Development and validation of a deep-learning algorithm for the detection of neovascular age-related macular degeneration from colour fundus photographs

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Abstract

Importance: Detection of early onset neovascular age-related macular degeneration (AMD) is critical to protecting vision.

Background: To describe the development and validation of a deep-learning algorithm (DLA) for the detection of neovascular age-related macular degeneration.

Design: Development and validation of a DLA using retrospective datasets.

Participants: We developed and trained the DLA using 56,113 retinal images and an additional 86,162 images from an independent dataset to externally validate the DLA. All images were non-stereoscopic and retrospectively collected.

Methods: The internal validation dataset was derived from real-world clinical settings in China. Gold standard grading was assigned when consensus was reached by three individual ophthalmologists. The DLA classified 31,247 images asgradable and 24,866 as ungradable (poor quality or poor field definition). These ungradable images were used to create a classification model for image quality. Efficiency and diagnostic accuracy were tested using 86,162 images derived from the Melbourne Collaborative Cohort Study. Neovascular AMD and/or ungradable outcome in one or both eyes was considered referable.

Main Outcome Measures: Area under the receiver operating characteristic curve (AUC), sensitivity and specificity.

Results: In the internal validation dataset, the AUC, sensitivity and specificity of the DLA for neovascular AMD was 0.995, 96.7%, 96.4%, respectively. Testing against the independent external dataset achieved an AUC, sensitivity and specificity of 0.967, 100% and 93.4%, respectively. More than 60% of false positive cases displayed other macular pathologies. Amongst the false negative cases (internal validation dataset only), over half (57.2%) proved to be undetected detachment of the neurosensory retina or RPE layer.
Conclusions and Relevance: This DLA shows robust performance for the detection of neovascular AMD amongst retinal images from a multi-ethnic sample and under different imaging protocols. Further research is warranted to investigate where this technology could be best utilized within screening and research settings.

KEYWORDS
deep-learning algorithm, age-related macular degeneration, retinal-imaging

1 | INTRODUCTION

Age-related macular degeneration (AMD) is a leading cause of severe vision loss amongst developed nations, particularly in those aged 65 years and older. With the aging population, it is projected that 288 million people globally will have AMD by 2040. There are two types of advanced AMD, atrophic (“dry”) and neovascular (“wet”), with the latter having been responsible for an estimated 90% of severe vision loss cases. Effective therapeutic options are currently only available for neovascular AMD, with anti-vascular endothelial growth factor (VEGF) agents effective in reducing vision loss and, in many cases, restoring vision.

Detection of early onset neovascular AMD and timely treatment is essential for the protection of visual function. Colour fundus photography is a common imaging tool utilized in primary eye care and screening settings and is effective for the diagnosis of AMD. Despite this, accurate interpretation of photographs is dependent on highly trained personnel, limiting its utility in underserved settings, such as developing countries and minority underserved populations. Furthermore, amongst high throughput settings (eg, screening, epidemiological research), most images captured are normal and therefore manual grading of each image is a cumbersome task requiring an immense number of human graders.

Convolutional neural networks (CNN), a popular deep-learning model, have recently been applied to common ophthalmic diseases, including AMD, with promising results of disease identification from fundus photos. Despite this, most previously reported systems adopt conservative definitions for referable AMD (eg, intermediate AMD), which, given the lack of treatment options for atrophic and earlier stages of the disease, may create strain on eye care systems in low resource countries (eg, China, India, most developing countries). Furthermore, these systems have rarely been validated amongst fundus image datasets that closely resemble real-world screening, where image quality varies considerably with different imaging protocols and retinal pigmentation differs substantially amongst ethnicities. These factors constitute a notable source of potential error for deep-learning algorithms (DLA), and must therefore be considered when planning robust evaluations.

The objective of the present study is to describe the development and validation of a DLA for the detection of neovascular AMD using a dataset of over 50 000 retinal photographs, collected from a range of retinal camera models and clinical settings in China. Additionally, we evaluated the accuracy and efficiency of the DLA in a large (>80 000 images), external dataset of non-mydriatic images from a predominantly Caucasian ethnicity.

