740-1747.
41. Fleming AD, Philip S, Goatman KA, et al. Automated detection of exudates for diabetic retinopathy screening. *Phys Med Biol*. 2006;52:7385-7396.
42. Olson JA, Sharp PF, Fleming A, et al. Evaluation of a system for automatic detection of diabetic retinopathy from color fundus photographs in a large population of patients with diabetes. *Diabetes Care*. 2008;31:63-64.
43. Philip S, Fleming AD, Goatman KA, et al. The efficacy of automated disease/no disease grading for diabetic retinopathy in a systematic screening programme. *Br J Ophthalmol*. 2007;91:1512-1517.
44. Giancardo L, Meridaudeau F, Karnowski TP, et al. Exudate-based diabetic macular edema detection in fundus images using publicly available datasets. *Med Image Anal*. 2012;16:216-226.
45. Cree MJ, Olson JA, McHardy KC, et al. Automated microaneurysms detection. In: *IEEE International Conference on Image Processing*. Lausanne, Switzerland: IEEE Press; 1996.
46. Ege BM, Hejlesen OK, Larsen OV, et al. Screening for diabetic retinopathy using computer based image analysis and statistical classification. *Comput Methods Programs Biomed*. 2000;62:165-175.
47. Soares JVB, Leandro JGG, Cesar Júnior RM, et al. Retinal vessel segmentation using the 2-D Gabor wavelet and supervised classification. *IEEE Trans Med Imaging*. 2006;25:1214-1222.
48. Niemeijer M, van Ginneken B, Russell SR, Suttrop-Schulten MSA, Abramoff MD. Automated detection and differentiation of drusen, exudates, and cotton-wool spots in digital color fundus photographs for diabetic retinopathy diagnosis. *Invest Ophthalmol Vis Sci*. 2007;48:2260-2267.
49. Niemeijer M, Ginneken B, Cree MJ, et al. Retinopathy online challenge: automatic detection of microaneurysms in digital color fundus photographs. *IEEE Trans Med Imaging*. 2010;29:185-195.
50. Niemeijer M, van Ginneken B, Staal J, Suttrop-Schulten MSA, Abramoff MD. Automatic detection of red lesions in digital color fundus photographs. *IEEE Trans Med Imaging*. 2005;24:584-592.
51. Giancardo L, Meridaudeau F, Karnowski TP, Li Y, Tobin KW Jr, Chaum E. Microaneurysm detection with Radon transform-based classification on retina images. *Conf Proc IEEE Eng Med Biol Soc*. 2011;2011:5939-5942.