2 | METHODS

This study was approved by the Institutional Review Boards of the Zhongshan Ophthalmic Center, China (2017KYPJ049), the Royal Victorian Eye and Ear Hospital Human Research Ethics Committee (HREC-14/1199H) and conducted in accordance with the Declaration of Helsinki as revised in 2013.

2.1 Development of the DLA

The DLA was developed using 56 113 deidentified, original colour fundus photographs acquired from a web-based, cloud sourcing platform (http://www.labelme.org, Guangzhou, China). These macular and disc-centred images were acquired from clinic-based settings and contributed by a total of 36 ophthalmology departments, optometry clinics and screening settings in China. All retinal photographs in the training dataset were captured with a variety of common conventional desktop retinal cameras (eg, Topcon, Canon, Heidelberg and Digital Retinography System), using a variety of imaging protocols. Twenty-one ophthalmologists were enrolled as graders only after meeting a high level of agreement (unweighted Kappa ≥0.70) with an experienced ophthalmologist in a test set of 60 images (20 images of normal fundi, 20 early to late dry AMD and 20 neovascular AMD).

Retinal photographs were graded between October 2016 and March 2017. To ensure an accurate diagnosis of AMD, a multi-step method was undertaken. First, images from the total dataset (n = 56 113) were randomly assigned to a single ophthalmologist for grading. Once grading was complete, the image was assigned to a separate ophthalmologist until three consistent grading outcomes were achieved for a given image. At this time, a consensus grading outcome was
given to the image. Graders were masked to the previous image grading outcomes and a given image could only be assigned to a specific grader once. AMD was graded according to the Beckman clinical classification system that has been described in detail elsewhere. In brief, patients were categorized as no AMD, early, intermediate AMD or late AMD, atrophic or neovascular. An image was defined as “poor quality” if the vessels within the macular region could not be identified or ≥50% of the macular region was obscured. Images that did not include the macular region were classified as poor field definition. Figure 1 and Table 1 describe the process of image grading and the classification of AMD using the online LabelMe platform.

The deep-learning classification approach used to train our DLA from image pre-processing to AMD classification is shown in Figure 2. Firstly, several automated pre-processing steps were performed for normalization to control for variations in image size and resolution. This included; (a) applying local space average colour for colour constancy; (b) downsizing images to a resolution of 299 × 299 pixels; and (c) online data augmentation to enlarge heterogeneity but keep the prognostic features in the image by a random horizontal shift of approximately 0-3 pixels and 90°, 180° or 270° random rotation. The study included three deep-learning models, all using inception-v3 architecture. This included networks for the (a) classification for referable AMD (late-wet AMD), (b) assessment of image quality and (c) assessment of the visual availability of the macular region (field definition).

2.2 Validation datasets

Using identical pre-processing procedures, the performance of the DLA was internally and externally assessed. Using our local LabelMe dataset, using an internal hold-out method, a total of 5554 images from the original dataset of 56,113 were used for internal validation. The consensus grading outcome from the ophthalmologists provided the gold standard for which the DLA was compared to. An experienced ophthalmologist (Z.L.) classified false positive and negative images into subgroups. External validation of the DLA was also assessed on an independent external dataset of 86,162 images of 21,777

<table>
<thead>
<tr>
<th>Classification</th>
<th>Presence of clinical features</th>
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<tbody>
<tr>
<td>Absent</td>
<td>Does not meet any of the following criteria</td>
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<tr>
<td>Early/intermediate</td>
<td>Any criterion of the following:</td>
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<tr>
<td></td>
<td>Drusen &gt; 63 μm</td>
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<td></td>
<td>RPE abnormalities (ie, hyperpigmentation or depigmentation)</td>
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<tr>
<td>Late dry</td>
<td>Any geographic atrophy</td>
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<tr>
<td>Late wet</td>
<td>Any criterion of the following:</td>
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<td></td>
<td>Serous detachment of the sensory retina or RPE</td>
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<td></td>
<td>Sub-RPE retinal haemorrhage</td>
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<tr>
<td></td>
<td>Sub-retinal/sub-RPE fibrovascular proliferation</td>
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<tr>
<td></td>
<td>Disciform scar</td>
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<tr>
<td>Poor quality</td>
<td>Any criterion of the following:</td>
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<tr>
<td></td>
<td>Vessels within the macular area cannot be identified</td>
</tr>
<tr>
<td></td>
<td>≥50% of the area is obscured</td>
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<tr>
<td>Poor field definition</td>
<td>Not macular-centred photographs</td>
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*Lesions were assessed in the circle area within 2-disc diameters of fovea. Abbreviation: RPE, retinal pigment epithelium.

**FIGURE 1** Image grading process of internal validation dataset. AMD, age-related macular degeneration.
participants, derived from the Melbourne Collaborative Cohort Study (MCCS). The participants assessed for AMD in the MCCS in 2003 to 2007 were aged 47 to 86 years (mean age 65.3 years, 60.1% female), 14% were born in Southern Europe (Greece, Italy or Malta), with the remaining 86% of Anglo-Celtic origin, born in Australia, the United Kingdom or New Zealand. In the MCCS, digital, non-stereoscopic 45° fundus photography of the macular and optic disc were taken of each eye, using a Canon CR6-45NM non-mydriatic retinal camera with a digital Canon (D60) camera back (Canon Inc., Kanagawa, Japan). All images were non mydriatic. Experienced professional graders (K.Z.A. and G.M.) from the Centre for Eye Research Australia graded each MCCS image and were masked to the identity and clinical characteristics of study participants. Any uncertain cases were adjudicated by a senior retinal specialist (RG) from Australia. For all participants, a single manual grading outcome based on worse affected eye was provided per participant. Overall, 2694 (12.7%) had early stages of AMD (one or more drusen ≥ 125 μm in size or one or more drusen with 63-124 μm in size with pigmentary abnormalities in a 600-μm diameter grading grid), 122 (0.6%) had late AMD (geographic atrophy or neovascular AMD). Participants with an ungradable manual grading result in both eyes were excluded from the dataset for the purpose of analysis.

DLA grading of MCCS images was undertaken independently of the research team involved in the development of the software. First, images without the manual grading label were transferred by MCCS investigators to the research team, who subsequently processed DLA grading on three computers operating concurrently, using a custom DLA software that allows automated classification consecutively on a set of images. Following this, DLA grading outcomes were transferred to MCCS investigators for preliminary evaluation and identification of discordant cases. Lastly, all discordant cases were assessed by experienced ophthalmologists (X.X.Y. and L.R.) and any cases identified as suspects for neovascular AMD were presented at a consensus meeting for adjudication by the senior ophthalmologists (R.G. and M.H.). Given, in the majority of cases, multiple images were available per eye, we adopted the following logic to consolidate a single automated grading result for the right and left eye; (a) positive late-wet AMD = any image for a given eye was found to be positive on automated grading; (b) negative late-wet AMD = no image was found to be positive on automated grading and at least one image for a given eye was found to be negative; and (c) ungradable = all images for a given eye were ungradable on automated grading. Referable AMD was defined as neovascular AMD and/or ungradable outcome in one or both eyes. The inclusion of ungradable cases as referable is in line with previous reports and more closely resembles real-word circumstances, where both positive and ungradable cases go on to manual verification.

**FIGURE 2** The deep convolutional neural network used in this study. Data stream is from left to right. A fundus photograph is firstly preprocessed by scaling, subtraction of local space average colour, downsizing the image to a 299 × 299 matrix and data augmentation for normalization. The image is sequentially warped into probability distributions over whether referable AMD is present using Inception-v3 convolutional neural network architecture training from scratch on the training dataset and validation dataset.
2.3 | Convolutional neural network visualization

To visualize the learning procedure of our networks, we applied an Adaptive Kernel Visualization technique. In brief, this involved applying a sliding window size of 28 x 28 pixels, with stride of three pixels, to crop images into smaller sub-images and produce a \((544-28)/3 \times (544-28)/3\) = 172*172 feature map. A random sample of 100 true positive images from the internal validation dataset were selected and utilized as inputs for the trained neovascular AMD deep-learning models. The threshold was set at 0.5 for the late-wet AMD model, meaning that discriminative image regions were highlighted if the classification possibility output of being diagnosed was greater than 50%.