52. Mizutani A, Muramatsu C, Hatanaka Y, et al. Automated microaneurysm detection method based on double ring filter in retinal fundus images. In: *Proc SPIE: Medical Imaging*. 2009;7260.
53. Quéllec G, Lamard M, Josselin PM, Cazuguel G, Cochener B, Roux C. Optimal wavelet transform for the detection of microaneurysms in retina photographs. *IEEE Trans Med Imaging*. 2008;27:1230-1241.
54. Walter T, Massin P, Erginay A, Ordonez R, Jeulin C, Klein JC. Automatic detection of microaneurysms in color fundus images. *Med Image Anal*. 2007;11:555-566.
55. Huang K, Yan M, Aviyente S. Edge-directed inference for microaneurysms detection in digital fundus images. In: *SPIE Medical Imaging: Image Processing*. 2007;6512.
56. Fleming AD, Philip S, Goatman KA, Olson JA, Sharp PF. Automated microaneurysm detection using local contrast normalization and local vessel detection. *IEEE Trans Medical Imaging*. 2006;25:1223-1232.
57. Quéllec G, Lamard M, Josselin PM, et al. Detection of lesions in retina photographs based on the wavelet transform. In: *Proceedings from the IEEE EMBC International Conference on Engineering in Medicine and Biology Society*. New York; 2006.
58. Pallawala PMDS, Hsu W, Lee ML, et al. Automated microaneurysm segmentation and detection using generalized eigenvectors. In: *Proceedings from the IEEE Workshop on Application of Computer Vision (WACV)*. Breckenridge, CO; 2005.
59. Serrano C, Acha B, Revuelto S. 2D adaptive filtering and region growing algorithm for the detection of microaneurysms. In: *SPIE Medical Imaging: Image Processing*. 2004; 5370:1924-1931.
60. Larsen M, Godt J, Larsen N, et al. Automated detection of fundus photographic red lesions in diabetic retinopathy. *Invest Ophthalmol Vis Sci*. 2003;44:671-766.
61. Rapantzikos K, Zervakis M, Balas K. Detection and segmentation of drusen deposits on human retina: potential in the diagnosis of age-related macular degeneration. *Med Image Anal*. 2003;7:95-108.
62. Sinthanayothin C, Boyce JF, Williamson TH, et al. Automated detection of diabetic retinopathy on digital fundus images. *Diabetic Med*. 2002;19:105-112.
63. Hsu W, Pallawala PMDS, Lee ML, Kah-Guan AE. The role of domain knowledge in the detection of retinal hard exudates. In: *IEEE Conference on Computer Vision and Pattern Recognition (CVPR)*. 2001:246-251.
64. Yang G, Gagnon L, Wang S, Boucher MC. Algorithm for detecting microaneurysms in low-resolution color retinal images. In: *Proceedings of Vision Interface*. 2001:265-271.
65. Wang H, Hsu W, Goh KG, Lee ML. An effective approach to detect lesions in color retinal images. In: *Proceedings from the IEEE Conference on Computer Vision and Pattern Recognition (CVPR)*. Hilton Head, SC; 2000:2181-2186.
66. Hipwell JH, Strachan F, Olson JA, et al. Automated detection of microaneurysms in digital red-free photographs: a diabetic retinopathy screening tool. *Diabetic Med*. 2000;17:588-594.
67. Lee SC, Wang Y, Lee ET. Computer algorithm for automated detection and quantification of microaneurysms and hemorrhages (HMAs) in color retinal images. In: *SPIE Medical Imaging: Image Perception and Performance*. 1999;3663.
68. Cree MJ, Olson JA, McHardy KC, et al. A fully automated comparative microaneurysm digital detection system. *Eye*. 1997;11:622-629.
69. Gardner GG, Keating D, Williamson TH, Elliott AT. Automatic detection of diabetic retinopathy using an artificial neural network: a screening tool. *Br J Ophthalmol*. 1996;80:940.
70. Spencer T, Phillips RP, Sharp PF, Forrester JV. Automated detection and quantification of microaneurysms in fluorescein angiograms. *Graefes Arch Clin Exp Ophthalmol*. 1992; 230:36-41.
71. Abramoff MD, Alward WLM, Greenlee EC, et al. Automated segmentation of the optic disc from stereo color photographs using physiologically plausible features. *Invest Ophthalmol Vis Sci*. 2007;48:1665-1673.
72. Muramatsu C, Nakagawa T, Sawada A, et al. Determination of cup-to-disc ratio of optical nerve head for diagnosis of glaucoma on stereo retinal fundus image pairs. In: *SPIE*. 2009:096009.
73. Wong DWK, Liu J, Lim JH, et al. Level-set based automatic cup-to-disc ratio determination using retinal fundus images in Argali. In: *Proceedings from the 30th Annual International Conference of the IEEE Engineering in Medicine and Biology Society*. 2008;1:2266-2269.
74. Wong DWK, Liu J, Lim JH, et al. Intelligent fusion of cup-to-disc ratio determination methods for glaucoma detection in ARGALI. In: *Proceedings from the IEEE EMBC International Conference on Engineering in Medicine and Biology*. Minneapolis; 2009;1-20:5777-5780.
75. Liu J, Wong DWK, Lim JH, et al. ARGALI: an automatic cup-to-disc ratio measurement system for glaucoma analysis using level-set image processing. In: *Proceedings from the 13th IEEE EMBC International Conference on Biomedical Engineering*. Minneapolis; 2009;23:559-562.
76. Joshi GD, Sivaswamy J, Karan K, Krishnadas SR. Optic disk and cup boundary detection using regional information. In: *Proceedings from the 7th IEEE EMBC International Conference on Biomedical Imaging*. Buenos Aires; 2010: 948-951.
77. Hatanaka Y, Noudo A, Muramatsu C, et al. Vertical cup-to-disc ratio measurement for diagnosis of glaucoma on fundus images. In: *Proc SPIE: Medical Imaging: Computer-Aided Diagnosis*. 2010;7624.
78. Muramatsu C, Nakagawa T, Sawada A, et al. Automated segmentation of optic disc region on retinal fundus photographs: comparison of contour modeling and pixel classification methods. *Comput Methods Programs Biomed*. 2011;101:23-32.
79. Bock R, Meier J, Nyul LG, Hornegger J, Michelson G. Glaucoma risk index: automated glaucoma detection from color fundus images. *Med Image Anal*. 2010;14:471-481.
80. Liu J, Wong WK, Damon, Zhang, Z, et al. AGLAIA system and development status. Presented at the Asia Association for Research in Vision and Ophthalmology; 2011.
81. Liu J, Zhang Z, Wong WK, et al. Automatic glaucoma diagnosis and its genetic association study through Medical Image Informatics (AGLAIA). In: *Proceedings from the DICOM International Conference and Seminar*. Rio de Janeiro; 2010.
82. Liu J, Wong WK, Tan NM, et al. AGLAIA system architecture for glaucoma diagnosis. In: *Proceedings from the Asia Pacific Signal And Information Processing Association Summit and Conference*. Singapore; 2010.
83. Xu Y, Liu J, Lin S, et al. Efficient optic cup detection from intra-image learning with retinal structure priors. In: *Proceedings from the MICCAI International Conference on Medical Image Computing and Computer-Assisted Intervention (MICCAI)*. 2012;15:58-65.
84. Zhang Z, Yin F, et al. ORIGA(-light): an online retinal fundus image database for glaucoma analysis and research. In: *Proceedings from the Intern IEEE EMBC Conf on Engineering in Medicine and Biology*. Buenos Aires; 2010:3065-3068.