2.4 | Statistical analysis

The sensitivity, specificity, accuracy and area under the curve (AUC) of the DLA in detecting neovascular AMD was performed compared to the reference standard (local validation = retinal specialist; external validation = professional graders) for each participant. The 95% confidence intervals (CIs) were also calculated. STATA version 14.0 (College Station, Texas) was used for all statistical analyses in this study.

3 | RESULTS

Each image in the training and internal validation dataset was graded between three and 10 times before consensus agreement was reached, with a mean agreement rate of 86.5% (95% CI, 84.5%-88.5%) for the 21 ophthalmologists. Each ophthalmologist graded between 397 and 33513 (median, 4135) fundus photographs, with 12 ophthalmologists individually grading more than 5000 fundus photographs. This considerable variation between the number of images graded existed because ophthalmologists volunteered their time to perform retinal image grading. Of the total 56113 images, 7723 (13.8%) were labelled as poor quality and 17143 (30.5%) labelled as poor field definition (i.e., macular region not in view), leaving 31247 images with a conclusive AMD classification. Using a simple random sampling method, a total of 27397 images were assigned to the training dataset and the remaining 3850 images were held-out for internal validation. A subset of 18704 images from the entire dataset were used to develop the network for image quality and field definition. Amongst the 27397 images in the training dataset, 22553 (82.3%) had no AMD, 1338 (4.9%) had early or intermediate AMD, 72 (0.3%) had atrophic AMD and 3434 (12.5%) had neovascular AMD. Investigators purposefully oversampled neovascular AMD cases through targeted image collection in patients undergoing fundus fluorescein angiography in Chinese tertiary hospitals. Table 2 summarizes the characteristics of fundus photographs in the training and validation datasets.

3.1 | Internal hold-out-validation

In the internal hold out validation dataset (reference to ophthalmologist standard), the AUC, sensitivity, specificity and accuracy of the DLA for neovascular AMD was 0.995 (95% CI, 0.993-0.997), 98.0%, 94.0% and 94.7%, respectively (Figure 3). The AUC, sensitivity, specificity and accuracy for image quality and field definition was 0.995 (95% CI, 0.992-0.997), 96.7%, 96.4% 96.5%, respectively. Typical examples of the visualization maps for true positive cases are shown in Figure 4.

The most common clinical features of false negative cases (n = 14) included serous detachment of the sensory retina or retinal pigment epithelium (RPE) (n = 8, 57.2%), followed by sub-retinal/sub-RPE fibrovascular proliferation (n = 3, 21.4%). An analysis of false positive cases revealed that 145 (76.3%) had other eye disorders such as diabetic retinopathy (n = 38, 20%) and myopic maculopathy

<table>
<thead>
<tr>
<th>Characteristics of fundus photographs in the training set and randomized validation sample a</th>
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<tbody>
<tr>
<td>Training set</td>
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<tr>
<td>----------------</td>
</tr>
<tr>
<td>No AMD</td>
</tr>
<tr>
<td>Early or intermediate AMD</td>
</tr>
<tr>
<td>Late dry AMD</td>
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<tr>
<td>Late wet AMD</td>
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<tr>
<td>Total</td>
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aData are presented as n (%) unless otherwise indicated. AMD, age-related macular degeneration.

**FIGURE 3** The performance of this deep-learning algorithm in detection age-related macular disease. The area under receiver operating characteristic curve (AUC) of 0.995 (95% confidence interval [CI], 0.993-0.997) was obtained.
(n = 22, 11.5%). Normal fundus photographs without artefacts (n = 34, 17.9%) and those with artefacts (n = 11, 5.8%) made up the remaining false positive cases. Examples of the typical false negative and false positive images can be found in Figures 5 and 6.

3.2 | External validation

The external dataset contained 86 202 retinal images of 21 777 participants from the MCCS. Of these, there were 335 (1.54%) participants who had missing or ungradable manual grading outcomes in both eyes that were excluded from analysis. Amongst the remaining 21 327 participants included in the external validation dataset 48 (0.2%) had late-wet AMD.

When adopting the aforementioned definition for referable AMD (ie, neovascular AMD and/or ungradable outcome in one or both eyes), the AUC, sensitivity and specificity of the DLA in the external validation dataset was 0.967, 100% and 92.6%, respectively. This consisted of

**FIGURE 4** AMD true positive. Images A1, B1 & C1 show original images without heat-map. A2 shows heatmap predominately visualizing the macular, temporal side of the optic nerve head (ONH) and retinal vessels (superiorly). B2 shows heatmap visualizing the macular region and C2 heatmap shows an area of atrophy superior to the central macular and superior vessels predominantly being visualized. AMD, age-related macular degeneration

**FIGURE 5** Typical cases of false negative findings in the detection age-related macular disease. A, Serous detachment of the sensory retina or RPE; B, sub-retinal/sub-RPE fibrovascular proliferation; C, sub-RPE retinal haemorrhage with serous detachment of the sensory retina or RPE. RPE, retinal pigment epithelium
true positives, 322 false positive cases and 1281 ungradable cases in one or both eyes. In total, five neovascular AMD cases were labelled as ungradable in DLA analysis. Although not in line with real-world practices, if we were to class these five cases as false negatives, the adjusted sensitivity metric would reduce from 100% to 89.6%. Of note, adjudication of discordant manual and DLA grading outcomes via consensus of senior ophthalmologists, revealed that the DLA correctly classified four cases as negative that had originally been labelled as positive for neovascular AMD by MCCS manual graders. This misclassification was confirmed in the adjudication consensus meeting attended by senior ophthalmologists.

An analysis of false positives revealed that 62.7% (n = 202/322) of cases displayed signs of other macular pathology, of which the most common findings were early AMD (n = 84/322, 26.1%), atrophic AMD (n = 24/322, 7.5%), myopic maculopathy (n = 16/322, 5.0%), retinal scarring or irregular macular lesions (n = 16/322, 5%) and vitreous opacity (n = 5/322, 1.6%). The remaining 37.3% (n = 120/322) of false positive cases had no abnormal macular findings, amongst them, 46.7% (n = 56/120) of these display image artefacts.

In total, it took about 48 hours (eg, six working days of 8 hours) to complete DLA grading using three computers operating concurrently. To assess repeatability and reliability of the DLA, automated grading was repeated on a random sample of 2000 images from the MCCS dataset. In this subset evaluation, 100% grading consistency was observed.

**FIGURE 6** Typical cases of false positive findings. A, Diabetic retinopathy; B, myopic maculopathy; C, early or intermediate AMD; D, choroiditis; E, normal retina; F, normal image with artefacts. AMD, age-related macular degeneration

**DISCUSSION**

This paper describes the development of a DLA for the detection of neovascular AMD based on a large specialist annotated dataset of 56113 images collected amongst a Chinese population. Amongst an independent, local validation dataset of 5554 images (including 18.4% neovascular AMD cases) derived from a range of providers and camera models, our DLA achieved robust performance for neovascular AMD (AUC = 0.995). Furthermore, the DLA showed excellent efficiency and diagnostic performance (AUC = 0.967) in a large (>80 000 images), external dataset of non-mydriatic images from participants of Southern European and Anglo-Celtic origin.

Over the past decade, automated techniques for the assessment of AMD, via feature extraction from small retinal image datasets (<1000), have been reported with variable accuracy. More recent reports provide novel data on the accuracy of deep-learning systems for the detection of AMD. The majority of these studies have utilized the Age-Related Eye Disease Study (AREDS) participants to develop and test the accuracy of their DLAs. The AREDS dataset is useful for training and internally validating DLAs as the images are of high-quality (pupillary dilation ≥5 mm), are collected within research settings and have already been gold standard graded by experts. However, AREDS participants were excluded at recruitment if they had unrelated sight threatening disease. It has been suggested that many CNN’s perform sub optimally when tested on unrelated datasets because of technical
differences including camera setup illumination and inclusion criteria.\footnote{21} In comparison, our DLA was trained using images from multiple clinical settings using various retinal cameras and imaging protocols, and was able to achieve excellent diagnostic accuracy.