85. Yin F, Liu J, Wong DWK, et al. Automated segmentation of optic disc and optic cup in fundus images for glaucoma diagnosis. In: *Proceedings from the International IEEE CBMS Symposium on Computer-Based Medical Systems*. Rome; 2012:1-6.
86. Joshi G, Sivaswamy J, Krishnadas SR. Depth discontinuity-based cup segmentation from multiview color retinal images. *IEEE Trans Biomed Eng*. 2012;59:1523-1531.
87. Xu Y, Xu D, et al. Sliding window and regression based cup detection in digital fundus images for glaucoma diagnosis. International Conference on Medical Image Computing and Computer-Assisted Intervention (MICCAI). 2011:1-8.
88. Liu J, Yin FS, Wong DKW, et al. Automatic glaucoma diagnosis from fundus images. In: *Proc Intl IEEE EMBC Conf on Engineering in Medicine and Biology*. Boston; 2011:3383-3386.
89. Joshi GD, Sivaswamy J, Krishnadas SR. Optic disk and cup segmentation from monocular color retinal images for glaucoma assessment. *IEEE Trans Med Imaging*. 2011;30:1192-1205.
90. Acharya UR, Dua S, et al. Automated diagnosis of glaucoma using texture and higher order spectra features. *IEEE Trans Inf Technol Biomed*. 2011;15:449-455.
91. Muramatsu C, Nakagawa T, Sawada A, et al. Automated determination of cup-to-disc ratio for classification of glaucomatous and normal eyes on stereo retinal fundus images. *J Biomed Opt*. 2011;16:3387-3390.
92. Bock R, Meier J, et al. Glaucoma risk index: automated glaucoma detection from color fundus images. *Med Image Anal*. 2010;14:471-481.
93. Hatanaka Y, Noudo A, Muramatsu C, et al. Vertical cup-to-disc ratio measurement for diagnosis of glaucoma on fundus images. In: *Proc SPIE: Medical Imaging: Computer-Aided Diagnosis*. Vol 7624. 2010:76243C.
94. Joshi GD, Sivaswamy J, Krishnadas, SR, et al. Optic disk and cup boundary detection using regional information. In: *Proceedings from the IEEE International Symposium on Biomedical Imaging*. Buenos Aires; 2010:948-951.
95. Wong DWK, Liu J, Zhang Z, et al. Level-set based automatic cup-to-disc ratio determination using retinal fundus images in ARGALI. In: *Proceedings from the IEEE EMBC International Conference on Engineering in Medicine and Biology*. Vancouver; 2008:2266-2269.
96. Muramatsu C, Nakagawa T, Sawada A, et al. Determination of cup-to-disc ratio of optical nerve head for diagnosis of glaucoma on stereo retinal fundus image pairs. In: *Proc SPIE: Medical Imaging: Computer-Aided Diagnosis*. 2009;7260:1:5777-5780.
97. Wong DWK, Liu J, et al. Automated detection of kinks from blood vessels for optic cup segmentation in retinal images. In: *Proc SPIE: Medical Imaging: Computer-Aided Diagnosis*. 2009;7260:72601J.
98. Liu J, Wong DWK, Lim JH, et al. Optic cup and disk extraction from retinal fundus images for determination of cup-to-disc ratio. In: *Proceedings from the IEEE International Conference on Industrial Electronics and Automation*. Singapore: 2008:1828-1832.
99. Xu J, Chutatape O. Optic disk feature extraction via modified deformable model technique for glaucoma analysis. *Pattern Recognit*. 2007;40:2063-2076.
100. Abramoff MD, Alward WLM, Greenlee EC, et al. Automated segmentation of the optic disc from stereo color photographs using physiologically plausible features. *Invest Ophthalmol Vis Sci*. 2007;48:1665-1673.
101. Inoue N, Yanashima K, et al. Development of a simple diagnostic method for the glaucoma using ocular fundus pictures. In: *Proceedings of the IEEE EMBC International Conference on Engineering in Medicine and Biology*. Chicago: [PUBLISHER]; 2005:3355-3358.