Performing external validation is therefore essential given that image quality imperfections are common in real-world screening settings and various camera models are used. The study by Ting et al (2017)\textsuperscript{11} did not perform an external validation for their referable AMD DLA and of those studies that used the AREDS dataset, only Grassman et al (2018)\textsuperscript{5} performed an external validation, albeit on a small (5555) independent dataset. For the external validation, the algorithm was able to detect 84.2\% of all fundus images with signs of early or late AMD. However, this was only achieved when those aged under 55 years (due to visualization of the macular reflex causing false positives) and images with pathology not related to AMD were excluded. In the current study, only those with missing or ungradable manual grading outcomes in both eyes were excluded from the analysis making the dataset more generalizable to real-world screening settings.

Direct comparison between recent reports relating to AMD detection using DLAs is made difficult due to differing classification criteria. Burlina et al (2018),\textsuperscript{20} Grassman et al (2018)\textsuperscript{5} and Burlina et al (2017)\textsuperscript{3} utilized multi-step approaches using classifications developed for AREDS. For example, Grassman et al\textsuperscript{5} defined 13 classes, one indicating little or no AMD, grades 2 to 9 representing changes associated with early or intermediate AMD and grades 10 to 12 covering late-stage AMD such as geographic atrophy and neovascular AMD. Greater numbers of classification groups led to lower kappa scores in the study by Burlina et al (2018)\textsuperscript{20} (0.77 for four-step approach and 0.74 in the nine-step approach) and an overall accuracy of 63\% in the study by Grassman et al (2018).\textsuperscript{5} Similarly, this was shown by Burlina et al (2017)\textsuperscript{3} who obtained accuracy values of 79.4\%, 81.5\% and 93.4\% for 4-class, 3-class and 2-class classifications, respectively. Rather than using a multi-step approach, Peng et al (2018)\textsuperscript{19} detected individual AMD risk factors including drusen, pigmentedary changes and late AMD. Although they used late AMD as a classifier, this included geographic atrophy which our DLA does not classify as referable AMD. A binary classification was used to determine referable AMD (≥intermediate AMD) in the investigations by Ting et al (2017)\textsuperscript{11} and Burlina et al (2017)\textsuperscript{4} who both reported AUC’s between 0.93 and 0.96. Whilst these findings are similar to the current study, they are not directly comparable as only those classified as neovascular AMD we considered referable. The choice to only include neovascular AMD as referable was due to the lack of effective treatment options for atrophic and earlier forms of AMD. Therefore, if deployed in low recourse settings, previously reported DLAs may increase the strain on eye care resources by over-referring many cases.

Taking the current literature into consideration, we developed a DLA to detect neovascular AMD and robustly evaluated its performance in two datasets of images that were taken under varying imaging protocols and across different ethnicities. In order to exhaust all possible variations of neovascular AMD phenotypes, we adopted a training dataset that included a large number (n = 3434) of neovascular AMD cases collected from over 30 clinical settings using different fundus camera and imaging protocols. In the large external validation dataset, the DLA demonstrated reliable automated image analysis under a non-mydriatic retinal protocol, achieving robust diagnostic accuracy (AUC = 0.967) and an ungradable image rate of only 5.1\%. We hypothesize that the ungradable image rate may be lower if the DLA was deployed within a prospective screening setting, given the software has built in automated classifiers for image quality and field definition that would prompt real-time image re-capture. Our DLA was also successfully validated amongst two ethnic groups (Caucasian AUC = 0.967; Chinese AUC = 0.995) with distinct retinal pigmentation. While we acknowledge that further evaluation is warranted amongst darker skinned ethnic groups, these results provide evidence that the performance of our DLA is likely generalizable to a large number of populations globally.