102. Koreen S, Gelman R, Martinez-Perez ME, et al. Evaluation of a computer-based system for plus disease diagnosis in retinopathy of prematurity. *Ophthalmology*. 2007;114:59-67.
103. Wilson CM, Cocker KD, Moseley MJ, et al. Computerized analysis of retinal vessel width and tortuosity in premature infants. *Invest Ophthalmol Vis Sci*. 2008;49:3577-3585.
104. Wallace DK, Freedman SF, Zhao Z, et al. Accuracy of ROPTool vs individual examiners in assessing retinal vascular tortuosity. *Arch Ophthalmol*. 2007;125:1523-1530.
105. Wallace DK, Freedman SF, Zhao Z. A pilot study using ROPTool to measure retinal vascular dilation. *Retina*. 2009;29:1182-1187.
106. Johnson KS, Mills MD, Karp KA, et al. Semiautomated analysis of retinal vessel diameter in retinopathy of prematurity patients with and without plus disease. *Am J Ophthalmol*. 2007;143:723-725.
107. Jelinek HF, Cree MJ, eds. *Automated Detection of Retinal Pathology*. Boca Raton, FL: CRC Press; 2009.
108. Hubschman JP, Trucco E, et al. Assessment of retinal non-perfusion in vascular retinal diseases using single versus summarized ultra wide-field fluorescein angiography images. In: *Proceedings for the Association for Research in Vision and Ophthalmology*. Fort Lauderdale, FL: Association for Research in Vision and Ophthalmology; May 6-9, 2012.
109. Reshef DS. *Evaluation of Generalized Arteriolar Narrowing Expressed as Central Retinal Artery/Vein Equivalents Ratio (CRAVER) Using ECG Synchronized Retinal Photography*. Baltimore, MD: Johns Hopkins University; 1999. Thesis.
110. Chen HC, Patel V, Wiek J, Rassam SM, Kohner EM. Vessel diameter changes during the cardiac cycle. *Eye*. 1994;8:97-103.
111. Knudtson MD, Klein BEK, Klein R, et al. Variation associated with measurement of retinal vessel diameters at different points in the pulse cycle. *Br J Ophthalmol*. 2004;88:57-61.
112. Wong TY, Islam F, Klein R, et al. Retinal vascular caliber, cardiovascular risk factors, and inflammation: the multi-ethnic study of atherosclerosis (MESA). *Invest Ophthalmol Vis Sci*. 2006;47:2341.
113. Kumar DK, Hao H, Aliahmad B, Wong T, Kawasaki R. Does retinal vascular geometry vary with cardiac cycle? *Invest Ophthalmol Vis Sci*. 2012;53:5799-5805.
114. Moret F, Poloschek C, Lagrèze W, Bach M. Visualization of fundus vessel pulsation using principal component analysis. *Invest Ophthalmol Vis Sci*. 2011;52:5457-5464.
115. Nguyen TT, Kawasaki R, Wang JJ, et al. Flicker lights induced retinal vasodilation in diabetes and diabetic retinopathy. *Diabetes Care*. 2009;32:2075.
116. Staal J, Abramoff MD, et al. Ridge-based vessel segmentation in color images of the retina. *IEEE Trans Med Imaging*. 2004;23:501-509.
117. Hoover A, Kouznetsova V, Goldbaum M. Locating blood vessels in retinal images by piecewise threshold probing of a matched filter response. *IEEE Trans Med Imaging*. 2000;19:203-210.
118. Cree MJ, Leandro JGG, Soares JVB, et al. Comparison of various methods to delineate blood vessels in retinal images. In: *Proceedings of the 16th Australian Institute of Physics Congress*. Canberra: Australia Institute of Physics; 2005.
119. Queller G, Lamard M, Cazuguel G, Roux C, Cochener B. Case retrieval in medical datasets by fusing heterogeneous information. *IEEE Trans Med Imaging*. 2011;30:108-118.
120. Queller G, Lamard M, et al. Automated assessment of diabetic retinopathy severity using content-based image retrieval in

multimodal fundus photographs. *Invest Ophthalmol Vis Sci*. 2011;52:8342–8348.

121. Scharstein D, Szeliski R. A taxonomy and evaluation of dense two-frame stereo correspondence algorithms. *Int J Comput Vis*. 2002;47:7–42.
122. Aldington SJ, Kohner EM, Meuer S, Klein R, Sjølie AK. Methodology for retinal photography and assessment of diabetic retinopathy: the EURODIAB IDDM complications study. *Diabetologia*. 1995;38:437–444.

## APPENDIX A

### A LIST OF PUBLIC DATA REPOSITORIES FOR RETINAL IMAGE ANALYSIS

This appendix lists the public retinal datasets known to us. Unless otherwise stated, all datasets listed are easily reachable by a Google search. Most descriptions are excerpts from the Web sites listed.

Structured Analysis of the Retina (STARE; available in the public domain at <http://www.ces.clemson.edu/~ahoover/stare/>) is one of the earliest and most often cited test sets in the ARIA literature, created for validating OD location. It consists of 31 images of healthy retinas and 50 images of retinas with disease, acquired using a Topcon TRV-50 fundus camera at 35° FOV and subsequently digitized at 605 × 700 pixels in resolution, 24 bits per pixel (standard RGB). The nerve is visible in all 81 images, although partially visible in 14 as appearing on the image border. In 5 images the nerve is completely obscured by hemorrhaging.