An important potential benefit of deep-learning technology relates to the ability to grade retinal images at an extraordinary speed and scale. In the present study, it took 48 hours to complete DLA grading of the 86 162 images from the MCCS dataset using three computers operating concurrently. In comparison, experienced human graders took several months of dedicated time to complete this task, albeit adopting a greater depth of classification (manual grading = 5-class vs DLA = 2-class). In total, the DLA correctly classified 93.8\% of participants with healthy fundus images and only 6.8\% of participants, including all neovascular AMD cases, were classified as positive (n = 43) or ungradable (n = 5) by the DLA. This represents a significant potential workload and cost saving if the DLA were deployed as a pre-screening tool in high throughput screening or research settings, where only positive and ungradable cases flagged by automated grading go on to manual verification. In addition, our finding that the DLA correctly classified five cases in the external validation dataset that had been previously misclassified by manual graders as positive for neovascular AMD is noteworthy.

Despite the promising accuracy and efficiency of the DLA, we recognize that one of the major challenges to clinical adoption relates to a major mind-set shift in how clinicians entrust clinical care to machines. To assist in the clinical acceptance,\textsuperscript{22} we explored characteristics of misclassification (false negatives and false positives) of the DLA. We found that over 60\% of false positive cases in the internal and external validation datasets displayed other macular pathologies, suggesting that most of these false positive cases may have in fact benefited from referral. Amongst the false negative cases
(internal validation dataset only), over half (57.2%) proved to be undetected sensory detachment of neurosensory retina or RPE layer. This is perhaps not a surprising finding given the relative inconspicuous nature of this lesion when compared to other neovascular AMD phenotypes such as fibrovascular or scarring changes. Further training of the DLA with more image examples of this lesion may improve the diagnostic accuracy. In addition to exploring reasons for misclassification, we developed an Adaptive Kernel Visualization method that enables the most discriminative image regions of the DLA to be discerned. This tool offers great potential to alleviate the previously existing tension between accuracy and interpretability of these systems by enabling clinicians to understand important exposure variables in real-time.

The strengths of this study include the utilization of a large training dataset (>50,000 images) of gold standard labelled images, the independent assessment of the DLA on an external dataset, and the robust performance (ie, accuracy and efficiency) of the DLA on diverse datasets containing images from multiple cameras, under different imaging protocols and across two ethnicities. Some limitations must also be considered. First, there was a relatively small representation of neovascular AMD cases in the external validation dataset (n = 48), which may have resulted in an unstable estimate of diagnostic accuracy. Despite this, similar metrics were observed on the internal hold-out validation dataset that contained in excess of 700 neovascular AMD cases. Second, we suspect that a large proportion of DLA screen-positive cases in the internal and external validation datasets would have end-stage lesions (eg, fibrosis and atrophy), and therefore be ineligible for treatment. Identification of new onset neovascular AMD cases or the existence of choroidal neovascularisation for treatment would rely on confirmation via optical coherence tomography (OCT). Lastly, given that it is not considered cost-effective to screen for AMD in the general population, the usefulness of the evaluated DLA in screening settings may be questioned. It is important to note that we have also developed DLAs to detect retinal infarction (GON),23 the usefulness of the DLA in screening settings may be questioned. It is important to note that we have also developed DLAs to detect referable diabetic retinopathy (DR) and glaucomatous optic neuropathy (GON),24-26 which can be deployed concurrently with the DLA described in the present report. Therefore, given the evaluation of AMD is typically included within manual screening programs amongst the diabetic population, we speculate that, at a minimum, the DLA in question could be discerned. This tool offers great potential to alleviate the previously existing tension between accuracy and interpretability of these systems by enabling clinicians to understand important exposure variables in real-time.

In conclusion, this DLA shows robust performance for the detection of neovascular AMD amongst retinal images from a multi-ethnic sample and under different imaging protocols. The results suggest that this system may provide an efficient and cost-effective “pre-screener” when applied to large research datasets. Future efforts will focus on investigating the effectiveness of our DLA for neovascular AMD as a screening tool in high risk populations, when combined with our other fundus-image based DLAs for GON and DR, and what impact the introduction OCT imaging has as a second-line screening tool amongst screen-positive patients.

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CONFLICT OF INTEREST

The authors have no financial or other conflicts of interest concerning this study. Mingguang He and Wei Meng report a patent on managing colour fundus images using deep-learning models. The patent application number was ZL201510758675.5 and patent filing date was 31 May 2017.

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