Digital Retinal Images for Vessel Extraction (DRIVE; available in the public domain at <http://www.isi.uu.nl/Research/Databases/DRIVE/>) is another much cited test set; it was created to enable comparative studies on segmentation of blood vessels in retinal images. The photographs were obtained from a DR screening program in The Netherlands. The screening population consisted of 400 diabetic subjects between 25 and 90 years of age. Forty photographs were randomly selected, 33 without and 7 with DR signs. The images were acquired using a Canon CR5 nonmydriatic 3CCD camera with a 45° FOV. Each image was captured using 8 bits per color plane at 768 by 584 pixels. The FOV of each image is circular with a diameter of approximately 540 pixels.

Diabetic Retinopathy Database and Evaluation Protocol (DiaRetDB1; available in the public domain at <http://www2.it.lut.fi/project/imageret/diaretdb1/>) consists of 89 color fundus images. Eighty-four contain at least mild nonproliferative DR signs (microaneurysms) and 5 are considered normal, not containing DR signs according to all experts who participated in the evaluation. Images were captured using the same 50° FOV digital fundus camera with varying imaging settings. The data correspond to a good (not necessarily typical) practical situation, where images are comparable and can be used to evaluate the general performance of diagnostic methods. Four medical experts were asked to mark the areas related to the microaneurysms, hemorrhages, and hard and soft exudates. Ground truth confidence levels (<50%, ~>50%, ~100%), representing the certainty of the decision that a marked finding is correct, are included.

Méthodes d'Évaluation de Systèmes de Segmentation et d'Indexation Dédiées à l'Ophthalmologie Rétinienne (MESSIDOR; available in the public domain at <http://messidor.crihan.fr/download-en.php>) contains 1200 eye fundus color digital images of the posterior pole, acquired by three ophthalmologic departments using a color video 3CCD camera on a Topcon TRC NW6 nonmydriatic retinograph

with a 45° FOV, 8 bits per color plane, and resolutions of 1440 × 960, 2240 × 1488, or 2304 × 1536 pixels. Eight hundred images were acquired with pupil dilation (one drop of Tropicamide at 0.5%) and 400 without dilation. The 1200 images are packaged in three sets, one per ophthalmologic department. Each set is divided into four zipped subsets each containing 100 images in TIFF format and an Excel file (Microsoft, Redmond, WA) with medical diagnoses for each image. Currently there are no annotations (markings) on the images. Annotations by a single clinician for OD diameter and the fovea center, for the whole MESSIDOR set, have been made available by the Department of Electronic, Computer Systems and Automatic Engineering, University of Huelva, Spain (available in the public domain at [www.uhu.es/retinopathy/muestras/Provided\\_Information.zip](http://www.uhu.es/retinopathy/muestras/Provided_Information.zip)).

REVIEW (available in the public domain at <http://reviewdb.lincoln.ac.uk/>) contains several subsets, with a mix of patients with disease and no disease, and has a focus on validating accurate measurements. It includes 16 images with 193 vessel segments, demonstrating a variety of pathologies and vessel types. These image sets contain 5066 manually marked profiles. Images were assessed by three independent experts, who marked the vessel edges.

AREDS (Age-Related Eye Disease Study; available in the public domain at <https://web.emmes.com/study/areds/>, <http://www.areds2.org/>) is a major clinical trial sponsored by the National Eye Institute (NEI) at the National Institutes of Health (<http://www.nei.nih.gov/amd/>), involving several US centers working on AMD. The dataset includes several thousands of analog and digitized fundus images showing various stages of AMD. Longitudinal studies were performed over 10 years showing disease progression. The images were graded by national centers for AMD as well as for lens opacity. The ground truth does not include image-level delineation of drusen. The fundus photographs consist principally of 30° images including stereo images centered on temporal margin of the disc and an oblique view of the center of the macula near the temporal margin of the field, stereo images centered on the center of the macula, and monoscopic images centered temporal to the macula and offering an oblique view of the center of the macula near the nasal margin of the field.

ARIA (available in the public domain at [http://www.eyecharity.com/aria\\_online/](http://www.eyecharity.com/aria_online/)) contains color fundus images collected at St Paul's Eye Unit and the University of Liverpool, United Kingdom, as part of the ARIA project. All subjects were adults. All images were taken using a Zeiss FF450+ fundus camera, originally stored as uncompressed TIFF files and converted to compressed JPG files for World Wide Web publication. All photographs were taken at a 50° FOV. Blood vessel masks created by trained image analysis experts are available. The optic disc and fovea, where relevant, are outlined in separate file sets. The data are organized into three categories, namely, age-related macular degeneration subjects, healthy control group subjects, and diabetic subjects.

ROC (available in the public domain at <http://roc.healthcare.uiowa.edu/>) is a set of 100 digital color fundus photographs selected from a large dataset (150,000 images) acquired at multiple sites within the EyeCheck DR screening program (see ROC Web site references), marked as gradable by the screening program ophthalmologists and including microaneurysms. Three different types of images with different resolutions are included, acquired by a Topcon NW 100, a Topcon NW200, or a Canon CR5-45NM and resulting in two differently shaped FOVs. All images are JPEG, and compression was set in the camera. The substantial black background around the FOV present in the original type II and III images was cut off using specialized software. This complete set was

randomly split into a training and a test set, each containing 50 images. Four retinal experts, all from the Department of Ophthalmology at the University of Iowa, were asked to annotate all microaneurysms and all irrelevant lesions in all 100 images in the test and training sets.

BIOIMLAB (<http://bioimlab.dei.unipd.it/Data%20Sets.htm>) at the University of Padova, Italy, maintains a number of publicly available datasets for several measurements, including vessel tortuosity (60 images from normal and hypertensive patients; 30 images of retinal arteries of similar length and caliber, 30 images of retinal veins of similar length and caliber, MATLAB [Mathworks, Natick, MA] data structures).

HEI-MED (available in the public domain at <http://vibot.u-bourgogne.fr/luca/heimed.php>) is a collection of 169 fundus images to train and test image-processing algorithms for the detection of exudates and diabetic macular edema. The images have been collected as part of a telemedicine network for DR diagnosis. The images contain manual segmentation of exudation, and include a machine segmentation of the vascular tree and optic nerve locations. The dataset contains a mixture of ethnic groups, with roughly 60% African American, 25% Caucasian, and 11% Hispanic.

## APPENDIX B

This paper is the result of an international collaboration to which a total of 41 people contributed. Editorial limits on the number of authors prevented us from listing the following contributors: Adria Perez Rovira, Kristis Zutis, Khai Sing Chin (VAMPIRE project, School of Computing, University of Dundee, Dundee, United Kingdom); Kenneth W. Tobin, who pioneered ARIA work at ORNL, and Hector Santos-Villalobos (Oak Ridge National Laboratory, Oak Ridge, TN); Enrico Grisan, Enea Poletti (Department of Information Engineering, University of Padova, Padova, Italy); David Reed, Christopher Gee (Jules Stein Eye Institute, Los Angeles, CA); Andrew Hunter (REVIEW Group, University of Lincoln, Lincoln, United Kingdom); Tien Yin Wong, M. Kamran Ikram (SERI, Singapore); Jiang Liu, Ngan-Meng Tan (A\*STAR, Singapore); Meindert Nijemeier (Department of Biomedical Engineering, University of Iowa, Iowa City, IA); Wynne Hsu, Mong Li Lee (National University of Singapore, Singapore); Hao Hao, Behzad Alihamad, Ganesh Naik (RMIT University, Melbourne, Australia); David E. Freund (Applied Physics Laboratory, Johns Hopkins University, Laurel, MD); Juan Xu (Department of Ophthalmology, University of Pittsburgh School of Medicine, Pittsburgh, PA).